Pearson New International Edition

Computer Graphics with Open GL
Hearn    Baker    Carithers
Fourth Edition
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The power and utility of computer graphics is widely recognized, and a broad range of graphics hardware and software systems is now available for applications in virtually all fields. Graphics capabilities for both two-dimensional and three-dimensional applications are now common, even on general-purpose computers and handheld calculators. With personal computers, we can use a variety of interactive input devices and graphics software packages. For higher-quality applications, we can choose from a number of sophisticated special-purpose graphics hardware systems and technologies. In this chapter, we explore the basic features of graphics hardware components and graphics software packages.
Video Display Devices

Typically, the primary output device in a graphics system is a video monitor. Historically, the operation of most video monitors was based on the standard cathode-ray tube (CRT) design, but several other technologies exist. In recent years, flat-panel displays have become significantly more popular due to their reduced power consumption and thinner designs.

Refresh Cathode-Ray Tubes

Figure 1 illustrates the basic operation of a CRT. A beam of electrons (cathode rays), emitted by an electron gun, passes through focusing and deflection systems that direct the beam toward specified positions on the phosphor-coated screen. The phosphor then emits a small spot of light at each position contacted by the electron beam. Because the light emitted by the phosphor fades very rapidly, some method is needed for maintaining the screen picture. One way to do this is to store the picture information as a charge distribution within the CRT. This charge distribution can then be used to keep the phosphors activated. However, the most common method now employed for maintaining phosphor glow is to redraw the picture repeatedly by quickly directing the electron beam back over the same screen points. This type of display is called a refresh CRT, and the frequency at which a picture is redrawn on the screen is referred to as the refresh rate.

The primary components of an electron gun in a CRT are the heated metal cathode and a control grid (Fig. 2). Heat is supplied to the cathode by directing a current through a coil of wire, called the filament, inside the cylindrical cathode structure. This causes electrons to be "boiled off" the hot cathode surface. In
the vacuum inside the CRT envelope, the free, negatively charged electrons are then accelerated toward the phosphor coating by a high positive voltage. The accelerating voltage can be generated with a positively charged metal coating on the inside of the CRT envelope near the phosphor screen, or an accelerating anode, as in Figure 2, can be used to provide the positive voltage. Sometimes the electron gun is designed so that the accelerating anode and focusing system are within the same unit.

Intensity of the electron beam is controlled by the voltage at the control grid, which is a metal cylinder that fits over the cathode. A high negative voltage applied to the control grid will shut off the beam by repelling electrons and stopping them from passing through the small hole at the end of the control-grid structure. A smaller negative voltage on the control grid simply decreases the number of electrons passing through. Since the amount of light emitted by the phosphor coating depends on the number of electrons striking the screen, the brightness of a display point is controlled by varying the voltage on the control grid. This brightness, or intensity level, is specified for individual screen positions with graphics software commands.

The focusing system in a CRT forces the electron beam to converge to a small cross section as it strikes the phosphor. Otherwise, the electrons would repel each other, and the beam would spread out as it approaches the screen. Focusing is accomplished with either electric or magnetic fields. With electrostatic focusing, the electron beam is passed through a positively charged metal cylinder so that electrons along the center line of the cylinder are in an equilibrium position. This arrangement forms an electrostatic lens, as shown in Figure 2, and the electron beam is focused at the center of the screen in the same way that an optical lens focuses a beam of light at a particular focal distance. Similar lens focusing effects can be accomplished with a magnetic field set up by a coil mounted around the outside of the CRT envelope, and magnetic lens focusing usually produces the smallest spot size on the screen.

Additional focusing hardware is used in high-precision systems to keep the beam in focus at all screen positions. The distance that the electron beam must travel to different points on the screen varies because the radius of curvature for most CRTs is greater than the distance from the focusing system to the screen center. Therefore, the electron beam will be focused properly only at the center of the screen. As the beam moves to the outer edges of the screen, displayed images become blurred. To compensate for this, the system can adjust the focusing according to the screen position of the beam.

As with focusing, deflection of the electron beam can be controlled with either electric or magnetic fields. Cathode-ray tubes are now commonly constructed with magnetic-deflection coils mounted on the outside of the CRT envelope, as illustrated in Figure 1. Two pairs of coils are used for this purpose. One pair is mounted on the top and bottom of the CRT neck, and the other pair is mounted on opposite sides of the neck. The magnetic field produced by each pair of coils results in a transverse deflection force that is perpendicular to both the direction of the magnetic field and the direction of travel of the electron beam. Horizontal deflection is accomplished with one pair of coils, and vertical deflection with the other pair. The proper deflection amounts are attained by adjusting the current through the coils. When electrostatic deflection is used, two pairs of parallel plates are mounted inside the CRT envelope. One pair of plates is mounted horizontally to control vertical deflection, and the other pair is mounted vertically to control horizontal deflection (Fig. 3).

Spots of light are produced on the screen by the transfer of the CRT beam energy to the phosphor. When the electrons in the beam collide with the phosphor
coating, they are stopped and their kinetic energy is absorbed by the phosphor. Part of the beam energy is converted by friction into heat energy, and the remainder causes electrons in the phosphor atoms to move up to higher quantum-energy levels. After a short time, the “excited” phosphor electrons begin dropping back to their stable ground state, giving up their extra energy as small quanta of light energy called photons. What we see on the screen is the combined effect of all the electron light emissions: a glowing spot that quickly fades after all the excited phosphor electrons have returned to their ground energy level. The frequency (or color) of the light emitted by the phosphor is in proportion to the energy difference between the excited quantum state and the ground state.

Different kinds of phosphors are available for use in CRTs. Besides color, a major difference between phosphors is their persistence: how long they continue to emit light (that is, how long it is before all excited electrons have returned to the ground state) after the CRT beam is removed. Persistence is defined as the time that it takes the emitted light from the screen to decay to one-tenth of its original intensity. Lower-persistence phosphors require higher refresh rates to maintain a picture on the screen without flicker. A phosphor with low persistence can be useful for animation, while high-persistence phosphors are better suited for displaying highly complex, static pictures. Although some phosphors have persistence values greater than 1 second, general-purpose graphics monitors are usually constructed with persistence in the range from 10 to 60 microseconds.

Figure 4 shows the intensity distribution of a spot on the screen. The intensity is greatest at the center of the spot, and it decreases with a Gaussian distribution out to the edges of the spot. This distribution corresponds to the cross-sectional electron density distribution of the CRT beam.

The maximum number of points that can be displayed without overlap on a CRT is referred to as the resolution. A more precise definition of resolution is the number of points per centimeter that can be plotted horizontally and vertically, although it is often simply stated as the total number of points in each direction. Spot intensity has a Gaussian distribution (Fig. 4), so two adjacent spots will appear distinct as long as their separation is greater than the diameter at which each spot has an intensity of about 60 percent of that at the center of the spot. This overlap position is illustrated in Figure 5. Spot size also depends on intensity. As more electrons are accelerated toward the phosphor per second, the diameters of the CRT beam and the illuminated spot increase. In addition, the increased excitation energy tends to spread to neighboring phosphor atoms not directly in the path of the beam, which further increases the spot diameter. Thus, resolution of a CRT is dependent on the type of phosphor, the intensity to be displayed, and the focusing and deflection systems. Typical resolution on high-quality systems is 1280 by 1024, with higher resolutions available on many systems. High-resolution systems are often referred to as high-definition systems.
The physical size of a graphics monitor, on the other hand, is given as the length of
the screen diagonal, with sizes varying from about 12 inches to 27 inches or more.
A CRT monitor can be attached to a variety of computer systems, so the number
of screen points that can actually be plotted also depends on the capabilities of
the system to which it is attached.

Raster-Scan Displays
The most common type of graphics monitor employing a CRT is the raster-scan
display, based on television technology. In a raster-scan system, the electron beam
is swept across the screen, one row at a time, from top to bottom. Each row is
referred to as a scan line. As the electron beam moves across a scan line, the beam
intensity is turned on and off (or set to some intermediate value) to create a pattern
of illuminated spots. Picture definition is stored in a memory area called the
refresh buffer or frame buffer, where the term frame refers to the total screen area.
This memory area holds the set of color values for the screen points. These stored
color values are then retrieved from the refresh buffer and used to control the
intensity of the electron beam as it moves from spot to spot across the screen. In this
way, the picture is “painted” on the screen one scan line at a time, as demonstrated
in Figure 6. Each screen spot that can be illuminated by the electron beam
is referred to as a pixel or pel (shortened forms of picture element). Since the
refresh buffer is used to store the set of screen color values, it is also sometimes
called a color buffer. Also, other kinds of pixel information, besides color, are
stored in buffer locations, so all the different buffer areas are sometimes referred
to collectively as the “frame buffer.” The capability of a raster-scan system to
store color information for each screen point makes it well suited for the realistic
display of scenes containing subtle shading and color patterns. Home television
sets and printers are examples of other systems using raster-scan methods.

Raster systems are commonly characterized by their resolution, which is the
number of pixel positions that can be plotted. Another property of video monitors

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**FIGURE 6**
A raster-scan system displays an object as a set of discrete points across each
scan line.
is aspect ratio, which is now often defined as the number of pixel columns divided by the number of scan lines that can be displayed by the system. (Sometimes this term is used to refer to the number of scan lines divided by the number of pixel columns.) Aspect ratio can also be described as the number of horizontal points to vertical points (or vice versa) necessary to produce equal-length lines in both directions on the screen. Thus, an aspect ratio of 4/3, for example, means that a horizontal line plotted with four points has the same length as a vertical line plotted with three points, where line length is measured in some physical units such as centimeters. Similarly, the aspect ratio of any rectangle (including the total screen area) can be defined to be the width of the rectangle divided by its height.

The range of colors or shades of gray that can be displayed on a raster system depends on both the types of phosphor used in the CRT and the number of bits per pixel available in the frame buffer. For a simple black-and-white system, each screen point is either on or off, so only one bit per pixel is needed to control the intensity of screen positions. A bit value of 1, for example, indicates that the electron beam is to be turned on at that position, and a value of 0 turns the beam off. Additional bits allow the intensity of the electron beam to be varied over a range of values between “on” and “off.” Up to 24 bits per pixel are included in high-quality systems, which can require several megabytes of storage for the frame buffer, depending on the resolution of the system. For example, a system with 24 bits per pixel and a screen resolution of 1024 by 1024 requires 3 MB of storage for the refresh buffer. The number of bits per pixel in a frame buffer is sometimes referred to as either the depth of the buffer area or the number of bit planes. A frame buffer with one bit per pixel is commonly called a bitmap, and a frame buffer with multiple bits per pixel is a pixmap, but these terms are also used to describe other rectangular arrays, where a bitmap is any pattern of binary values and a pixmap is a multicolor pattern.

As each screen refresh takes place, we tend to see each frame as a smooth continuation of the patterns in the previous frame, so long as the refresh rate is not too low. Below about 24 frames per second, we can usually perceive a gap between successive screen images, and the picture appears to flicker. Old silent films, for example, show this effect because they were photographed at a rate of 16 frames per second. When sound systems were developed in the 1920s, motion-picture film rates increased to 24 frames per second, which removed flickering and the accompanying jerky movements of the actors. Early raster-scan computer systems were designed with a refresh rate of about 30 frames per second. This produces reasonably good results, but picture quality is improved, up to a point, with higher refresh rates on a video monitor because the display technology on the monitor is basically different from that of film. A film projector can maintain the continuous display of a film frame until the next frame is brought into view. But on a video monitor, a phosphor spot begins to decay as soon as it is illuminated. Therefore, current raster-scan displays perform refreshing at the rate of 60 to 80 frames per second, although some systems now have refresh rates of up to 120 frames per second. And some graphics systems have been designed with a variable refresh rate. For example, a higher refresh rate could be selected for a stereoscopic application so that two views of a scene (one from each eye position) can be alternately displayed without flicker. But other methods, such as multiple frame buffers, are typically used for such applications.

Sometimes, refresh rates are described in units of cycles per second, or hertz (Hz), where a cycle corresponds to one frame. Using these units, we would describe a refresh rate of 60 frames per second as simply 60 Hz. At the end of each scan line, the electron beam returns to the left side of the screen to begin displaying the next scan line. The return to the left of the screen, after refreshing
each scan line, is called the **horizontal retrace** of the electron beam. And at the end of each frame (displayed in \( \frac{1}{50} \) to \( \frac{1}{60} \) of a second), the electron beam returns to the upper-left corner of the screen (**vertical retrace**) to begin the next frame.

On some raster-scan systems and TV sets, each frame is displayed in two passes using an **interlaced** refresh procedure. In the first pass, the beam sweeps across every other scan line from top to bottom. After the vertical retrace, the beam then sweeps out the remaining scan lines (Fig. 7). Interlacing of the scan lines in this way allows us to see the entire screen displayed in half the time that it would have taken to sweep across all the lines at once from top to bottom. This technique is primarily used with slower refresh rates. On an older, 30 frame-per-second, non-interlaced display, for instance, some flicker is noticeable. But with interlacing, each of the two passes can be accomplished in \( \frac{1}{60} \) of a second, which brings the refresh rate nearer to 60 frames per second. This is an effective technique for avoiding flicker—provided that adjacent scan lines contain similar display information.

**Random-Scan Displays**

When operated as a **random-scan display** unit, a CRT has the electron beam directed only to those parts of the screen where a picture is to be displayed. Pictures are generated as line drawings, with the electron beam tracing out the component lines one after the other. For this reason, random-scan monitors are also referred to as **vector displays** (or **stroke-writing displays** or **calligraphic displays**). The component lines of a picture can be drawn and refreshed by a random-scan system in any specified order (Fig. 8). A pen plotter operates in a similar way and is an example of a random-scan, hard-copy device.

Refresh rate on a random-scan system depends on the number of lines to be displayed on that system. Picture definition is now stored as a set of line-drawing commands in an area of memory referred to as the **display list**, **refresh display file**, **vector file**, or **display program**. To display a specified picture, the system cycles through the set of commands in the display file, drawing each component line in turn. After all line-drawing commands have been processed, the system cycles back to the first line command in the list. Random-scan displays are designed to draw all the component lines of a picture 30 to 60 times each second, with up to 100,000 “short” lines in the display list. When a small set of lines is to be displayed, each refresh cycle is delayed to avoid very high refresh rates, which could burn out the phosphor.

Random-scan systems were designed for line-drawing applications, such as architectural and engineering layouts, and they cannot display realistic shaded scenes. Since picture definition is stored as a set of line-drawing instructions rather than as a set of intensity values for all screen points, vector displays generally have higher resolutions than raster systems. Also, vector displays produce smooth line
drawings because the CRT beam directly follows the line path. A raster system, by contrast, produces jagged lines that are plotted as discrete point sets. However, the greater flexibility and improved line-drawing capabilities of raster systems have resulted in the abandonment of vector technology.

Color CRT Monitors

A CRT monitor displays color pictures by using a combination of phosphors that emit different-colored light. The emitted light from the different phosphors merges to form a single perceived color, which depends on the particular set of phosphors that have been excited.

One way to display color pictures is to coat the screen with layers of different-colored phosphors. The emitted color depends on how far the electron beam penetrates into the phosphor layers. This approach, called the beam-penetration method, typically used only two phosphor layers: red and green. A beam of slow electrons excites only the outer red layer, but a beam of very fast electrons penetrates the red layer and excites the inner green layer. At intermediate beam speeds, combinations of red and green light are emitted to show two additional colors: orange and yellow. The speed of the electrons, and hence the screen color at any point, is controlled by the beam acceleration voltage. Beam penetration has been an inexpensive way to produce color, but only a limited number of colors are possible, and picture quality is not as good as with other methods.

Shadow-mask methods are commonly used in raster-scan systems (including color TV) because they produce a much wider range of colors than the beam-penetration method. This approach is based on the way that we seem to perceive colors as combinations of red, green, and blue components, called the RGB color model. Thus, a shadow-mask CRT uses three phosphor color dots at each pixel position. One phosphor dot emits a red light, another emits a green light, and the third emits a blue light. This type of CRT has three electron guns, one for each color dot, and a shadow-mask grid just behind the phosphor-coated screen. The
light emitted from the three phosphors results in a small spot of color at each pixel position, since our eyes tend to merge the light emitted from the three dots into one composite color. Figure 9 illustrates the *delta-delta* shadow-mask method, commonly used in color CRT systems. The three electron beams are deflected and focused as a group onto the shadow mask, which contains a series of holes aligned with the phosphor-dot patterns. When the three beams pass through a hole in the shadow mask, they activate a dot triangle, which appears as a small color spot on the screen. The phosphor dots in the triangles are arranged so that each electron beam can activate only its corresponding color dot when it passes through the shadow mask. Another configuration for the three electron guns is an *in-line* arrangement in which the three electron guns, and the corresponding RGB color dots on the screen, are aligned along one scan line instead of in a triangular pattern. This in-line arrangement of electron guns is easier to keep in alignment and is commonly used in high-resolution color CRTs.

We obtain color variations in a shadow-mask CRT by varying the intensity levels of the three electron beams. By turning off two of the three guns, we get only the color coming from the single activated phosphor (red, green, or blue). When all three dots are activated with equal beam intensities, we see a white color. Yellow is produced with equal intensities from the green and red dots only, magenta is produced with equal blue and red intensities, and cyan shows up when blue and green are activated equally. In an inexpensive system, each of the three electron beams might be restricted to either on or off, limiting displays to eight colors. More sophisticated systems can allow intermediate intensity levels to be set for the electron beams, so that several million colors are possible.

Color graphics systems can be used with several types of CRT display devices. Some inexpensive home-computer systems and video games have been designed for use with a color TV set and a radio-frequency (RF) modulator. The purpose of the RF modulator is to simulate the signal from a broadcast TV station. This means that the color and intensity information of the picture must be combined and superimposed on the broadcast-frequency carrier signal that the TV requires as input. Then the circuitry in the TV takes this signal from the RF modulator, extracts the picture information, and paints it on the screen. As we might expect, this extra handling of the picture information by the RF modulator and TV circuitry decreases the quality of displayed images.
Computer Graphics Hardware

Composite monitors are adaptations of TV sets that allow bypass of the broadcast circuitry. These display devices still require that the picture information be combined, but no carrier signal is needed. Since picture information is combined into a composite signal and then separated by the monitor, the resulting picture quality is still not the best attainable.

Color CRTs in graphics systems are designed as RGB monitors. These monitors use shadow-mask methods and take the intensity level for each electron gun (red, green, and blue) directly from the computer system without any intermediate processing. High-quality raster-graphics systems have 24 bits per pixel in the frame buffer, allowing 256 voltage settings for each electron gun and nearly 17 million color choices for each pixel. An RGB color system with 24 bits of storage per pixel is generally referred to as a full-color system or a true-color system.

Flat-Panel Displays

Although most graphics monitors are still constructed with CRTs, other technologies are emerging that may soon replace CRT monitors. The term flat-panel display refers to a class of video devices that have reduced volume, weight, and power requirements compared to a CRT. A significant feature of flat-panel displays is that they are thinner than CRTs, and we can hang them on walls or wear them on our wrists. Since we can even write on some flat-panel displays, they are also available as pocket notepads. Some additional uses for flat-panel displays are as small TV monitors, calculator screens, pocket video-game screens, laptop computer screens, armrest movie-viewing stations on airlines, advertisement boards in elevators, and graphics displays in applications requiring rugged, portable monitors.

We can separate flat-panel displays into two categories: emissive displays and nonemissive displays. The emissive displays (or emitters) are devices that convert electrical energy into light. Plasma panels, thin-film electroluminescent displays, and light-emitting diodes are examples of emissive displays. Flat CRTs have also been devised, in which electron beams are accelerated parallel to the screen and then deflected 90° onto the screen. But flat CRTs have not proved to be as successful as other emissive devices. Nonemissive displays (or nonemitters) use optical effects to convert sunlight or light from some other source into graphics patterns. The most important example of a nonemissive flat-panel display is a liquid-crystal device.

Plasma panels, also called gas-discharge displays, are constructed by filling the region between two glass plates with a mixture of gases that usually includes neon. A series of vertical conducting ribbons is placed on one glass panel, and a set of horizontal conducting ribbons is built into the other glass panel (Fig. 10). Firing voltages applied to an intersecting pair of horizontal and vertical conductors cause the gas at the intersection of the two conductors to break down into a glowing plasma of electrons and ions. Picture definition is stored in a refresh buffer, and the firing voltages are applied to refresh the pixel positions (at the intersections of the conductors) 60 times per second. Alternating-current methods are used to provide faster application of the firing voltages and, thus, brighter displays. Separation between pixels is provided by the electric field of the conductors. One disadvantage of plasma panels has been that they were strictly monochromatic devices, but systems are now available with multicolor capabilities.

Thin-film electroluminescent displays are similar in construction to plasma panels. The difference is that the region between the glass plates is filled with a phosphor, such as zinc sulfide doped with manganese, instead of a gas (Fig. 11). When a sufficiently high voltage is applied to a pair of crossing electrodes, the
phosphor becomes a conductor in the area of the intersection of the two electrodes. Electrical energy is absorbed by the manganese atoms, which then release the energy as a spot of light similar to the glowing plasma effect in a plasma panel. Electroluminescent displays require more power than plasma panels, and good color displays are harder to achieve.

A third type of emissive device is the light-emitting diode (LED). A matrix of diodes is arranged to form the pixel positions in the display, and picture definition is stored in a refresh buffer. As in scan-line refreshing of a CRT, information is read from the refresh buffer and converted to voltage levels that are applied to the diodes to produce the light patterns in the display.

Liquid-crystal displays (LCDs) are commonly used in small systems, such as laptop computers and calculators (Fig. 12). These nonemissive devices produce a picture by passing polarized light from the surroundings or from an internal light source through a liquid-crystal material that can be aligned to either block or transmit the light.

The term liquid crystal refers to the fact that these compounds have a crystalline arrangement of molecules, yet they flow like a liquid. Flat-panel displays commonly use nematic (threadlike) liquid-crystal compounds that tend to keep the long axes of the rod-shaped molecules aligned. A flat-panel display can then be constructed with a nematic liquid crystal, as demonstrated in Figure 13. Two glass plates, each containing a light polarizer that is aligned at a right angle to the other plate, sandwich the liquid-crystal material. Rows of horizontal, transparent conductors are built into one glass plate, and columns of vertical conductors are put into the other plate. The intersection of two conductors defines a pixel position. Normally, the molecules are aligned as shown in the “on state” of Figure 13. Polarized light passing through the material is twisted so that it will pass through the opposite polarizer. The light is then reflected back to the viewer. To turn off the pixel, we apply a voltage to the two intersecting conductors to align the molecules so that the light is not twisted. This type of flat-panel device is referred to as a passive-matrix LCD. Picture definitions are stored in a refresh buffer, and the screen is refreshed at the rate of 60 frames per second, as in the emissive
FIGURE 13
The light-twisting, shutter effect used in the design of most LCD devices.

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devices. Backlighting is also commonly applied using solid-state electronic devices, so that the system is not completely dependent on outside light sources. Colors can be displayed by using different materials or dyes and by placing a triad of color pixels at each screen location. Another method for constructing LCDs is to place a transistor at each pixel location, using thin-film transistor technology. The transistors are used to control the voltage at pixel locations and to prevent charge from gradually leaking out of the liquid-crystal cells. These devices are called active-matrix displays.

Three-Dimensional Viewing Devices

Graphics monitors for the display of three-dimensional scenes have been devised using a technique that reflects a CRT image from a vibrating, flexible mirror (Fig. 14). As the varifocal mirror vibrates, it changes focal length. These vibrations are synchronized with the display of an object on a CRT so that each point on the object is reflected from the mirror into a spatial position corresponding to the distance of that point from a specified viewing location. This allows us to walk around an object or scene and view it from different sides.

In addition to displaying three-dimensional images, these systems are often capable of displaying two-dimensional cross-sectional “slices” of objects selected at different depths, such as in medical applications to analyze data from ultrasonography and CAT scan devices, in geological applications to analyze topological and seismic data, in design applications involving solid objects, and in three-dimensional simulations of systems, such as molecules and terrain.
Stereoscopic and Virtual-Reality Systems

Another technique for representing a three-dimensional object is to display stereoscopic views of the object. This method does not produce true three-dimensional images, but it does provide a three-dimensional effect by presenting a different view to each eye of an observer so that scenes do appear to have depth.

To obtain a stereoscopic projection, we must obtain two views of a scene generated with viewing directions along the lines from the position of each eye (left and right) to the scene. We can construct the two views as computer-generated scenes with different viewing positions, or we can use a stereo camera pair to photograph an object or scene. When we simultaneously look at the left view with the left eye and the right view with the right eye, the two views merge into a single image and we perceive a scene with depth.

One way to produce a stereoscopic effect on a raster system is to display each of the two views on alternate refresh cycles. The screen is viewed through glasses, with each lens designed to act as a rapidly alternating shutter that is synchronized to block out one of the views. One such design (Figure 15) uses liquid-crystal shutters and an infrared emitter that synchronizes the glasses with the views on the screen.

Stereoscopic viewing is also a component in virtual-reality systems, where users can step into a scene and interact with the environment. A headset containing an optical system to generate the stereoscopic views can be used in conjunction with interactive input devices to locate and manipulate objects in the scene. A sensing system in the headset keeps track of the viewer’s position, so that the front and back of objects can be seen as the viewer “walks through” and interacts with the display. Another method for creating a virtual-reality environment
is to use projectors to generate a scene within an arrangement of walls, where a viewer interacts with a virtual display using stereoscopic glasses and data gloves (Section 4).

Lower-cost, interactive virtual-reality environments can be set up using a graphics monitor, stereoscopic glasses, and a head-tracking device. The tracking device is placed above the video monitor and is used to record head movements, so that the viewing position for a scene can be changed as head position changes.

## 2 Raster-Scan Systems

Interactive raster-graphics systems typically employ several processing units. In addition to the central processing unit (CPU), a special-purpose processor, called the **video controller** or **display controller**, is used to control the operation of the display device. Organization of a simple raster system is shown in Figure 16. Here, the frame buffer can be anywhere in the system memory, and the video controller accesses the frame buffer to refresh the screen. In addition to the video controller, more sophisticated raster systems employ other processors as coprocessors and accelerators to implement various graphics operations.

### Video Controller

Figure 17 shows a commonly used organization for raster systems. A fixed area of the system memory is reserved for the frame buffer, and the video controller is given direct access to the frame-buffer memory.

Frame-buffer locations, and the corresponding screen positions, are referenced in Cartesian coordinates. In an application program, we use the commands
within a graphics software package to set coordinate positions for displayed objects relative to the origin of the Cartesian reference frame. Often, the coordinate origin is referenced at the lower-left corner of a screen display area by the software commands, although we can typically set the origin at any convenient location for a particular application. Figure 18 shows a two-dimensional Cartesian reference frame with the origin at the lower-left screen corner. The screen surface is then represented as the first quadrant of a two-dimensional system, with positive $x$ values increasing from left to right and positive $y$ values increasing from the bottom of the screen to the top. Pixel positions are then assigned integer $x$ values that range from 0 to $x_{\text{max}}$ across the screen, left to right, and integer $y$ values that vary from 0 to $y_{\text{max}}$, bottom to top. However, hardware processes such as screen refreshing, as well as some software systems, reference the pixel positions from the top-left corner of the screen.

In Figure 19, the basic refresh operations of the video controller are diagrammed. Two registers are used to store the coordinate values for the screen pixels. Initially, the $x$ register is set to 0 and the $y$ register is set to the value for the top scan line. The contents of the frame buffer at this pixel position are then retrieved and used to set the intensity of the CRT beam. Then the $x$ register is incremented by 1, and the process is repeated for the next pixel on the top scan line. This procedure continues for each pixel along the top scan line. After the last pixel on the top scan line has been processed, the $x$ register is reset to 0 and the $y$ register is set to the value for the next scan line down from the top of the screen. Pixels along this scan line are then processed in turn, and the procedure is repeated for each successive scan line. After cycling through all pixels along the bottom scan line, the video controller resets the registers to the first pixel position on the top scan line and the refresh process starts over.

Since the screen must be refreshed at a rate of at least 60 frames per second, the simple procedure illustrated in Figure 19 may not be accommodated by typical RAM chips if the cycle time is too slow. To speed up pixel processing,
video controllers can retrieve multiple pixel values from the refresh buffer on each pass. The multiple pixel intensities are then stored in a separate register and used to control the CRT beam intensity for a group of adjacent pixels. When that group of pixels has been processed, the next block of pixel values is retrieved from the frame buffer.

A video controller can be designed to perform a number of other operations. For various applications, the video controller can retrieve pixel values from different memory areas on different refresh cycles. In some systems, for example, multiple frame buffers are often provided so that one buffer can be used for refreshing while pixel values are being loaded into the other buffers. Then the current refresh buffer can switch roles with one of the other buffers. This provides a fast mechanism for generating real-time animations, for example, since different views of moving objects can be successively loaded into a buffer without interrupting a refresh cycle. Another video-controller task is the transformation of blocks of pixels, so that screen areas can be enlarged, reduced, or moved from one location to another during the refresh cycles. In addition, the video controller often contains a lookup table, so that pixel values in the frame buffer are used to access the lookup table instead of controlling the CRT beam intensity directly. This provides a fast method for changing screen intensity values. Finally, some systems are designed to allow the video controller to mix the frame-buffer image with an input image from a television camera or other input device.

**Raster-Scan Display Processor**

Figure 20 shows one way to organize the components of a raster system that contains a separate display processor, sometimes referred to as a graphics controller or a display coprocessor. The purpose of the display processor is to free the CPU from the graphics chores. In addition to the system memory, a separate display-processor memory area can be provided.

A major task of the display processor is digitizing a picture definition given in an application program into a set of pixel values for storage in the frame buffer. This digitization process is called scan conversion. Graphics commands specifying straight lines and other geometric objects are scan converted into a set of discrete points, corresponding to screen pixel positions. Scan converting a straight-line segment, for example, means that we have to locate the pixel positions closest to the line path and store the color for each position in the frame buffer.
buffer. Similar methods are used for scan converting other objects in a picture definition. Characters can be defined with rectangular pixel grids, as in Figure 21, or they can be defined with outline shapes, as in Figure 22. The array size for character grids can vary from about 5 by 7 to 9 by 12 or more for higher-quality displays. A character grid is displayed by superimposing the rectangular grid pattern into the frame buffer at a specified coordinate position. For characters that are defined as outlines, the shapes are scan-converted into the frame buffer by locating the pixel positions closest to the outline.

Display processors are also designed to perform a number of additional operations. These functions include generating various line styles (dashed, dotted, or solid), displaying color areas, and applying transformations to the objects in a scene. Also, display processors are typically designed to interface with interactive input devices, such as a mouse.

In an effort to reduce memory requirements in raster systems, methods have been devised for organizing the frame buffer as a linked list and encoding the color information. One organization scheme is to store each scan line as a set of number pairs. The first number in each pair can be a reference to a color value, and the second number can specify the number of adjacent pixels on the scan line that are to be displayed in that color. This technique, called run-length encoding, can result in a considerable saving in storage space if a picture is to be constructed Mostly with long runs of a single color each. A similar approach can be taken when pixel colors change linearly. Another approach is to encode the raster as a set of rectangular areas (cell encoding). The disadvantages of encoding runs are that color changes are difficult to record and storage requirements increase as the lengths of the runs decrease. In addition, it is difficult for the display controller to process the raster when many short runs are involved. Moreover, the size of the frame buffer is no longer a major concern, because of sharp declines in memory costs. Nevertheless, encoding methods can be useful in the digital storage and transmission of picture information.

3 Graphics Workstations and Viewing Systems

Most graphics monitors today operate as raster-scan displays, and both CRT and flat-panel systems are in common use. Graphics workstations range from small general-purpose computer systems to multi-monitor facilities, often with ultra-large viewing screens. For a personal computer, screen resolutions vary from about 640 by 480 to 1280 by 1024, and diagonal screen lengths measure from 12 inches to over 21 inches. Most general-purpose systems now have considerable color capabilities, and many are full-color systems. For a desktop workstation specifically designed for graphics applications, the screen resolution can vary from 1280 by 1024 to about 1600 by 1200, with a typical screen diagonal of 18 inches or more. Commercial workstations can also be obtained with a variety of devices for specific applications.

High-definition graphics systems, with resolutions up to 2560 by 2048, are commonly used in medical imaging, air-traffic control, simulation, and CAD. Many high-end graphics workstations also include large viewing screens, often with specialized features.

Multi-panel display screens are used in a variety of applications that require "wall-sized" viewing areas. These systems are designed for presenting graphics displays at meetings, conferences, conventions, trade shows, retail stores, museums, and passenger terminals. A multi-panel display can be used to show a large
view of a single scene or several individual images. Each panel in the system displays one section of the overall picture. Color Plate 7 shows a 360° paneled viewing system in the NASA control-tower simulator, which is used for training and for testing ways to solve air-traffic and runway problems at airports. Large graphics displays can also be presented on curved viewing screens. A large, curved-screen system can be useful for viewing by a group of people studying a particular graphics application, such as the example in Color Plate 8. A control center, featuring a battery of standard monitors, allows an operator to view sections of the large display and to control the audio, video, lighting, and projection systems using a touch-screen menu. The system projectors provide a seamless, multichannel display that includes edge blending, distortion correction, and color balancing. And a surround-sound system is used to provide the audio environment.

4 Input Devices

Graphics workstations can make use of various devices for data input. Most systems have a keyboard and one or more additional devices specifically designed for interactive input. These include a mouse, trackball, spaceball, and joystick. Some other input devices used in particular applications are digitizers, dials, button boxes, data gloves, touch panels, image scanners, and voice systems.

Keyboards, Button Boxes, and Dials

An alphanumeric keyboard on a graphics system is used primarily as a device for entering text strings, issuing certain commands, and selecting menu options. The keyboard is an efficient device for inputting such nongraphic data as picture labels associated with a graphics display. Keyboards can also be provided with features to facilitate entry of screen coordinates, menu selections, or graphics functions.

Cursor-control keys and function keys are common features on general-purpose keyboards. Function keys allow users to select frequently accessed operations with a single keystroke, and cursor-control keys are convenient for selecting a displayed object or a location by positioning the screen cursor. A keyboard can also contain other types of cursor-positioning devices, such as a trackball or joystick, along with a numeric keypad for fast entry of numeric data. In addition to these features, some keyboards have an ergonomic design that provides adjustments for relieving operator fatigue.

For specialized tasks, input to a graphics application may come from a set of buttons, dials, or switches that select data values or customized graphics operations. Buttons and switches are often used to input predefined functions, and dials are common devices for entering scalar values. Numerical values within some defined range are selected for input with dial rotations. A potentiometer is used to measure dial rotation, which is then converted to the corresponding numerical value.

Mouse Devices

A mouse is a small handheld unit that is usually moved around on a flat surface to position the screen cursor. One or more buttons on the top of the mouse provide a mechanism for communicating selection information to the computer; wheels or rollers on the bottom of the mouse can be used to record the amount and direction of movement. Another method for detecting mouse motion is with an optical sensor. For some optical systems, the mouse is moved over a special mouse pad that has a grid of horizontal and vertical lines. The optical sensor detects
movement across the lines in the grid. Other optical mouse systems can operate on any surface. Some mouse systems are cordless, communicating with computer processors using digital radio technology.

Since a mouse can be picked up and put down at another position without change in cursor movement, it is used for making relative changes in the position of the screen cursor. One, two, three, or four buttons are included on the top of the mouse for signaling the execution of operations, such as recording cursor position or invoking a function. Most general-purpose graphics systems now include a mouse and a keyboard as the primary input devices.

Additional features can be included in the basic mouse design to increase the number of allowable input parameters and the functionality of the mouse. The Logitech G700 wireless gaming mouse in Figure 23 features 13 separately-programmable control inputs. Each input can be configured to perform a wide range of actions, from traditional single-click inputs to macro operations containing multiple keystrokes, mouse events, and pre-programmed delays between operations. The laser-based optical sensor can be configured to control the degree of sensitivity to motion, allowing the mouse to be used in situations requiring different levels of control over cursor movement. In addition, the mouse can hold up to five different configuration profiles to allow the configuration to be switched easily when changing applications.

**Trackballs and Spaceballs**

A *trackball* is a ball device that can be rotated with the fingers or palm of the hand to produce screen-cursor movement. Potentiometers, connected to the ball, measure the amount and direction of rotation. Laptop keyboards are often equipped with a trackball to eliminate the extra space required by a mouse. A trackball also can be mounted on other devices, or it can be obtained as a separate add-on unit that contains two or three control buttons.

An extension of the two-dimensional trackball concept is the *spaceball*, which provides six degrees of freedom. Unlike the trackball, a spaceball does not actually move. Strain gauges measure the amount of pressure applied to the spaceball to provide input for spatial positioning and orientation as the ball is pushed or pulled in various directions. Spaceballs are used for three-dimensional positioning and selection operations in virtual-reality systems, modeling, animation, CAD, and other applications.

**Joysticks**

Another positioning device is the *joystick*, which consists of a small, vertical lever (called the stick) mounted on a base. We use the joystick to steer the screen cursor around. Most joysticks select screen positions with actual stick movement; others
respond to pressure on the stick. Some joysticks are mounted on a keyboard, and some are designed as stand-alone units.

The distance that the stick is moved in any direction from its center position corresponds to the relative screen-cursor movement in that direction. Potentiometers mounted at the base of the joystick measure the amount of movement, and springs return the stick to the center position when it is released. One or more buttons can be programmed to act as input switches to signal actions that are to be executed once a screen position has been selected.

In another type of movable joystick, the stick is used to activate switches that cause the screen cursor to move at a constant rate in the direction selected. Eight switches, arranged in a circle, are sometimes provided so that the stick can select any one of eight directions for cursor movement. Pressure-sensitive joysticks, also called isometric joysticks, have a non-movable stick. A push or pull on the stick is measured with strain gauges and converted to movement of the screen cursor in the direction of the applied pressure.

Data Gloves
A data glove is a device that fits over the user’s hand and can be used to grasp a “virtual object.” The glove is constructed with a series of sensors that detect hand and finger motions. Electromagnetic coupling between transmitting antennas and receiving antennas are used to provide information about the position and orientation of the hand. The transmitting and receiving antennas can each be structured as a set of three mutually perpendicular coils, forming a three-dimensional Cartesian reference system. Input from the glove is used to position or manipulate objects in a virtual scene. A two-dimensional projection of the scene can be viewed on a video monitor, or a three-dimensional projection can be viewed with a headset.

Digitizers
A common device for drawing, painting, or interactively selecting positions is a digitizer. These devices can be designed to input coordinate values in either a two-dimensional or a three-dimensional space. In engineering or architectural applications, a digitizer is often used to scan a drawing or object and to input a set of discrete coordinate positions. The input positions are then joined with straight-line segments to generate an approximation of a curve or surface shape.

One type of digitizer is the graphics tablet (also referred to as a data tablet), which is used to input two-dimensional coordinates by activating a hand cursor or stylus at selected positions on a flat surface. A hand cursor contains crosshairs for sighting positions, while a stylus is a pencil-shaped device that is pointed at positions on the tablet. The tablet size varies from 12 by 12 inches for desktop models to 44 by 60 inches or larger for floor models. Graphics tablets provide a highly accurate method for selecting coordinate positions, with an accuracy that varies from about 0.2 mm on desktop models to about 0.05 mm or less on larger models.

Many graphics tablets are constructed with a rectangular grid of wires embedded in the tablet surface. Electromagnetic pulses are generated in sequence along the wires, and an electric signal is induced in a wire coil in an activated stylus or hand-cursor to record a tablet position. Depending on the technology, signal strength, coded pulses, or phase shifts can be used to determine the position on the tablet.

An acoustic (or sonic) tablet uses sound waves to detect a stylus position. Either strip microphones or point microphones can be employed to detect the sound
emitted by an electrical spark from a stylus tip. The position of the stylus is calculated by timing the arrival of the generated sound at the different microphone positions. An advantage of two-dimensional acoustic tablets is that the microphones can be placed on any surface to form the “tablet” work area. For example, the microphones could be placed on a book page while a figure on that page is digitized.

Three-dimensional digitizers use sonic or electromagnetic transmissions to record positions. One electromagnetic transmission method is similar to that employed in the data glove: A coupling between the transmitter and receiver is used to compute the location of a stylus as it moves over an object surface. As the points are selected on a nonmetallic object, a wire-frame outline of the surface is displayed on the computer screen. Once the surface outline is constructed, it can be rendered using lighting effects to produce a realistic display of the object.

Image Scanners

Drawings, graphs, photographs, or text can be stored for computer processing with an image scanner by passing an optical scanning mechanism over the information to be stored. The gradations of grayscale or color are then recorded and stored in an array. Once we have the internal representation of a picture, we can apply transformations to rotate, scale, or crop the picture to a particular screen area. We can also apply various image-processing methods to modify the array representation of the picture. For scanned text input, various editing operations can be performed on the stored documents. Scanners are available in a variety of sizes and capabilities, including small handheld models, drum scanners, and flatbed scanners.

Touch Panels

As the name implies, touch panels allow displayed objects or screen positions to be selected with the touch of a finger. A typical application of touch panels is for the selection of processing options that are represented as a menu of graphical icons. Some monitors are designed with touch screens. Other systems can be adapted for touch input by fitting a transparent device containing a touch-sensing mechanism over the video monitor screen. Touch input can be recorded using optical, electrical, or acoustical methods.

Optical touch panels employ a line of infrared light-emitting diodes (LEDs) along one vertical edge and along one horizontal edge of the frame. Light detectors are placed along the opposite vertical and horizontal edges. These detectors are used to record which beams are interrupted when the panel is touched. The two crossing beams that are interrupted identify the horizontal and vertical coordinates of the screen position selected. Positions can be selected with an accuracy of about 1/4 inch. With closely spaced LEDs, it is possible to break two horizontal or two vertical beams simultaneously. In this case, an average position between the two interrupted beams is recorded. The LEDs operate at infrared frequencies so that the light is not visible to a user.

An electrical touch panel is constructed with two transparent plates separated by a small distance. One of the plates is coated with a conducting material, and the other plate is coated with a resistive material. When the outer plate is touched, it is forced into contact with the inner plate. This contact creates a voltage drop across the resistive plate that is converted to the coordinate values of the selected screen position.

In acoustical touch panels, high-frequency sound waves are generated in horizontal and vertical directions across a glass plate. Touching the screen causes
part of each wave to be reflected from the finger to the emitters. The screen position at the point of contact is calculated from a measurement of the time interval between the transmission of each wave and its reflection to the emitter.

**Light Pens**

Light pens are pencil-shaped devices used to select screen positions by detecting the light coming from points on the CRT screen. They are sensitive to the short burst of light emitted from the phosphor coating at the instant the electron beam strikes a particular point. Other light sources, such as the background light in the room, are usually not detected by a light pen. An activated light pen, pointed at a spot on the screen as the electron beam lights up that spot, generates an electrical pulse that causes the coordinate position of the electron beam to be recorded. As with cursor-positioning devices, recorded light-pen coordinates can be used to position an object or to select a processing option.

Although light pens are still with us, they are not as popular as they once were because they have several disadvantages compared to other input devices that have been developed. For example, when a light pen is pointed at the screen, part of the screen image is obscured by the hand and pen. In addition, prolonged use of the light pen can cause arm fatigue, and light pens require special implementations for some applications because they cannot detect positions within black areas. To be able to select positions in any screen area with a light pen, we must have some nonzero light intensity emitted from each pixel within that area. In addition, light pens sometimes give false readings due to background lighting in a room.

**Voice Systems**

Speech recognizers are used with some graphics workstations as input devices for voice commands. The voice system input can be used to initiate graphics operations or to enter data. These systems operate by matching an input against a predefined dictionary of words and phrases.

A dictionary is set up by speaking the command words several times. The system then analyzes each word and establishes a dictionary of word frequency patterns, along with the corresponding functions that are to be performed. Later, when a voice command is given, the system searches the dictionary for a frequency-pattern match. A separate dictionary is needed for each operator using the system. Input for a voice system is typically spoken into a microphone mounted on a headset; the microphone is designed to minimize input of background sounds. Voice systems have some advantage over other input devices because the attention of the operator need not switch from one device to another to enter a command.

**5 Hard-Copy Devices**

We can obtain hard-copy output for our images in several formats. For presentations or archiving, we can send image files to devices or service bureaus that will produce overhead transparencies, 35mm slides, or film. Also, we can put our pictures on paper by directing graphics output to a printer or plotter.

The quality of the pictures obtained from an output device depends on dot size and the number of dots per inch, or lines per inch, that can be displayed. To produce smooth patterns, higher-quality printers shift dot positions so that adjacent dots overlap.
Printers produce output by either impact or nonimpact methods. **Impact** printers press formed character faces against an inked ribbon onto the paper. A line printer is an example of an impact device, with the typefaces mounted on bands, chains, drums, or wheels. **Nonimpact** printers and plotters use laser techniques, ink-jet sprays, electrostatic methods, and electrothermal methods to get images onto paper.

Character impact printers often have a **dot-matrix** print head containing a rectangular array of protruding wire pins, with the number of pins varying depending upon the quality of the printer. Individual characters or graphics patterns are obtained by retracting certain pins so that the remaining pins form the pattern to be printed. Figure 24 shows a picture printed on a dot-matrix printer.

In a **laser** device, a laser beam creates a charge distribution on a rotating drum coated with a photoelectric material, such as selenium. Toner is applied to the drum and then transferred to paper. **Ink-jet** methods produce output by squirting ink in horizontal rows across a roll of paper wrapped on a drum. The electrically charged ink stream is deflected by an electric field to produce dot-matrix patterns. An **electrostatic** device places a negative charge on the paper, one complete row at a time across the sheet. Then the paper is exposed to a positively charged toner. This causes the toner to be attracted to the negatively charged areas, where it adheres to produce the specified output. Another output technology is the **electrothermal** printer. With these systems, heat is applied to a dot-matrix print head to output patterns on heat-sensitive paper.

We can get limited color output on some impact printers by using different-colored ribbons. Nonimpact devices use various techniques to combine three different color pigments (cyan, magenta, and yellow) to produce a range of color patterns. Laser and electrostatic devices deposit the three pigments on separate passes; ink-jet methods shoot the three colors simultaneously on a single pass along each print line.

Drafting layouts and other drawings are typically generated with ink-jet or pen plotters. A pen plotter has one or more pens mounted on a carriage, or crossbar, that spans a sheet of paper. Pens with varying colors and widths are used to produce a variety of shadings and line styles. Wet-ink, ballpoint, and felt-tip pens are all possible choices for use with a pen plotter. Plotter paper can lie flat or it can be rolled onto a drum or belt. Crossbars can be either movable or stationary, while the pen moves back and forth along the bar. The paper is held in position using clamps, a vacuum, or an electrostatic charge.
6 Graphics Networks

So far, we have mainly considered graphics applications on an isolated system with a single user. However, multiuser environments and computer networks are now common elements in many graphics applications. Various resources, such as processors, printers, plotters, and data files, can be distributed on a network and shared by multiple users.

A graphics monitor on a network is generally referred to as a **graphics server**, or simply a **server**. Often, the monitor includes standard input devices such as a keyboard and a mouse or trackball. In that case, the system can provide input, as well as being an output server. The computer on the network that is executing a graphics application program is called the **client**, and the output of the program is displayed on a server. A workstation that includes processors, as well as a monitor and input devices, can function as both a server and a client.

When operating on a network, a client computer transmits the instructions for displaying a picture to the monitor (server). Typically, this is accomplished by collecting the instructions into packets before transmission instead of sending the individual graphics instructions one at a time over the network. Thus, graphics software packages often contain commands that affect packet transmission, as well as the commands for creating pictures.

7 Graphics on the Internet

A great deal of graphics development is now done on the global collection of computer networks known as the **Internet**. Computers on the Internet communicate using **transmission control protocol/internet protocol** (TCP/IP). In addition, the **World Wide Web** provides a hypertext system that allows users to locate and view documents that can contain text, graphics, and audio. Resources, such as graphics files, are identified by a **uniform (or universal) resource locator** (URL). Each URL contains two parts: (1) the protocol for transferring the document, and (2) the server that contains the document and, optionally, the location (directory) on the server. For example, the URL `http://www.siggraph.org/` indicates a document that is to be transferred with the hypertext transfer protocol (http) and that the server is www.siggraph.org, which is the home page of the Special Interest Group in Graphics (SIGGRAPH) of the Association for Computing Machinery. Another common type of URL begins with `ftp://`. This identifies a site that accepts file transfer protocol (FTP) connections, through which programs or other files can be downloaded.

Documents on the Internet can be constructed with the **Hypertext Markup Language** (HTML). The development of HTML provided a simple method for describing a document containing text, graphics, and references (hyperlinks) to other documents. Although resources could be made available using HTML and URL addressing, it was difficult originally to find information on the Internet. Subsequently, the National Center for Supercomputing Applications (NCSA) developed a “browser” called Mosaic, which made it easier for users to search for Web resources. The Mosaic browser later evolved into the browser called Netscape Navigator. In turn, Netscape Navigator inspired the creation of the Mozilla family of browsers, whose most well-known member is, perhaps, Firefox.

HTML provides a simple method for developing graphics on the Internet, but it has limited capabilities. Therefore, other languages have been developed for Internet graphics applications.
Summary

In this chapter, we surveyed the major hardware and software features of computer-graphics systems. Hardware components include video monitors, hardcopy output devices, various kinds of input devices, and components for interacting with virtual environments.

The predominant graphics display device is the raster refresh monitor, based on television technology. A raster system uses a frame buffer to store the color value for each screen position (pixel). Pictures are then painted onto the screen by retrieving this information from the frame buffer (also called a refresh buffer) as the electron beam in the CRT sweeps across each scan line from top to bottom. Older vector displays construct pictures by drawing straight-line segments between specified endpoint positions. Picture information is then stored as a set of line-drawing instructions.

Many other video display devices are available. In particular, flat-panel display technology is developing at a rapid rate, and these devices are now used in a variety of systems, including both desktop and laptop computers. Plasma panels and liquid-crystal devices are two examples of flat-panel displays. Other display technologies include three-dimensional and stereoscopic-viewing systems. Virtual-reality systems can include either a stereoscopic headset or a standard video monitor.

For graphical input, we have a range of devices to choose from. Keyboards, button boxes, and dials are used to input text, data values, or programming options. The most popular “pointing” device is the mouse, but trackballs, spaceballs, joysticks, cursor-control keys, and thumbwheels are also used to position the screen cursor. In virtual-reality environments, data gloves are commonly used. Other input devices are image scanners, digitizers, touch panels, light pens, and voice systems.

Hardcopy devices for graphics workstations include standard printers and plotters, in addition to devices for producing slides, transparencies, and film output. Printers produce hardcopy output using dot-matrix, laser, ink-jet, electrostatic, or electrothermal methods. Graphs and charts can be produced with an ink-pen plotter or with a combination printer-plotter device.

REFERENCES

A general treatment of electronic displays is available in Tannas (1985) and in Sherr (1993). Flat-panel devices are discussed in Depp and Howard (1993). Additional information on raster-graphics architecture can be found in Foley et al. (1990). Three-dimensional and stereoscopic displays are discussed in Johnson (1982) and in Grotch (1983). Head-mounted displays and virtual-reality environments are discussed in Chung et al. (1989).

EXERCISES

1. List the operating characteristics for the following display technologies: raster refresh systems, vector refresh systems, plasma panels, and LCDs.
2. List some applications appropriate for each of the display technologies in the previous question.
3. Determine the resolution (pixels per centimeter) in the x and y directions for the video monitor in use on your system. Determine the aspect ratio, and explain how relative proportions of objects can be maintained on your system.
4. Consider three different raster systems with resolutions of 800 by 600, 1280 by 960, and 1680 by 1050. What size frame buffer (in bytes) is needed for each of these systems to store 16 bits per pixel? How much storage is required for each system if 32 bits per pixel are to be stored?
5. Suppose an RGB raster system is to be designed using an 8 inch by 10 inch screen with a resolution of 100 pixels per inch in each direction. If we want to store 6 bits per pixel in the frame buffer, how much storage (in bytes) do we need for the frame buffer?
6. How long would it take to load an 800 by 600 frame buffer with 16 bits per pixel, if 105 bits can be transferred per second? How long would it take to
load a 32-bit-per-pixel frame buffer with a resolution of 1680 by 1050 using this same transfer rate?

7 Suppose we have a computer with 32 bits per word and a transfer rate of 1 mips (one million instructions per second). How long would it take to fill the frame buffer of a 300 dpi (dot per inch) laser printer with a page size of 8.5 inches by 11 inches?

8 Consider two raster systems with resolutions of 800 by 600 and 1680 by 1050. How many pixels could be accessed per second in each of these systems by a display controller that refreshes the screen at a rate of 60 frames per second? What is the access time per pixel in each system?

9 Suppose we have a video monitor with a display area that measures 12 inches across and 9.6 inches high. If the resolution is 1280 by 1024 and the aspect ratio is 1, what is the diameter of each screen point?

10 How much time is spent scanning across each row of pixels during screen refresh on a raster system with a resolution of 1680 by 1050 and a refresh rate of 30 frames per second?

11 Consider a noninterlaced raster monitor with a resolution of \( n \) by \( m \) (\( m \) scan lines and \( n \) pixels per scan line), a refresh rate of \( r \) frames per second, a horizontal retrace time of \( t_{\text{horiz}} \) and a vertical retrace time of \( t_{\text{vert}} \). What is the fraction of the total refresh time per frame spent in retrace of the electron beam?

12 What is the fraction of the total refresh time per frame spent in retrace of the electron beam for a noninterlaced raster system with a resolution of 1680 by 1050, a refresh rate of 65 Hz, a horizontal retrace time of 4 microseconds, and a vertical retrace time of 400 microseconds?

13 Assuming that a certain full-color (24 bits per pixel) RGB raster system has a 1024 by 1024 frame buffer, how many distinct color choices (intensity levels) would we have available? How many different colors could we display at any one time?

14 Compare the advantages and disadvantages of a three-dimensional monitor using a varifocal mirror to those of a stereoscopic system.

15 List the different input and output components that are typically used with virtual-reality systems. Also, explain how users interact with virtual scenes displayed with different output devices, such as two-dimensional and stereoscopic monitors.

16 Explain how virtual-reality systems can be used in design applications. What are some other applications for virtual-reality systems?

17 List some applications for large-screen displays.

18 Explain the differences between a general graphics system designed for a programmer and one designed for a specific application, such as architectural design.

**IN MORE DEPTH**

1 In this course, you will design and build a graphics application incrementally. You should have a basic understanding of the types of applications for which computer graphics are used. Try to formulate a few ideas about one or more particular applications you may be interested in developing over the course of your studies. Keep in mind that you will be asked to incorporate techniques covered in this text, as well as to show your understanding of alternative methods for implementing those concepts. As such, the application should be simple enough that you can realistically implement it in a reasonable amount of time, but complex enough to afford the inclusion of each of the relevant concepts in the text. One obvious example is a video game of some sort in which the user interacts with a virtual environment that is initially displayed in two dimensions and later in three dimensions. Some concepts to consider would be two- and three-dimensional objects of different forms (some simple, some moderately complex with curved surfaces, etc.), complex shading of object surfaces, various lighting techniques, and animation of some sort. Write a report with at least three to four ideas that you will choose to implement you acquire more knowledge of the course material. Note that one type of application may be more suited to demonstrate a given concept than another.

2 Find details about the graphical capabilities of the graphics controller and the display device in your system by looking up their specifications. Record the following information:

- What is the maximum resolution your graphics controller is capable of rendering?
- What is the maximum resolution of your display device?
- What type of hardware does your graphics controller contain?
- What is the GPU's clock speed?
- How much of its own graphics memory does it have?

If you have a relatively new system, it is unlikely that you will be pushing the envelope of your graphics hardware in your application development for this text. However, knowing the capabilities of your graphics system will provide you with a sense of how much it will be able to handle.
Computer Graphics Hardware
Color Plates

Color Plate 7
The 360° viewing screen in the NASA airport control-tower simulator, called the FutureFlight Central Facility. (Courtesy of Silicon Graphics, Inc. and NASA. © 2003 SGI. All rights reserved.)

Color Plate 8
A geophysical visualization presented on a 25-foot semicircular screen, which provides a 160° horizontal and 40° vertical field of view. (Courtesy of Silicon Graphics, Inc., the Landmark Graphics Corporation, and Trimension Systems. © 2003 SGI. All rights reserved.)

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There are two broad classifications for computer-graphics software: special-purpose packages and general programming packages. Special-purpose packages are designed for nonprogrammers who want to generate pictures, graphs, or charts in some application area without worrying about the graphics procedures that might be needed to produce such displays. The interface to a special-purpose package is typically a set of menus that allow users to communicate with the programs in their own terms. Examples of such applications include artists’ painting programs and various architectural, business, medical, and engineering CAD systems. By contrast, a general programming package provides a library of graphics functions that can be used in a programming language such as C, C++, Java, or Fortran. Basic functions in a typical graphics library include those for specifying picture components (straight lines, polygons, spheres, and other objects), setting color values, selecting views of a scene, and applying rotations or other transformations. Some examples of general graphics programming packages are...
GL (Graphics Library), OpenGL, VRML (Virtual-Reality Modeling Language), Java 2D, and Java 3D. A set of graphics functions is often called a computer-graphics application programming interface (CG API) because the library provides a software interface between a programming language (such as C++) and the hardware. So when we write an application program in C++, the graphics routines allow us to construct and display a picture on an output device.

1 Coordinate Representations

To generate a picture using a programming package, we first need to give the geometric descriptions of the objects that are to be displayed. These descriptions determine the locations and shapes of the objects. For example, a box is specified by the positions of its corners (vertices), and a sphere is defined by its center position and radius. With few exceptions, general graphics packages require geometric descriptions to be specified in a standard, right-handed, Cartesian-coordinate reference frame. If coordinate values for a picture are given in some other reference frame (spherical, hyperbolic, etc.), they must be converted to Cartesian coordinates before they can be input to the graphics package. Some packages that are designed for specialized applications may allow use of other coordinate frames that are appropriate for those applications.

In general, several different Cartesian reference frames are used in the process of constructing and displaying a scene. First, we can define the shapes of individual objects, such as trees or furniture, within a separate reference frame for each object. These reference frames are called modeling coordinates, or sometimes local coordinates or master coordinates. Once the individual object shapes have been specified, we can construct (“model”) a scene by placing the objects into appropriate locations within a scene reference frame called world coordinates. This step involves the transformation of the individual modeling-coordinate frames to specified positions and orientations within the world-coordinate frame. As an example, we could construct a bicycle by defining each of its parts (wheels, frame, seat, handlebars, gears, chain, pedals) in a separate modeling-coordinate frame. Then, the component parts are fitted together in world coordinates. If both bicycle wheels are the same size, we need to describe only one wheel in a local-coordinate frame. Then the wheel description is fitted into the world-coordinate bicycle description in two places. For scenes that are not too complicated, object components can be set up directly within the overall world-coordinate object structure, bypassing the modeling-coordinate and modeling-transformation steps. Geometric descriptions in modeling coordinates and world coordinates can be given in any convenient floating-point or integer values, without regard for the constraints of a particular output device. For some scenes, we might want to specify object geometries in fractions of a foot, while for other applications we might want to use millimeters, or kilometers, or light-years.

After all parts of a scene have been specified, the overall world-coordinate description is processed through various routines onto one or more output-device reference frames for display. This process is called the viewing pipeline. World-coordinate positions are first converted to viewing coordinates corresponding to the view we want of a scene, based on the position and orientation of a hypothetical camera. Then object locations are transformed to a two-dimensional (2D) projection of the scene, which corresponds to what we will see on the output device. The scene is then stored in normalized coordinates, where each coordinate value is in the range from −1 to 1 or in the range from 0 to 1, depending on the system.
FIGURE 1
The transformation sequence from modeling coordinates to device coordinates for a three-dimensional scene. Object shapes can be individually defined in modeling-coordinate reference systems. Then the shapes are positioned within the world-coordinate scene. Next, world-coordinate specifications are transformed through the viewing pipeline to viewing and projection coordinates and then to normalized coordinates. At the final step, individual device drivers transfer the normalized-coordinate representation of the scene to the output devices for display.

Normalized coordinates are also referred to as normalized device coordinates, since using this representation makes a graphics package independent of the coordinate range for any specific output device. We also need to identify visible surfaces and eliminate picture parts outside the bounds for the view we want to show on the display device. Finally, the picture is scan-converted into the refresh buffer of a raster system for display. The coordinate systems for display devices are generally called device coordinates, or screen coordinates in the case of a video monitor. Often, both normalized coordinates and screen coordinates are specified in a left-handed coordinate reference frame so that increasing positive distances from the \(xy\) plane (the screen, or viewing plane) can be interpreted as being farther from the viewing position.

Figure 1 briefly illustrates the sequence of coordinate transformations from modeling coordinates to device coordinates for a display that is to contain a view of two three-dimensional (3D) objects. An initial modeling-coordinate position \((x_{mc}, y_{mc}, z_{mc})\) in this illustration is transferred to world coordinates, then to viewing and projection coordinates, then to left-handed normalized coordinates, and finally to a device-coordinate position \((x_{dc}, y_{dc})\) with the sequence:

\[
(x_{mc}, y_{mc}, z_{mc}) \rightarrow (x_{wc}, y_{wc}, z_{wc}) \rightarrow (x_{vc}, y_{vc}, z_{vc}) \rightarrow (x_{pc}, y_{pc}, z_{pc}) \\
\rightarrow (x_{nc}, y_{nc}, z_{nc}) \rightarrow (x_{dc}, y_{dc})
\]

Device coordinates \((x_{dc}, y_{dc})\) are integers within the range \((0, 0)\) to \((x_{max}, y_{max})\) for a particular output device. In addition to the two-dimensional positions \((x_{dc}, y_{dc})\) on the viewing surface, depth information for each device-coordinate position is stored for use in various visibility and surface-processing algorithms.

2 Graphics Functions
A general-purpose graphics package provides users with a variety of functions for creating and manipulating pictures. These routines can be broadly classified
according to whether they deal with graphics output, input, attributes, transformations, viewing, subdividing pictures, or general control.

The basic building blocks for pictures are referred to as **graphics output primitives**. They include character strings and geometric entities, such as points, straight lines, curved lines, filled color areas (usually polygons), and shapes defined with arrays of color points. In addition, some graphics packages provide functions for displaying more complex shapes such as spheres, cones, and cylinders. Routines for generating output primitives provide the basic tools for constructing pictures.

**Attributes** are properties of the output primitives; that is, an attribute describes how a particular primitive is to be displayed. This includes color specifications, line styles, text styles, and area-filling patterns.

We can change the size, position, or orientation of an object within a scene using **geometric transformations**. Some graphics packages provide an additional set of functions for performing **modeling transformations**, which are used to construct a scene where individual object descriptions are given in local coordinates. Such packages usually provide a mechanism for describing complex objects (such as an electrical circuit or a bicycle) with a tree (hierarchical) structure. Other packages simply provide the geometric-transformation routines and leave modeling details to the programmer.

After a scene has been constructed, using the routines for specifying the object shapes and their attributes, a graphics package projects a view of the picture onto an output device. **Viewing transformations** are used to select a view of the scene, the type of projection to be used, and the location on a video monitor where the view is to be displayed. Other routines are available for managing the screen display area by specifying its position, size, and structure. For three-dimensional scenes, visible objects are identified and the lighting conditions are applied.

Interactive graphics applications use various kinds of input devices, including a mouse, a tablet, and a joystick. **Input functions** are used to control and process the data flow from these interactive devices.

Some graphics packages also provide routines for subdividing a picture description into a named set of component parts. And other routines may be available for manipulating these picture components in various ways.

Finally, a graphics package contains a number of housekeeping tasks, such as clearing a screen display area to a selected color and initializing parameters. We can lump the functions for carrying out these chores under the heading **control operations**.

### 3 Software Standards

The primary goal of standardized graphics software is portability. When packages are designed with standard graphics functions, software can be moved easily from one hardware system to another and used in different implementations and applications. Without standards, programs designed for one hardware system often cannot be transferred to another system without extensive rewriting of the programs.

International and national standards-planning organizations in many countries have cooperated in an effort to develop a generally accepted standard for computer graphics. After considerable effort, this work on standards led to the development of the **Graphical Kernel System (GKS)** in 1984. This system was adopted as the first graphics software standard by the International Standards Organization (ISO) and by various national standards organizations, including
the American National Standards Institute (ANSI). Although GKS was originally designed as a two-dimensional graphics package, a three-dimensional GKS extension was soon developed. The second software standard to be developed and approved by the standards organizations was Programmer’s Hierarchical Interactive Graphics System (PHIGS), which is an extension of GKS. Increased capabilities for hierarchical object modeling, color specifications, surface rendering, and picture manipulations are provided in PHIGS. Subsequently, an extension of PHIGS, called PHIGS+, was developed to provide three-dimensional surface-rendering capabilities not available in PHIGS.

As the GKS and PHIGS packages were being developed, the graphics workstations from Silicon Graphics, Inc. (SGI), became increasingly popular. These workstations came with a set of routines called GL (Graphics Library), which very soon became a widely used package in the graphics community. Thus, GL became a de facto graphics standard. The GL routines were designed for fast, real-time rendering, and soon this package was being extended to other hardware systems. As a result, OpenGL was developed as a hardware-independent version of GL in the early 1990s. This graphics package is now maintained and updated by the OpenGL Architecture Review Board, which is a consortium of representatives from many graphics companies and organizations. The OpenGL library is specifically designed for efficient processing of three-dimensional applications, but it can also handle two-dimensional scene descriptions as a special case of three dimensions where all the $z$ coordinate values are 0.

Graphics functions in any package are typically defined as a set of specifications independent of any programming language. A language binding is then defined for a particular high-level programming language. This binding gives the syntax for accessing the various graphics functions from that language. Each language binding is defined to make best use of the corresponding language capabilities and to handle various syntax issues, such as data types, parameter passing, and errors. Specifications for implementing a graphics package in a particular language are set by the ISO. The OpenGL bindings for the C and C++ languages are the same. Other OpenGL bindings are also available, such as those for Java and Python.

Later in this book, we use the C/C++ binding for OpenGL as a framework for discussing basic graphics concepts and the design and application of graphics packages. Example programs in C++ illustrate applications of OpenGL and the general algorithms for implementing graphics functions.

4 Other Graphics Packages

Many other computer-graphics programming libraries have been developed. Some provide general graphics routines, and some are aimed at specific applications or particular aspects of computer graphics, such as animation, virtual reality, or graphics on the Internet.

A package called Open Inventor furnishes a set of object-oriented routines for describing a scene that is to be displayed with calls to OpenGL. The Virtual-Reality Modeling Language (VRML), which began as a subset of Open Inventor, allows us to set up three-dimensional models of virtual worlds on the Internet. We can also construct pictures on the Web using graphics libraries developed for the Java language. With Java 2D, we can create two-dimensional scenes within Java applets, for example; or we can produce three-dimensional web displays with Java 3D. With the RenderMan Interface from the Pixar Corporation, we can generate scenes
using a variety of lighting models. Finally, graphics libraries are often provided in other types of systems, such as Mathematica, MatLab, and Maple.

5 Introduction to OpenGL

A basic library of functions is provided in OpenGL for specifying graphics primitives, attributes, geometric transformations, viewing transformations, and many other operations. As we noted in the last section, OpenGL is designed to be hardware independent, so many operations, such as input and output routines, are not included in the basic library. However, input and output routines and many additional functions are available in auxiliary libraries that have been developed for OpenGL programs.

Basic OpenGL Syntax

Function names in the OpenGL basic library (also called the OpenGL core library) are prefixed with gl, and each component word within a function name has its first letter capitalized. The following examples illustrate this naming convention:

    glBegin. glClear. glCopyPixels. glPolygonMode

Certain functions require that one (or more) of their arguments be assigned a symbolic constant specifying, for instance, a parameter name, a value for a parameter, or a particular mode. All such constants begin with the uppercase letters GL. In addition, component words within a constant name are written in capital letters, and the underscore (_) is used as a separator between all component words in the name. The following are a few examples of the several hundred symbolic constants available for use with OpenGL functions:

GL_2D, GL_RGB, GL_CCW, GL_POLYGON, GL_AMBIENT_AND_DIFFUSE

The OpenGL functions also expect specific data types. For example, an OpenGL function parameter might expect a value that is specified as a 32-bit integer. But the size of an integer specification can be different on different machines. To indicate a specific data type, OpenGL uses special built-in, data-type names, such as

    GLbyte, GLshort, GLint, GLfloat, GLdouble, GLboolean

Each data-type name begins with the capital letters GL, and the remainder of the name is a standard data-type designation written in lowercase letters.

Some arguments of OpenGL functions can be assigned values using an array that lists a set of data values. This is an option for specifying a list of values as a pointer to an array, rather than specifying each element of the list explicitly as a parameter argument. A typical example of the use of this option is in specifying xyz coordinate values.

Related Libraries

In addition to the OpenGL basic (core) library, there are a number of associated libraries for handling special operations. The OpenGL Utility (GLU) provides routines for setting up viewing and projection matrices, describing complex objects with line and polygon approximations, displaying quadrics and B-splines.
using linear approximations, processing the surface-rendering operations, and other complex tasks. Every OpenGL implementation includes the GLU library, and all GLU function names start with the prefix glu. There is also an object-oriented toolkit based on OpenGL, called Open Inventor, which provides routines and predefined object shapes for interactive three-dimensional applications. This toolkit is written in C++.

To create a graphics display using OpenGL, we first need to set up a display window on our video screen. This is simply the rectangular area of the screen in which our picture will be displayed. We cannot create the display window directly with the basic OpenGL functions since this library contains only device-independent graphics functions, and window-management operations depend on the computer we are using. However, there are several window-system libraries that support OpenGL functions for a variety of machines. The OpenGL Extension to the X Window System (GLX) provides a set of routines that are prefixed with the letters glX. Apple systems can use the Apple GL (AGL) interface for window-management operations. Function names for this library are prefixed with agl. For Microsoft Windows systems, the WGL routines provide a Windows-to-OpenGL interface. These routines are prefixed with the letters wg1. The Presentation Manager to OpenGL (PGL) is an interface for the IBM OS/2, which uses the prefix pg1 for the library routines. The OpenGL Utility Toolkit (GLUT) provides a library of functions for interacting with any screen-windowing system. The GLUT library functions are prefixed with glut, and this library also contains methods for describing and rendering quadric curves and surfaces.

Since GLUT is an interface to other device-specific window systems, we can use it so that our programs will be device-independent. Information regarding the latest version of GLUT and download procedures for the source code are available at the following web site:

http://www.opengl.org/resources/libraries/glut/

Header Files

In all of our graphics programs, we will need to include the header file for the OpenGL core library. For most applications we will also need GLU, and on many systems we will need to include the header file for the window system. For instance, with Microsoft Windows, the header file that accesses the WGL routines is windows.h. This header file must be listed before the OpenGL and GLU header files because it contains macros needed by the Microsoft Windows version of the OpenGL libraries. So the source file in this case would begin with

```c
#include <windows.h>
#include <GL/gl.h>
#include <GL/glu.h>
```

However, if we use GLUT to handle the window-managing operations, we do not need to include gl.h and glu.h because GLUT ensures that these will be included correctly. Thus, we can replace the header files for OpenGL and GLU with

```c
#include <GL/glut.h>
```

(We could include gl.h and glu.h as well, but doing so would be redundant and could affect program portability.) On some systems, the header files for OpenGL and GLUT routines are found in different places in the filesystem. For instance, on Apple OS X systems, the header file inclusion statement would be

```c
#include <GLUT/glut.h>
```
In addition, we will often need to include header files that are required by the C++ code. For example,

```c
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
```

With the ISO/ANSI standard for C++, these header files are called `cstdio`, `cstdlib`, and `cmath`.

Display-Window Management Using GLUT

To get started, we can consider a simplified, minimal number of operations for displaying a picture. Since we are using the OpenGL Utility Toolkit, our first step is to initialize GLUT. This initialization function could also process any command-line arguments, but we will not need to use these parameters for our first example programs. We perform the GLUT initialization with the statement

```c
glutInit (&argc, argv);
```

Next, we can state that a display window is to be created on the screen with a given caption for the title bar. This is accomplished with the function

```c
glutCreateWindow ("An Example OpenGL Program");
```

where the single argument for this function can be any character string that we want to use for the display-window title.

Then we need to specify what the display window is to contain. For this, we create a picture using OpenGL functions and pass the picture definition to the GLUT routine `glutDisplayFunc`, which assigns our picture to the display window. As an example, suppose we have the OpenGL code for describing a line segment in a procedure called `lineSegment`. Then the following function call passes the line-segment description to the display window:

```c
glutDisplayFunc (lineSegment);
```

But the display window is not yet on the screen. We need one more GLUT function to complete the window-processing operations. After execution of the following statement, all display windows that we have created, including their graphic content, are now activated:

```c
glutMainLoop ( );
```

This function must be the last one in our program. It displays the initial graphics and puts the program into an infinite loop that checks for input from devices such as a mouse or keyboard. Our first example will not be interactive, so the program will just continue to display our picture until we close the display window. In later chapters, we consider how we can modify our OpenGL programs to handle interactive input.

Although the display window that we created will be in some default location and size, we can set these parameters using additional GLUT functions. We use the `glutInitWindowPosition` function to give an initial location for the upper-left corner of the display window. This position is specified in integer screen
coordinates, whose origin is at the upper-left corner of the screen. For instance, the following statement specifies that the upper-left corner of the display window should be placed 50 pixels to the right of the left edge of the screen and 100 pixels down from the top edge of the screen:

    glutInitWindowPosition (50, 100);

Similarly, the glutInitWindowSize function is used to set the initial pixel width and height of the display window. Thus, we specify a display window with an initial width of 400 pixels and a height of 300 pixels (Fig. 2) with the statement

    glutInitWindowSize (400, 300);

After the display window is on the screen, we can reposition and resize it.

We can also set a number of other options for the display window, such as buffering and a choice of color modes, with the glutInitDisplayMode function. Arguments for this routine are assigned symbolic GLUT constants. For example, the following command specifies that a single refresh buffer is to be used for the display window and that we want to use the color mode which uses red, green, and blue (RGB) components to select color values:

    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);

The values of the constants passed to this function are combined using a logical or operation. Actually, single buffering and RGB color mode are the default options. But we will use the function now as a reminder that these are the options that are set for our display. Later, we discuss color modes in more detail, as well as other display options, such as double buffering for animation applications and selecting parameters for viewing three-dimensional scenes.

A Complete OpenGL Program

There are still a few more tasks to perform before we have all the parts that we need for a complete program. For the display window, we can choose a background color. And we need to construct a procedure that contains the appropriate OpenGL functions for the picture that we want to display.
Using RGB color values, we set the background color for the display window to be white, as in Figure 2, with the OpenGL function:

```c
glClearColor (1.0, 1.0, 1.0, 0.0);
```

The first three arguments in this function set the red, green, and blue component colors to the value 1.0, giving us a white background color for the display window. If, instead of 1.0, we set each of the component colors to 0.0, we would get a black background. And if all three of these components were set to the same intermediate value between 0.0 and 1.0, we would get some shade of gray. The fourth parameter in the `glClearColor` function is called the alpha value for the specified color. One use for the alpha value is as a “blending” parameter. When we activate the OpenGL blending operations, alpha values can be used to determine the resulting color for two overlapping objects. An alpha value of 0.0 indicates a totally transparent object, and an alpha value of 1.0 indicates an opaque object. Blending operations will not be used for a while, so the value of alpha is irrelevant to our early example programs. For now, we will simply set alpha to 0.0.

Although the `glClearColor` command assigns a color to the display window, it does not put the display window on the screen. To get the assigned window color displayed, we need to invoke the following OpenGL function:

```c
glClear (GL_COLOR_BUFFER_BIT);
```

The argument `GL_COLOR_BUFFER_BIT` is an OpenGL symbolic constant specifying that it is the bit values in the color buffer (refresh buffer) that are to be set to the values indicated in the `glClearColor` function. (OpenGL has several different kinds of buffers that can be manipulated.

In addition to setting the background color for the display window, we can choose a variety of color schemes for the objects we want to display in a scene. For our initial programming example, we will simply set the object color to be a dark green:

```c
glColor3f (0.0, 0.4, 0.2);
```

The suffix `3f` on the `glColor` function indicates that we are specifying the three RGB color components using floating-point (`f`) values. This function requires that the values be in the range from 0.0 to 1.0, and we have set red = 0.0, green = 0.4, and blue = 0.2.

For our first program, we simply display a two-dimensional line segment. To do this, we need to tell OpenGL how we want to “project” our picture onto the display window because generating a two-dimensional picture is treated by OpenGL as a special case of three-dimensional viewing. So, although we only want to produce a very simple two-dimensional line, OpenGL processes our picture through the full three-dimensional viewing operations. We can set the projection type (mode) and other viewing parameters that we need with the following two functions:

```c
glMatrixMode (GL_PROJECTION);
gluOrtho2D (0.0, 200.0, 0.0, 150.0);
```

This specifies that an orthogonal projection is to be used to map the contents of a two-dimensional rectangular area of world coordinates to the screen, and that the x-coordinate values within this rectangle range from 0.0 to 200.0 with y-coordinate values ranging from 0.0 to 150.0. Whatever objects we define
within this world-coordinate rectangle will be shown within the display window. Anything outside this coordinate range will not be displayed. Therefore, the GLU function `gluOrtho2D` defines the coordinate reference frame within the display window to be (0.0, 0.0) at the lower-left corner of the display window and (200.0, 150.0) at the upper-right window corner. Since we are only describing a two-dimensional object, the orthogonal projection has no other effect than to “paste” our picture into the display window that we defined earlier. For now, we will use a world-coordinate rectangle with the same aspect ratio as the display window, so that there is no distortion of our picture. Later, we will consider how we can maintain an aspect ratio that does not depend upon the display-window specification.

Finally, we need to call the appropriate OpenGL routines to create our line segment. The following code defines a two-dimensional, straight-line segment with integer, Cartesian endpoint coordinates (180, 15) and (10, 145).

```
  glBegin (GL_LINES);
    glVertex2i (180, 15);
    glVertex2i (10, 145);
  glEnd ( );
```

Now we are ready to put all the pieces together. The following OpenGL program is organized into three functions. We place all initializations and related one-time parameter settings in function `init`. Our geometric description of the “picture” that we want to display is in function `lineSegment`, which is the function that will be referenced by the GLUT function `glutDisplayFunc`. And the `main` function contains the GLUT functions for setting up the display window and getting our line segment onto the screen. Figure 3 shows the display window and line segment generated by this program.

**FIGURE 3**
The display window and line segment produced by the example program.
At the end of procedure lineSegment is a function, glFlush, that we have not yet discussed. This is simply a routine to force execution of our OpenGL functions, which are stored by computer systems in buffers in different locations, depending on how OpenGL is implemented. On a busy network, for example, there could be delays in processing some buffers. But the call to glFlush forces all such buffers to be emptied and the OpenGL functions to be processed.

The procedure lineSegment that we set up to describe our picture is referred to as a display callback function. And this procedure is described as being “registered” by glutDisplayFunc as the routine to invoke whenever the display window might need to be redisplayed. This can occur, for example, if the display window is moved. In subsequent chapters, we will look at other types of callback functions and the associated GLUT routines that we use to register them. In general, OpenGL programs are organized as a set of callback functions that are to be invoked when certain actions occur.
### Error Handling in OpenGL

Many of the features of the OpenGL API are extremely powerful. Unfortunately, they can also be very confusing, particularly for programmers who are just learning to use them. Although we would like to believe otherwise, it’s quite possible (if not likely) that our OpenGL programs may contain errors. Accordingly, it is worth spending a few moments discussing error handling in OpenGL programs.

The OpenGL and GLU libraries have a relatively simple method of recording errors. When OpenGL detects an error in a call to a base library routine or a GLU routine, it records an error code internally, and the routine which caused the error is ignored (so that it has no effect on either the internal OpenGL state or the contents of the frame buffer). However, OpenGL only records one error code at a time. Once an error occurs, no other error code will be recorded until your program explicitly queries the OpenGL error state:

```c
GLenum code;

code = glGetError();
```

This call returns the current error code and clears the internal error flag. If the returned value is equal to the OpenGL symbolic constant `GL_NO_ERROR`, everything is fine. Any other return value indicates that a problem has occurred.

The base OpenGL library defines a number of symbolic constants which represent different error conditions; the ones which occur most often are listed in Table 1. The GLU library also defines a number of error codes, but most of them have almost meaningless names such as `GLU_NURBS_ERROR1`, `GLU_NURBS_ERROR2`, and so on. (These are not actually meaningless names, but their meaning won’t become clear until we discuss more advanced concepts in later chapters.)

These symbolic constants are helpful, but printing them out directly is not particularly informative. Fortunately, the GLU library contains a function that returns a descriptive string for each of the GLU and GL errors. To use it, we first retrieve the current error code, and then pass it as a parameter to this function. The return value can be printed out using, for example, the C standard library `fprintf` function:

```c
#include <stdio.h>

GLenum code;

#include <stdio.h>
GLenum code;
```
const GLubyte *string;

code = glGetError ();
string = gluErrorString (code);
fprintf( stderr, "OpenGL error: %s\n", string );

The value returned by gluErrorString points to a string located inside the GLU library. It is not a dynamically-allocated string, so it must not be deallocated by our program. It also must not be modified by our program (hence the const modifier on the declaration of string).

We can easily encapsulate these function calls into a general error-reporting function in our program. The following function will retrieve the current error code, print the descriptive error string, and return the code to the calling routine:

```c
#include <stdio.h>

GLenum errorCheck ()
{
    GLenum code;
    const GLubyte *string;
    code = glGetError ();
    if (code != GL_NO_ERROR)
    {
        string = gluErrorString (code);
        fprintf( stderr, "OpenGL error: %s\n", string );
    }
    return code;
}
```

We encourage you to use a function like this in the OpenGL programs you develop. It is usually a good idea to check for errors at least once in your display callback routine, and more frequently if you are using OpenGL features you have never used before, or if you are seeing unusual or unexpected results in the image your program produces.

### 6 Summary

In this chapter, we surveyed the major features of graphics software systems. Some software systems, such as CAD packages and paint programs, are designed for particular applications. Other software systems provide a library of general graphics routines that can be used within a programming language such as C++ to generate pictures for any application.

Standard graphics-programming packages developed and approved through ISO and ANSI are GKS, 3D GKS, PHIGS, and PHIGS+. Other packages that have evolved into standards are GL and OpenGL. Many other graphics libraries are available for use in a programming language, including Open Inventor, VRML, RenderMan, Java 2D, and Java 3D. Other systems, such as Mathematica, MatLab, and Maple, often provide a set of graphics-programming functions.

Normally, graphics-programming packages require coordinate specifications to be given in Cartesian reference frames. Each object for a scene can be defined
in a separate modeling Cartesian-coordinate system, which is then mapped to
a world-coordinate location to construct the scene. From world coordinates,
three-dimensional objects are projected to a two-dimensional plane, converted
to normalized device coordinates, and then transformed to the final display-
device coordinates. The transformations from modeling coordinates to normal-
ized device coordinates are independent of particular output devices that might
be used in an application. Device drivers are then used to convert normalized
coordinates to integer device coordinates.

Functions that are available in graphics programming packages can be
divided into the following categories: graphics output primitives, attributes, geo-
metric and modeling transformations, viewing transformations, input functions,
picture-structuring operations, and control operations.

The OpenGL system consists of a device-independent set of routines (called
the core library), the utility library (GLU), and the utility toolkit (GLUT). In the
auxiliary set of routines provided by GLU, functions are available for generat-
ing complex objects, for parameter specifications in two-dimensional viewing
applications, for dealing with surface-rendering operations, and for performing
some other supporting tasks. In GLUT, we have an extensive set of functions for
managing display windows, interacting with screen-window systems, and for
generating some three-dimensional shapes. We can use GLUT to interface with
any computer system, or we can use GLX, Apple GL, WGL, or another system-
specific software package.

REFERENCES
Standard sources for information on OpenGL are Woo,
et al. (1999), Shreiner (2000), and Shreiner (2010). Open
Inventor is explored in Wernecke (1994). McCarthy and
Descartes (1998) can be consulted for discussions of VRML.
Presentations on RenderMan can be found in Upstill (1989) and Apodaca and Gritz (2000). Examples of
graphics programming in Java 2D are given in Knudsen
Graphics programming using Java 3D is explored in
Sowizral, et al. (2000); Palmer (2001); Selman (2002); and
Walsh and Gehringer (2002).
For information on PHIGS and PHIGS+, see
Howard, et al. (1991); Hopgood and Duce (1991);
Gaskins (1992); and Blake (1993). Information on the
two-dimensional GKS standard and on the evolution of
graphics standards is available in Hopgood, et al.

EXERCISES
1 What command could we use to set the color of an
OpenGL display window to dark gray? What command
would we use to set the color of the display window to white?
2 List the statements needed to set up an OpenGL
display window whose lower-left corner is at pixel
position (75, 200) and has a window width of 200
pixels and a height of 150 pixels.
3 List the OpenGL statements needed to draw a line
segment from the upper-right corner of a display
window of width 150 and height 250 to the lower-
left corner of the window.
4 Suppose we have written a function called rectangle
whose purpose is to draw a rectangle in the
middle of a given display window. What OpenGL
statement would be needed to make sure the rect-
angle is drawn correctly each time the display window
needs to be repainted?
5 Explain what is meant by the term “OpenGL display
callback function”.
6 Explain the difference between modeling coordi-

nates and world coordinates.
7 Explain what normalized coordinates are and why
they are useful for graphics software packages.

IN MORE DEPTH
1 Develop a few application ideas and iden-
tify and describe the objects that you will be
manipulating graphically in the application.
Explain desirable physical and visual proper-
ties of these objects in detail so that you
can concretely define what attributes you will
develop for them in later exercises. Consider the
following:
Computer Graphics Software

- Are the objects complex in shape or texture?
- Can the objects be approximated fairly well by simpler shapes?
- Are some of the objects comprised of fairly complex curved surfaces?
- Do the objects lend themselves to being represented two-dimensionally at first, even if unrealistically?
- Can the objects be represented as a hierarchically organized group of smaller objects, or parts?
- Will the objects change position and orientation dynamically in response to user input?
- Will the lighting conditions change in the application, and do the object appearances change under these varying conditions?

If all of the objects in your application yield a “no” answer to any of these questions, consider modifying the application or your design and implementation approach so that such properties are present in at least one of the objects. Alternatively, come up with two or more applications so that at least one of them contains objects that satisfy each of these properties. Draw a rough specification, using visual diagrams and/or text, listing the properties of the objects in your application. You will use and modify this specification as you continue to develop your application.

2 You will be developing your application using the OpenGL API. In order to do this, you will need to have a working programming environment in which you can edit, compile and run OpenGL programs. For this exercise, you should go through the necessary steps to get this environment up and running on your machine. Once you have the environment set up, create a new project with the source code given in the example in this chapter that draws a single line in the display window. Make sure you can compile and run this program. Information about getting an OpenGL environment up and running on your system can be found at http://www.opengl.org/ and http://www.opengl.org/wiki/Getting_started.

Make sure that you obtain all of the necessary libraries, including the GLU and GLUT libraries. The examples in this text are all written in C++. Follow your instructor’s guidelines on the use of other languages (provided OpenGL bindings can be obtained for them).
A general software package for graphics applications, sometimes referred to as a computer-graphics application programming interface (CG API), provides a library of functions that we can use within a programming language such as C++ to create pictures. The set of library functions can be subdivided into several categories. One of the first things we need to do when creating a picture is to describe the component parts of the scene to be displayed. Picture components could be trees and terrain, furniture and walls, storefronts and street scenes, automobiles and billboards, atoms and molecules, or stars and galaxies. For each type of scene, we need to describe the structure of the individual objects and their coordinate locations within the scene. Those functions in a graphics package that we use to describe the various picture components are called the graphics output primitives, or simply primitives. The output primitives describing the geometry of objects are typically referred to as geometric primitives. Point positions and straight-line segments are the simplest
geometric primitives. Additional geometric primitives that can be available in a graphics package include circles and other conic sections, quadric surfaces, spline curves and surfaces, and polygon color areas. Also, most graphics systems provide some functions for displaying character strings. After the geometry of a picture has been specified within a selected coordinate reference frame, the output primitives are projected to a two-dimensional plane, corresponding to the display area of an output device, and scan converted into integer pixel positions within the frame buffer.

In this chapter, we introduce the output primitives available in OpenGL, and discuss their use.

1 Coordinate Reference Frames

To describe a picture, we first decide upon a convenient Cartesian coordinate system, called the world-coordinate reference frame, which could be either two-dimensional or three-dimensional. We then describe the objects in our picture by giving their geometric specifications in terms of positions in world coordinates. For instance, we define a straight-line segment with two endpoint positions, and a polygon is specified with a set of positions for its vertices. These coordinate positions are stored in the scene description along with other information about the objects, such as their color and their coordinate extents, which are the minimum and maximum \( x \), \( y \), and \( z \) values for each object. A set of coordinate extents is also described as a bounding box for an object. For a two-dimensional figure, the coordinate extents are sometimes called an object's bounding rectangle. Objects are then displayed by passing the scene information to the viewing routines, which identify visible surfaces and ultimately map the objects to positions on the video monitor. The scan-conversion process stores information about the scene, such as color values, at the appropriate locations in the frame buffer, and the objects in the scene are displayed on the output device.

Screen Coordinates

Locations on a video monitor are referenced in integer screen coordinates, which correspond to the pixel positions in the frame buffer. Pixel coordinate values give the scan line number (the \( y \) value) and the column number (the \( x \) value along a scan line). Hardware processes, such as screen refreshing, typically address pixel positions with respect to the top-left corner of the screen. Scan lines are then referenced from 0, at the top of the screen, to some integer value, \( y_{\text{max}} \), at the bottom of the screen, and pixel positions along each scan line are numbered from 0 to \( x_{\text{max}} \), left to right. However, with software commands, we can set up any convenient reference frame for screen positions. For example, we could specify an integer range for screen positions with the coordinate origin at the lower-left of a screen area (Figure 1), or we could use noninteger Cartesian values for a picture description. The coordinate values we use to describe the geometry of a scene are then converted by the viewing routines to integer pixel positions within the frame buffer.

Scan-line algorithms for the graphics primitives use the defining coordinate descriptions to determine the locations of pixels that are to be displayed. For
example, given the endpoint coordinates for a line segment, a display algorithm must calculate the positions for those pixels that lie along the line path between the endpoints. Since a pixel position occupies a finite area of the screen, the finite size of a pixel must be taken into account by the implementation algorithms. For the present, we assume that each integer screen position references the center of a pixel area.

Once pixel positions have been identified for an object, the appropriate color values must be stored in the frame buffer. For this purpose, we will assume that we have available a low-level procedure of the form

```plaintext
setPixel (x, y);
```

This procedure stores the current color setting into the frame buffer at integer position \((x, y)\), relative to the selected position of the screen-coordinate origin. We sometimes also will want to be able to retrieve the current frame-buffer setting for a pixel location. So we will assume that we have the following low-level function for obtaining a frame-buffer color value:

```plaintext
getPixel (x, y, color);
```

In this function, parameter \(color\) receives an integer value corresponding to the combined red, green, and blue (RGB) bit codes stored for the specified pixel at position \((x, y)\).

Although we need only specify color values at \((x, y)\) positions for a two-dimensional picture, additional screen-coordinate information is needed for three-dimensional scenes. In this case, screen coordinates are stored as three-dimensional values, where the third dimension references the depth of object positions relative to a viewing position. For a two-dimensional scene, all depth values are 0.

**Absolute and Relative Coordinate Specifications**

So far, the coordinate references that we have discussed are stated as **absolute coordinate** values. This means that the values specified are the actual positions within the coordinate system in use.

However, some graphics packages also allow positions to be specified using **relative coordinates**. This method is useful for various graphics applications, such as producing drawings with pen plotters, artist’s drawing and painting systems, and graphics packages for publishing and printing applications. Taking this approach, we can specify a coordinate position as an offset from the last position that was referenced (called the **current position**). For example, if location \((3, 8)\) is the last position that has been referenced in an application program, a relative coordinate specification of \((2, -1)\) corresponds to an absolute position of \((5, 7)\). An additional function is then used to set a current position before any coordinates for primitive functions are specified. To describe an object, such as a series of connected line segments, we then need to give only a sequence of relative coordinates (offsets), once a starting position has been established. Options can be provided in a graphics system to allow the specification of locations using either relative or absolute coordinates. In the following discussions, we will assume that all coordinates are specified as absolute references unless explicitly stated otherwise.
2 Specifying A Two-Dimensional World-Coordinate Reference Frame in OpenGL

The `gluOrtho2D` command is a function we can use to set up any two-dimensional Cartesian reference frame. The arguments for this function are the four values defining the \( x \) and \( y \) coordinate limits for the picture we want to display. Since the `gluOrtho2D` function specifies an orthogonal projection, we need also to be sure that the coordinate values are placed in the OpenGL projection matrix. In addition, we could assign the identity matrix as the projection matrix before defining the world-coordinate range. This would ensure that the coordinate values were not accumulated with any values we may have previously set for the projection matrix. Thus, for our initial two-dimensional examples, we can define the coordinate frame for the screen display window with the following statements:

```c
glMatrixMode (GL_PROJECTION);
glLoadIdentity ();
gluOrtho2D (xmin, xmax, ymin, ymax);
```

The display window will then be referenced by coordinates \((x_{\min}, y_{\min})\) at the lower-left corner and by coordinates \((x_{\max}, y_{\max})\) at the upper-right corner, as shown in Figure 2.

We can then designate one or more graphics primitives for display using the coordinate reference specified in the `gluOrtho2D` statement. If the coordinate extents of a primitive are within the coordinate range of the display window, all of the primitive will be displayed. Otherwise, only those parts of the primitive within the display-window coordinate limits will be shown. Also, when we set up the geometry describing a picture, all positions for the OpenGL primitives must be given in absolute coordinates, with respect to the reference frame defined in the `gluOrtho2D` function.

![Video Screen
Display Window

**F I G U R E** 2
World-coordinate limits for a display window, as specified in the `gluOrtho2D` function.
3 OpenGL Point Functions

To specify the geometry of a point, we simply give a coordinate position in the world reference frame. Then this coordinate position, along with other geometric descriptions we may have in our scene, is passed to the viewing routines. Unless we specify other attribute values, OpenGL primitives are displayed with a default size and color. The default color for primitives is white, and the default point size is equal to the size of a single screen pixel.

We use the following OpenGL function to state the coordinate values for a single position:

```c
glVertex* ( );
```

where the asterisk (*) indicates that suffix codes are required for this function. These suffix codes are used to identify the spatial dimension, the numerical data type to be used for the coordinate values, and a possible vector form for the coordinate specification. Calls to `glVertex` functions must be placed between a `glBegin` function and a `glEnd` function. The argument of the `glBegin` function is used to identify the kind of output primitive that is to be displayed, and `glEnd` takes no arguments. For point plotting, the argument of the `glBegin` function is the symbolic constant `GL_POINTS`. Thus, the form for an OpenGL specification of a point position is

```c
glBegin (GL_POINTS);
    glVertex* ( );
    glEnd ( );
```

Although the term *vertex* strictly refers to a “corner” point of a polygon, the point of intersection of the sides of an angle, a point of intersection of an ellipse with its major axis, or other similar coordinate positions on geometric structures, the `glVertex` function is used in OpenGL to specify coordinates for any point position. In this way, a single function is used for point, line, and polygon specifications—and, most often, polygon patches are used to describe the objects in a scene.

Coordinate positions in OpenGL can be given in two, three, or four dimensions. We use a suffix value of 2, 3, or 4 on the `glVertex` function to indicate the dimensionality of a coordinate position. A four-dimensional specification indicates a homogeneous-coordinate representation, where the homogeneous parameter $h$ (the fourth coordinate) is a scaling factor for the Cartesian-coordinate values. Homogeneous-coordinate representations are useful for expressing transformation operations in matrix form. Because OpenGL treats two-dimensions as a special case of three dimensions, any $(x, y)$ coordinate specification is equivalent to a three-dimensional specification of $(x, y, 0)$. Furthermore, OpenGL represents vertices internally in four dimensions, so each of these specifications are equivalent to the four-dimensional specification $(x, y, 0, 1)$.

We also need to state which data type is to be used for the numerical-value specifications of the coordinates. This is accomplished with a second suffix code on the `glVertex` function. Suffix codes for specifying a numerical data type are `i` (integer), `s` (short), `f` (float), and `d` (double). Finally, the coordinate values can be listed explicitly in the `glVertex` function, or a single argument can be used that references a coordinate position as an array. If we use an array specification for a coordinate position, we need to append `v` (for “vector”) as a third suffix code.
In the following example, three equally spaced points are plotted along a two-
dimensional, straight-line path with a slope of 2 (see Figure 3). Coordinates are
given as integer pairs:

```c
glBegin (GL_POINTS);
glVertex2i (50, 100);
glVertex2i (75, 150);
glVertex2i (100, 200);
glEnd ( );
```

Alternatively, we could specify the coordinate values for the preceding points in
arrays such as

```c
int point1 [] = {50, 100};
int point2 [] = {75, 150};
int point3 [] = {100, 200};
```

and call the OpenGL functions for plotting the three points as

```c
glBegin (GL_POINTS);
glVertex2iv (point1);
glVertex2iv (point2);
glVertex2iv (point3);
glEnd ( );
```

In addition, here is an example of specifying two point positions in a three-
dimensional world reference frame. In this case, we give the coordinates as
explicit floating-point values:

```c
glBegin (GL_POINTS);
glVertex3f (-78.05, 909.72, 14.60);
glVertex3f (261.91, -5200.67, 188.33);
glEnd ( );
```

We could also define a C++ class or structure (struct) for specifying point
positions in various dimensions. For example,

```c
class wcPt2D {
public:
    GLfloat x, y;
};
```
Using this class definition, we could specify a two-dimensional, world-coordinate point position with the statements

```cpp
wcPt2D pointPos;

pointPos.x = 120.75;
pointPos.y = 45.30;
glBegin (GL_POINTS);
    glVertex2f (pointPos.x, pointPos.y);
glEnd ( );
```

Also, we can use the OpenGL point-plotting functions within a C++ procedure to implement the `setPixel` command.

---

### 4 OpenGL Line Functions

Graphics packages typically provide a function for specifying one or more straight-line segments, where each line segment is defined by two endpoint coordinate positions. In OpenGL, we select a single endpoint coordinate position using the `glVertex` function, just as we did for a point position. And we enclose a list of `glVertex` functions between the `glBegin`/`glEnd` pair. But now we use a symbolic constant as the argument for the `glBegin` function that interprets a list of positions as the endpoint coordinates for line segments. There are three symbolic constants in OpenGL that we can use to specify how a list of endpoint positions should be connected to form a set of straight-line segments. By default, each symbolic constant displays solid, white lines.

A set of straight-line segments between each successive pair of endpoints in a list is generated using the primitive line constant `GL_LINES`. In general, this will result in a set of unconnected lines unless some coordinate positions are repeated, because OpenGL considers lines to be connected only if they share a vertex; lines that cross but do not share a vertex are still considered to be unconnected. Nothing is displayed if only one endpoint is specified, and the last endpoint is not processed if the number of endpoints listed is odd. For example, if we have five coordinate positions, labeled `p1` through `p5`, and each is represented as a two-dimensional array, then the following code could generate the display shown in Figure 4(a):

```cpp
glBegin (GL_LINES);
    glVertex2iv (p1);
    glVertex2iv (p2);
    glVertex2iv (p3);
    glVertex2iv (p4);
    glVertex2iv (p5);
glEnd ( );
```

Thus, we obtain one line segment between the first and second coordinate positions and another line segment between the third and fourth positions. In this case, the number of specified endpoints is odd, so the last coordinate position is ignored.

With the OpenGL primitive constant `GL_LINE_STRIP`, we obtain a **polyline**. In this case, the display is a sequence of connected line segments between the first endpoint in the list and the last endpoint. The first line segment in the polyline is displayed between the first endpoint and the second endpoint; the second line segment is between the second and third endpoints; and so forth, up to the last line endpoint. Nothing is displayed if we do not list at least two coordinate positions.
FIGURE 4
Line segments that can be displayed in OpenGL using a list of five endpoint coordinates. (a) An unconnected set of lines generated with the primitive line constant GL_LINES. (b) A polyline generated with GL_LINE_STRIP. (c) A closed polyline generated with GL_LINE_LOOP.

Using the same five coordinate positions as in the previous example, we obtain the display in Figure 4(b) with the code

```c
glBegin (GL_LINE_STRIP);
glVertex2iv (p1);
glVertex2iv (p2);
glVertex2iv (p3);
glVertex2iv (p4);
glVertex2iv (p5);
glEnd ( );
```

The third OpenGL line primitive is GL_LINE_LOOP, which produces a **closed polyline**. Lines are drawn as with GL_LINE_STRIP, but an additional line is drawn to connect the last coordinate position and the first coordinate position. Figure 4(c) shows the display of our endpoint list when we select this line option, using the code

```c
glBegin (GL_LINE_LOOP);
glVertex2iv (p1);
glVertex2iv (p2);
glVertex2iv (p3);
glVertex2iv (p4);
glVertex2iv (p5);
glEnd ( );
```

As noted earlier, picture components are described in a world-coordinate reference frame that is eventually mapped to the coordinate reference for the output device. Then the geometric information about the picture is scan-converted to pixel positions.

### 5 OpenGL Curve Functions

Routines for generating basic curves, such as circles and ellipses, are not included as primitive functions in the OpenGL core library. But this library does contain functions for displaying Bézier splines, which are polynomials that are defined with a discrete point set. And the OpenGL Utility (GLU) library has routines for three-dimensional quadrics, such as spheres and cylinders, as well as
routines for producing rational B-splines, which are a general class of splines that include the simpler Bézier curves. Using rational B-splines, we can display circles, ellipses, and other two-dimensional quadrics. In addition, there are routines in the OpenGL Utility Toolkit (GLUT) that we can use to display some three-dimensional quadrics, such as spheres and cones, and some other shapes. However, all these routines are more involved than the basic primitives we introduce in this chapter.

Another method we can use to generate a display of a simple curve is to approximate it using a polyline. We just need to locate a set of points along the curve path and connect the points with straight-line segments. The more line sections we include in the polyline, the smoother the appearance of the curve. As an example, Figure 5 illustrates various polyline displays that could be used for a circle segment.

A third alternative is to write our own curve-generation functions based on the algorithms presented in following chapters.

6 Fill-Area Primitives

Another useful construct, besides points, straight-line segments, and curves, for describing components of a picture is an area that is filled with some solid color or pattern. A picture component of this type is typically referred to as a fill area or a filled area. Most often, fill areas are used to describe surfaces of solid objects, but they are also useful in a variety of other applications. Also, fill regions are usually planar surfaces, mainly polygons. But, in general, there are many possible shapes for a region in a picture that we might wish to fill with a color option. Figure 6 illustrates a few possible fill-area shapes. For the present, we assume that all fill areas are to be displayed with a specified solid color.

Although any fill-area shape is possible, graphics libraries generally do not support specifications for arbitrary fill shapes. Most library routines require that
a fill area be specified as a polygon. Graphics routines can more efficiently process polygons than other kinds of fill shapes because polygon boundaries are described with linear equations. Moreover, most curved surfaces can be approximated reasonably well with a set of polygon patches, just as a curved line can be approximated with a set of straight-line segments. In addition, when lighting effects and surface-shading procedures are applied, an approximated curved surface can be displayed quite realistically. Approximating a curved surface with polygon facets is sometimes referred to as surface tessellation, or fitting the surface with a polygon mesh. Figure 7 shows the side and top surfaces of a metal cylinder approximated in an outline form as a polygon mesh. Displays of such figures can be generated quickly as wire-frame views, showing only the polygon edges to give a general indication of the surface structure. Then the wire-frame model could be shaded to generate a display of a natural-looking material surface. Objects described with a set of polygon surface patches are usually referred to as standard graphics objects, or just graphics objects.

In general, we can create fill areas with any boundary specification, such as a circle or connected set of spline-curve sections. And some of the polygon methods discussed in the next section can be adapted to display fill areas with a nonlinear border.

## 7 Polygon Fill Areas

Mathematically defined, a **polygon** is a plane figure specified by a set of three or more coordinate positions, called **vertices**, that are connected in sequence by straight-line segments, called the **edges** or **sides** of the polygon. Further, in basic geometry, it is required that the polygon edges have no common point other than their endpoints. Thus, by definition, a polygon must have all its vertices within a single plane and there can be no edge crossings. Examples of polygons include triangles, rectangles, octagons, and decagons. Sometimes, any plane figure with a closed-polyline boundary is alluded to as a polygon, and one with no crossing edges is referred to as a **standard polygon** or a **simple polygon**. In an effort to avoid ambiguous object references, we will use the term **polygon** to refer only to those planar shapes that have a closed-polyline boundary and no edge crossings.

For a computer-graphics application, it is possible that a designated set of polygon vertices do not all lie exactly in one plane. This can be due to round-off error in the calculation of numerical values, to errors in selecting coordinate positions for the vertices, or, more typically, to approximating a curved surface with a set of polygonal patches. One way to rectify this problem is simply to divide the specified surface mesh into triangles. But in some cases, there may be reasons...
Another way to identify a concave polygon is to look at the polygon vertex positions relative to the extension line of any edge. If some vertices are on one side of the extension line and some vertices are on the other side, the polygon is concave.

Splitting Concave Polygons

Once we have identified a concave polygon, we can split it into a set of convex polygons. This can be accomplished using edge vectors and edge cross-products; or, we can use vertex positions relative to an edge extension line to determine which vertices are on one side of this line and which are on the other. For the following algorithms, we assume that all polygons are in the $xy$ plane. Of course, the original position of a polygon described in world coordinates may not be in the $xy$ plane, but we can always move it into that plane.

With the vector method for splitting a concave polygon, we first need to form the edge vectors. Given two consecutive vertex positions, $V_k$ and $V_{k+1}$, we define the edge vector between them as

$$E_k = V_{k+1} - V_k$$

Next we calculate the cross-products of successive edge vectors in order around the polygon perimeter. If the $z$ component of some cross-products is positive while other cross-products have a negative $z$ component, the polygon is concave. Otherwise, the polygon is convex. This assumes that no series of three successive vertices are collinear, in which case the cross-product of the two edge vectors for these vertices would be zero. If all vertices are collinear, we have a degenerate polygon (a straight line). We can apply the vector method by processing edge vectors in counterclockwise order. If any cross-product has a negative $z$ component (as in Figure 9), the polygon is concave and we can split it along the line of the first edge vector in the cross-product pair. The following example illustrates this method for splitting a concave polygon.

**Example 1**  The Vector Method for Splitting Concave Polygons

Figure 10 shows a concave polygon with six edges. Edge vectors for this polygon can be expressed as

$$E_1 = (1, 0, 0)$$  \hspace{1cm}  $$E_2 = (1, 1, 0)$$

$$E_3 = (1, -1, 0)$$  \hspace{1cm}  $$E_4 = (0, 2, 0)$$

$$E_5 = (-3, 0, 0)$$  \hspace{1cm}  $$E_6 = (0, -2, 0)$$
to retain the original shape of the mesh patches, so methods have been devised for approximating a nonplanar polygonal shape with a plane figure. We discuss how these plane approximations are calculated in the section on plane equations.

Polygon Classifications

An interior angle of a polygon is an angle inside the polygon boundary that is formed by two adjacent edges. If all interior angles of a polygon are less than or equal to 180°, the polygon is convex. An equivalent definition of a convex polygon is that its interior lies completely on one side of the infinite extension line of any one of its edges. Also, if we select any two points in the interior of a convex polygon, the line segment joining the two points is also in the interior. A polygon that is not convex is called a concave polygon. Figure 8 gives examples of convex and concave polygons.

The term degenerate polygon is often used to describe a set of vertices that are collinear or that have repeated coordinate positions. Collinear vertices generate a line segment. Repeated vertex positions can generate a polygon shape with extraneous lines, overlapping edges, or edges that have a length equal to 0. Sometimes the term degenerate polygon is also applied to a vertex list that contains fewer than three coordinate positions.

To be robust, a graphics package could reject degenerate or nonplanar vertex sets. But this requires extra processing to identify these problems, so graphics systems usually leave such considerations to the programmer.

Concave polygons also present problems. Implementations of fill algorithms and other graphics routines are more complicated for concave polygons, so it is generally more efficient to split a concave polygon into a set of convex polygons before processing. As with other polygon preprocessing algorithms, concave polygon splitting is often not included in a graphics library. Some graphics packages, including OpenGL, require all fill polygons to be convex. And some systems accept only triangular fill areas, which greatly simplifies many of the display algorithms.

Identifying Concave Polygons

A concave polygon has at least one interior angle greater than 180°. Also, the extension of some edges of a concave polygon will intersect other edges, and some pair of interior points will produce a line segment that intersects the polygon boundary. Therefore, we can use any one of these characteristics of a concave polygon as a basis for constructing an identification algorithm.

If we set up a vector for each polygon edge, then we can use the cross-product of adjacent edges to test for concavity. All such vector products will be of the same sign (positive or negative) for a convex polygon. Therefore, if some cross-products yield a positive value and some a negative value, we have a concave polygon. Figure 9 illustrates the edge-vector, cross-product method for identifying concave polygons.
where the $z$ component is 0, since all edges are in the $xy$ plane. The cross-product $E_j \times E_k$ for two successive edge vectors is a vector perpendicular to the $xy$ plane with $z$ component equal to $E_{jz}E_{ky} - E_{kx}E_{jy}$:

\[
E_1 \times E_2 = (0, 0, 1) \quad E_2 \times E_3 = (0, 0, -2) \\
E_3 \times E_4 = (0, 0, 2) \quad E_4 \times E_5 = (0, 0, 6) \\
E_5 \times E_6 = (0, 0, 6) \quad E_6 \times E_1 = (0, 0, 2)
\]

Since the cross-product $E_2 \times E_3$ has a negative $z$ component, we split the polygon along the line of vector $E_2$. The line equation for this edge has a slope of 1 and a $y$ intercept of $-1$. We then determine the intersection of this line with the other polygon edges to split the polygon into two pieces. No other edge cross-products are negative, so the two new polygons are both convex.

We can also split a concave polygon using a rotational method. Proceeding counterclockwise around the polygon edges, we shift the position of the polygon so that each vertex $V_k$ in turn is at the coordinate origin. Then, we rotate the polygon about the origin in a clockwise direction so that the next vertex $V_{k+1}$ is on the $x$ axis. If the following vertex, $V_{k+2}$, is below the $x$ axis, the polygon is concave. We then split the polygon along the $x$ axis to form two new polygons, and we repeat the concave test for each of the two new polygons. These steps are repeated until we have tested all vertices in the polygon list. Figure 11 illustrates the rotational method for splitting a concave polygon.

### Splitting a Convex Polygon into a Set of Triangles

Once we have a vertex list for a convex polygon, we could transform it into a set of triangles. This can be accomplished by first defining any sequence of three consecutive vertices to be a new polygon (a triangle). The middle triangle vertex is then deleted from the original vertex list. Then the same procedure is applied to this modified vertex list to strip off another triangle. We continue forming triangles in this manner until the original polygon is reduced to just three vertices, which define the last triangle in the set. A concave polygon can also be divided into a set of triangles using this approach, although care must be taken that the new diagonal edge formed by joining the first and third selected vertices does not cross the concave portion of the polygon, and that the three selected vertices at each step form an interior angle that is less than 180° (a “convex” angle).

### Inside-Outside Tests

Various graphics processes often need to identify interior regions of objects. Identifying the interior of a simple object, such as a convex polygon, a circle, or a sphere, is generally a straightforward process. But sometimes we must deal with more complex objects. For example, we may want to specify a complex fill region with intersecting edges, as in Figure 12. For such shapes, it is not always clear which regions of the $xy$ plane we should call “interior” and which regions we should designate as “exterior” to the object boundaries. Two commonly used algorithms for identifying interior areas of a plane figure are the odd-even rule and the nonzero winding-number rule.

We apply the **odd-even rule**, also called the odd-parity rule or the even-odd rule, by first conceptually drawing a line from any position $P$ to a distant point
outside the coordinate extents of the closed polyline. Then we count the number of line-segment crossings along this line. If the number of segments crossed by this line is odd, then \( P \) is considered to be an interior point. Otherwise, \( P \) is an exterior point. To obtain an accurate count of the segment crossings, we must be sure that the line path we choose does not intersect any line-segment endpoints. Figure 12(a) shows the interior and exterior regions obtained using the odd-even rule for a self-intersecting closed polyline. We can use this procedure, for example, to fill the interior region between two concentric circles or two concentric polygons with a specified color.

Another method for defining interior regions is the nonzero winding-number rule, which counts the number of times that the boundary of an object “winds” around a particular point in the counterclockwise direction. This count is called the winding number, and the interior points of a two-dimensional object can be defined to be those that have a nonzero value for the winding number. We apply the nonzero winding number rule by initializing the winding number to 0 and again imagining a line drawn from any position \( P \) to a distant point beyond the coordinate extents of the object. The line we choose must not pass through any endpoint coordinates. As we move along the line from position \( P \) to the distant point, we count the number of object line segments that cross the reference line in each direction. We add 1 to the winding number every time we intersect a segment that crosses the line in the direction from right to left, and we subtract 1 every time we intersect a segment that crosses from left to right. The final value of the winding number, after all boundary crossings have been counted, determines the relative position of \( P \). If the winding number is nonzero, \( P \) is considered to be an interior point. Otherwise, \( P \) is taken to be an exterior point. Figure 12(b) shows the interior and exterior regions defined by the nonzero winding-number rule for a self-intersecting, closed polyline. For simple objects, such as polygons and circles, the nonzero winding-number rule and the odd-even rule give the same results. But for more complex shapes, the two methods may yield different interior and exterior regions, as in the example of Figure 12.

One way to determine directional boundary crossings is to set up vectors along the object edges (or boundary lines) and along the reference line. Then we compute the vector cross-product of the vector \( \mathbf{u} \), along the line from \( P \) to a distant point, with an object edge vector \( \mathbf{E} \) for each edge that crosses the line. Assuming that we have a two-dimensional object in the \( xy \) plane, the direction of each vector cross-product will be either in the \( +z \) direction or in the \( -z \) direction. If the \( z \) component of a cross-product \( \mathbf{u} \times \mathbf{E} \) for a particular crossing is positive, that segment crosses from right to left and we add 1 to the winding number.
Otherwise, the segment crosses from left to right and we subtract 1 from the winding number.

A somewhat simpler way to compute directional boundary crossings is to use vector dot products instead of cross-products. To do this, we set up a vector that is perpendicular to vector $\mathbf{u}$ and that has a right-to-left direction as we look along the line from $\mathbf{P}$ in the direction of $\mathbf{u}$. If the components of $\mathbf{u}$ are denoted as $(u_x, u_y)$, then the vector that is perpendicular to $\mathbf{u}$ has components $(-u_y, u_x)$. Now, if the dot product of this perpendicular vector and a boundary-line vector is positive, that crossing is from right to left and we add 1 to the winding number. Otherwise, the boundary crosses our reference line from left to right, and we subtract 1 from the winding number.

The nonzero winding-number rule tends to classify as interior some areas that the odd-even rule deems to be exterior, and it can be more versatile in some applications. In general, plane figures can be defined with multiple, disjoint components, and the direction specified for each set of disjoint boundaries can be used to designate the interior and exterior regions. Examples include characters (such as letters of the alphabet and punctuation symbols), nested polygons, and concentric circles or ellipses. For curved lines, the odd-even rule is applied by calculating intersections with the curve paths. Similarly, with the nonzero winding-number rule, we need to calculate tangent vectors to the curves at the crossover intersection points with the reference line from position $\mathbf{P}$.

Variations of the nonzero winding-number rule can be used to define interior regions in other ways. For example, we could define a point to be interior if its winding number is positive or if it is negative; or we could use any other rule to generate a variety of fill shapes. Sometimes, Boolean operations are used to specify a fill area as a combination of two regions. One way to implement Boolean operations is by using a variation of the basic winding-number rule. With this scheme, we first define a simple, nonintersecting boundary for each of two regions. Then if we consider the direction for each boundary to be counterclockwise, the union of two regions would consist of those points whose winding number is positive (Figure 13). Similarly, the intersection of two regions with counterclockwise boundaries would contain those points whose winding number is greater than 1, as illustrated in Figure 14. To set up a fill area that is the difference of two regions (say, $A - B$), we can enclose region $A$ with a counterclockwise border and $B$ with a clockwise border. Then the difference region (Figure 15) is the set of all points whose winding number is positive.
A fill area defined as a region with a positive value for the winding number. This fill area is the difference, $A - B$, of two regions, where region A has a positive border direction (counterclockwise) and region B has a negative border direction (clockwise).

### Polygon Tables

Typically, the objects in a scene are described as sets of polygon surface facets. In fact, graphics packages often provide functions for defining a surface shape as a mesh of polygon patches. The description for each object includes coordinate information specifying the geometry for the polygon facets and other surface parameters such as color, transparency, and light-reflection properties. As information for each polygon is input, the data are placed into tables that are to be used in the subsequent processing, display, and manipulation of the objects in the scene. These polygon data tables can be organized into two groups: geometric tables and attribute tables. Geometric data tables contain vertex coordinates and parameters to identify the spatial orientation of the polygon surfaces. Attribute information for an object includes parameters specifying the degree of transparency of the object and its surface reflectivity and texture characteristics.

Geometric data for the objects in a scene are arranged conveniently in three lists: a vertex table, an edge table, and a surface-facet table. Coordinate values for each vertex in the object are stored in the vertex table. The edge table contains pointers back into the vertex table to identify the vertices for each polygon edge. And the surface-facet table contains pointers back into the edge table to identify the edges for each polygon. This scheme is illustrated in Figure 16 for two adjacent polygon surface facets, formed with six edges and five vertices.
adjacent polygon facets on an object surface. In addition, individual objects and
their component polygon faces can be assigned object and facet identifiers for
easy reference.

Listing the geometric data in three tables, as in Figure 16, provides a conve-
nient reference to the individual components (vertices, edges, and surface facets)
for each object. Also, the object can be displayed efficiently by using data from
the edge table to identify polygon boundaries. An alternative arrangement is to
use just two tables: a vertex table and a surface-facet table. But this scheme is
less convenient, and some edges could get drawn twice in a wire-frame dis-
play. Another possibility is to use only a surface-facet table, but this dupli-
cates coordinate information, since explicit coordinate values are listed for each
vertex in each polygon facet. Also the relationship between edges and facets
would have to be reconstructed from the vertex listings in the surface-facet
table.

We can add extra information to the data tables of Figure 16 for faster
information extraction. For instance, we could expand the edge table to include
forward pointers into the surface-facet table so that a common edge between
polygons could be identified more rapidly (Figure 17). This is particularly useful
for rendering procedures that must vary surface shading smoothly across the
dges from one polygon to the next. Similarly, the vertex table could be expanded
to reference corresponding edges, for faster information retrieval.

Additional geometric information that is usually stored in the data tables
includes the slope for each edge and the coordinate extents for polygon edges,
polygon facets, and each object in a scene. As vertices are input, we can calculate
derge slopes, and we can scan the coordinate values to identify the minimum and
maximum $x$, $y$, and $z$ values for individual lines and polygons. Edge slopes and
bounding-box information are needed in subsequent processing, such as surface
rendering and visible-surface identification algorithms.

Because the geometric data tables may contain extensive listings of vertices
and edges for complex objects and scenes, it is important that the data be checked
for consistency and completeness. When vertex, edge, and polygon definitions
are specified, it is possible, particularly in interactive applications, that certain
input errors could be made that would distort the display of the objects. The
more information included in the data tables, the easier it is to check for errors.
Therefore, error checking is easier when three data tables (vertex, edge, and sur-
fance facet) are used, since this scheme provides the most information. Some of
the tests that could be performed by a graphics package are (1) that every vertex
is listed as an endpoint for at least two edges, (2) that every edge is part of at
least one polygon, (3) that every polygon is closed, (4) that each polygon has at
least one shared edge, and (5) that if the edge table contains pointers to polygons,
every edge referenced by a polygon pointer has a reciprocal pointer back to the
polygon.

Plane Equations

To produce a display of a three-dimensional scene, a graphics system processes
the input data through several procedures. These procedures include transfor-
mation of the modeling and world-coordinate descriptions through the viewing
pipeline, identification of visible surfaces, and the application of rendering rou-
tines to the individual surface facets. For some of these processes, information
about the spatial orientation of the surface components of objects is needed. This
information is obtained from the vertex coordinate values and the equations that
describe the polygon surfaces.
Each polygon in a scene is contained within a plane of infinite extent. The general equation of a plane is

$$Ax + By + Cz + D = 0$$

where \((x, y, z)\) is any point on the plane, and the coefficients \(A, B, C,\) and \(D\) (called plane parameters) are constants describing the spatial properties of the plane. We can obtain the values of \(A, B, C,\) and \(D\) by solving a set of three plane equations using the coordinate values for three noncollinear points in the plane. For this purpose, we can select three successive convex-polygon vertices, \((x_1, y_1, z_1), (x_2, y_2, z_2),\) and \((x_3, y_3, z_3),\) in a counterclockwise order and solve the following set of simultaneous linear plane equations for the ratios \(A/D, B/D,\) and \(C/D:\)

$$(A/D)x_k + (B/D)y_k + (C/D)z_k = -1, \quad k = 1, 2, 3$$

The solution to this set of equations can be obtained in determinant form, using Cramer’s rule, as

$$A = \begin{vmatrix} 1 & y_1 & z_1 \\ 1 & y_2 & z_2 \\ 1 & y_3 & z_3 \end{vmatrix} \quad B = \begin{vmatrix} x_1 & 1 & z_1 \\ x_2 & 1 & z_2 \\ x_3 & 1 & z_3 \end{vmatrix} \quad C = \begin{vmatrix} x_1 & y_1 & 1 \\ x_2 & y_2 & 1 \\ x_3 & y_3 & 1 \end{vmatrix} \quad D = -\begin{vmatrix} x_1 & y_1 & z_1 \\ x_2 & y_2 & z_2 \\ x_3 & y_3 & z_3 \end{vmatrix}$$

Expanding the determinants, we can write the calculations for the plane coefficients in the form

$$A = y_1(z_2 - z_3) + y_2(z_3 - z_1) + y_3(z_1 - z_2)$$
$$B = z_1(x_2 - x_3) + z_2(x_3 - x_1) + z_3(x_1 - x_2)$$
$$C = x_1(y_2 - y_3) + x_2(y_3 - y_1) + x_3(y_1 - y_2)$$
$$D = -x_1(y_2 z_3 - y_3 z_2) - x_2(y_3 z_1 - y_1 z_3) - x_3(y_1 z_2 - y_2 z_1)$$

These calculations are valid for any three coordinate positions, including those for which \(D = 0.\) When vertex coordinates and other information are entered into the polygon data structure, values for \(A, B, C,\) and \(D\) can be computed for each polygon facet and stored with the other polygon data.

It is possible that the coordinates defining a polygon facet may not be contained within a single plane. We can solve this problem by dividing the facet into a set of triangles; or we could find an approximating plane for the vertex list. One method for obtaining an approximating plane is to divide the vertex list into subsets, where each subset contains three vertices, and calculate plane parameters \(A, B, C,\) and \(D\) for each subset. The approximating plane parameters are then obtained as the average value for each of the calculated plane parameters. Another approach is to project the vertex list onto the coordinate planes. Then we take parameter \(A\) proportional to the area of the polygon projection on the \(yz\) plane, parameter \(B\) proportional to the projection area on the \(xz\) plane, and parameter \(C\) proportional to the projection area on the \(xy\) plane. The projection method is often used in ray-tracing applications.

**Front and Back Polygon Faces**

Because we are usually dealing with polygon surfaces that enclose an object interior, we need to distinguish between the two sides of each surface. The side of a polygon that faces into the object interior is called the **back face**, and the visible, or outward, side is the **front face**. Identifying the position of points in space
relative to the front and back faces of a polygon is a basic task in many graphics algorithms, as, for example, in determining object visibility. Every polygon is contained within an infinite plane that partitions space into two regions. Any point that is not on the plane and that is visible to the front face of a polygon surface section is said to be in front of (or outside) the plane, and, thus, outside the object. And any point that is visible to the back face of the polygon is behind (or inside) the plane. A point that is behind (inside) all polygon surface planes is inside the object. We need to keep in mind that this inside/outside classification is relative to the plane containing the polygon, whereas our previous inside/outside tests using the winding-number or odd-even rule were in reference to the interior of some two-dimensional boundary.

Plane equations can be used to identify the position of spatial points relative to the polygon facets of an object. For any point \((x, y, z)\) not on a plane with parameters \(A, B, C, D\), we have

\[
Ax + By + Cz + D \neq 0
\]

Thus, we can identify the point as either behind or in front of a polygon surface contained within that plane according to the sign (negative or positive) of \(Ax + By + Cz + D\):

- if \(Ax + By + Cz + D < 0\), the point \((x, y, z)\) is behind the plane
- if \(Ax + By + Cz + D > 0\), the point \((x, y, z)\) is in front of the plane

These inequality tests are valid in a right-handed Cartesian system, provided the plane parameters \(A, B, C, D\) were calculated using coordinate positions selected in a strictly counterclockwise order when viewing the surface along a front-to-back direction. For example, in Figure 18, any point outside (in front of) the plane of the shaded polygon satisfies the inequality \(x - 1 > 0\), while any point inside (in back of) the plane has an \(x\)-coordinate value less than 1.

Orientation of a polygon surface in space can be described with the normal vector for the plane containing that polygon, as shown in Figure 19. This surface normal vector is perpendicular to the plane and has Cartesian components \((A, B, C)\), where parameters \(A, B, C\) are the plane coefficients calculated in Equations 4. The normal vector points in a direction from inside the plane to the outside; that is, from the back face of the polygon to the front face.

As an example of calculating the components of the normal vector for a polygon, which also gives us the plane parameters, we choose three of the vertices of the shaded face of the unit cube in Figure 18. These points are selected in a counterclockwise ordering as we view the cube from outside looking toward the origin. Coordinates for these vertices, in the order selected, are then used in Equations 4 to obtain the plane coefficients: \(A = 1, B = 0, C = 0, D = -1\). Thus, the normal vector for this plane is \(N = (1, 0, 0)\), which is in the direction of the positive \(x\) axis. That is, the normal vector is pointing from inside the cube to the outside and is perpendicular to the plane \(x = 1\).

The elements of a normal vector can also be obtained using a vector cross-product calculation. Assuming we have a convex-polygon surface facet and a right-handed Cartesian system, we again select any three vertex positions, \(V_1, V_2,\) and \(V_3\), taken in counterclockwise order when viewing from outside the object toward the inside. Forming two vectors, one from \(V_1\) to \(V_2\) and the second from \(V_1\) to \(V_3\), we calculate \(N\) as the vector cross-product:

\[
N = (V_2 - V_1) \times (V_3 - V_1)
\]

This generates values for the plane parameters \(A, B, C\). We can then obtain the value for parameter \(D\) by substituting these values and the coordinates for one of
the polygon vertices into Equation 1 and solving for $D$. The plane equation can be expressed in vector form using the normal $\mathbf{N}$ and the position $\mathbf{P}$ of any point in the plane as

$$\mathbf{N} \cdot \mathbf{P} = -D$$

For a convex polygon, we could also obtain the plane parameters using the cross-product of two successive edge vectors. And with a concave polygon, we can select the three vertices so that the two vectors for the cross-product form an angle less than $180^\circ$. Otherwise, we can take the negative of their cross-product to get the correct normal vector direction for the polygon surface.

### 8 OpenGL Polygon Fill-Area Functions

With one exception, the OpenGL procedures for specifying fill polygons are similar to those for describing a point or a polyline. A `glVertex` function is used to input the coordinates for a single polygon vertex, and a complete polygon is described with a list of vertices placed between a `glBegin/glEnd` pair. However, there is one additional function that we can use for displaying a rectangle that has an entirely different format.

By default, a polygon interior is displayed in a solid color, determined by the current color settings. As options, we can fill a polygon with a pattern and we can display polygon edges as line borders around the interior fill. There are six different symbolic constants that we can use as the argument in the `glBegin` function to describe polygon fill areas. These six primitive constants allow us to display a single fill polygon, a set of unconnected fill polygons, or a set of connected fill polygons.

In OpenGL, a fill area must be specified as a convex polygon. Thus, a vertex list for a fill polygon must contain at least three vertices, there can be no crossing edges, and all interior angles for the polygon must be less than $180^\circ$. And a single polygon fill area can be defined with only one vertex list, which precludes any specifications that contain holes in the polygon interior, such as that shown in Figure 20. We could describe such a figure using two overlapping convex polygons.

Each polygon that we specify has two faces: a back face and a front face. In OpenGL, fill color and other attributes can be set for each face separately, and back/front identification is needed in both two-dimensional and three-dimensional viewing routines. Therefore, polygon vertices should be specified in a counterclockwise order as we view the polygon from “outside.” This identifies the front face of that polygon.

![Figure 20](image-url)
Because graphics displays often include rectangular fill areas, OpenGL provides a special rectangle function that directly accepts vertex specifications in the $xy$ plane. In some implementations of OpenGL, the following routine can be more efficient than generating a fill rectangle using `glVertex` specifications:

```c
glRect* (x1, y1, x2, y2);
```

One corner of this rectangle is at coordinate position $(x_1, y_1)$, and the opposite corner of the rectangle is at position $(x_2, y_2)$. Suffix codes for `glRect` specify the coordinate data type and whether coordinates are to be expressed as array elements. These codes are $i$ (for integer), $s$ (for short), $f$ (for float), $d$ (for double), and $v$ (for vector). The rectangle is displayed with edges parallel to the $xy$ coordinate axes. As an example, the following statement defines the square shown in Figure 21:

```c
glRecti (200, 100, 50, 250);
```

If we put the coordinate values for this rectangle into arrays, we can generate the same square with the following code:

```c
int vertex1 [ ] = {200, 100};
int vertex2 [ ] = {50, 250};

glRectiv (vertex1, vertex2);
```

When a rectangle is generated with function `glRect`, the polygon edges are formed between the vertices in the order $(x_1, y_1)$, $(x_2, y_1)$, $(x_2, y_2)$, $(x_1, y_2)$, and then back to $(x_1, y_1)$. Thus, in our example, we produced a vertex list with a clockwise ordering. In many two-dimensional applications, the determination of front and back faces is unimportant. But if we do want to assign different properties to the front and back faces of the rectangle, then we should reverse the order of the two vertices in this example so that we obtain a counterclockwise ordering of the vertices.

Each of the other six OpenGL polygon fill primitives is specified with a symbolic constant in the `glBegin` function, along with a list of `glVertex` commands. With the OpenGL primitive constant `GL_POLYGON`, we can display a single polygon fill area such as that shown in Figure 22(a). For this example, we assume that we have a list of six points, labeled $p_1$ through $p_6$, specifying...
two-dimensional polygon vertex positions in a counterclockwise ordering. Each of the points is represented as an array of \((x, y)\) coordinate values:

```c
glBegin (GL_POLYGON);
glVertex2iv (p1);
glVertex2iv (p2);
glVertex2iv (p3);
glVertex2iv (p4);
glVertex2iv (p5);
glVertex2iv (p6);
glEnd ( );
```

A polygon vertex list must contain at least three vertices. Otherwise, nothing is displayed.

If we reorder the vertex list and change the primitive constant in the previous code example to GL_TRIANGLES, we obtain the two separated triangle fill areas in Figure 22(b):

```c
glBegin (GL_TRIANGLES);
glVertex2iv (p1);
glVertex2iv (p2);
glVertex2iv (p6);
glVertex2iv (p3);
glVertex2iv (p4);
glVertex2iv (p5);
glEnd ( );
```
In this case, the first three coordinate points define the vertices for one triangle, the next three points define the next triangle, and so forth. For each triangle fill area, we specify the vertex positions in a counterclockwise order. A set of unconnected triangles is displayed with this primitive constant unless some vertex coordinates are repeated. Nothing is displayed if we do not list at least three vertices; and if the number of vertices specified is not a multiple of 3, the final one or two vertex positions are not used.

By reordering the vertex list once more and changing the primitive constant to \texttt{GL\_TRIANGLE\_STRIP}, we can display the set of connected triangles shown in Figure 22(c):

```c
glBegin (GL\_TRIANGLE\_STRIP);
glVertex2iv (p1);
glVertex2iv (p2);
glVertex2iv (p6);
glVertex2iv (p3);
glVertex2iv (p5);
glVertex2iv (p4);
glEnd ( );
```

Assuming that no coordinate positions are repeated in a list of \(N\) vertices, we obtain \(N - 2\) triangles in the strip. Clearly, we must have \(N \geq 3\) or nothing is displayed. In this example, \(N = 6\) and we obtain four triangles. Each successive triangle shares an edge with the previously defined triangle, so the ordering of the vertex list must be set up to ensure a consistent display. One triangle is defined for each vertex position listed after the first two vertices. Thus, the first three vertices should be listed in counterclockwise order, when viewing the front (outside) surface of the triangle. After that, the set of three vertices for each subsequent triangle is arranged in a counterclockwise order within the polygon tables. This is accomplished by processing each position \(n\) in the vertex list in the order \(n = 1, n = 2, \ldots, n = N - 2\) and arranging the order of the corresponding set of three vertices according to whether \(n\) is an odd number or an even number. If \(n\) is odd, the polygon table listing for the triangle vertices is in the order \(n, n + 1, n + 2\). If \(n\) is even, the triangle vertices are listed in the order \(n + 1, n, n + 2\). In the preceding example, our first triangle \((n = 1)\) would be listed as having vertices \((p1, p2, p6)\). The second triangle \((n = 2)\) would have the vertex ordering \((p6, p2, p3)\). Vertex ordering for the third triangle \((n = 3)\) would be \((p6, p3, p5)\). And the fourth triangle \((n = 4)\) would be listed in the polygon tables with vertex ordering \((p5, p3, p4)\).

Another way to generate a set of connected triangles is to use the “fan” approach illustrated in Figure 22(d), where all triangles share a common vertex. We obtain this arrangement of triangles using the primitive constant \texttt{GL\_TRIANGLE\_FAN} and the original ordering of our six vertices:

```c
glBegin (GL\_TRIANGLE\_FAN);
glVertex2iv (p1);
glVertex2iv (p2);
glVertex2iv (p3);
glVertex2iv (p4);
glVertex2iv (p5);
glVertex2iv (p6);
glEnd ( );
```

For \(N\) vertices, we again obtain \(N - 2\) triangles, providing no vertex positions are repeated, and we must list at least three vertices. In addition, the vertices must
be specified in the proper order to define front and back faces for each triangle correctly. The first coordinate position listed (in this case, p1) is a vertex for each triangle in the fan. If we again enumerate the triangles and the coordinate positions listed as $n = 1, n = 2, \ldots, n = N - 2$, then vertices for triangle $n$ are listed in the polygon tables in the order $1, n + 1, n + 2$. Therefore, triangle 1 is defined with the vertex list $(p1, p2, p3)$; triangle 2 has the vertex ordering $(p1, p3, p4)$; triangle 3 has its vertices specified in the order $(p1, p4, p5)$; and triangle 4 is listed with vertices $(p1, p5, p6)$.

Besides the primitive functions for triangles and a general polygon, OpenGL provides for the specifications of two types of quadrilaterals (four-sided polygons). With the GL_QUADS primitive constant and the following list of eight vertices, specified as two-dimensional coordinate arrays, we can generate the display shown in Figure 23(a):

```c
glBegin (GL_QUADS);
    glVertex2iv (p1);
    glVertex2iv (p2);
    glVertex2iv (p3);
    glVertex2iv (p4);
    glVertex2iv (p5);
    glVertex2iv (p6);
    glVertex2iv (p7);
    glVertex2iv (p8);
glEnd ( );
```

![Figure 23](image-url)
The first four coordinate points define the vertices for one quadrilateral, the next four points define the next quadrilateral, and so on. For each quadrilateral fill area, we specify the vertex positions in a counterclockwise order. If no vertex coordinates are repeated, we display a set of unconnected four-sided fill areas. We must list at least four vertices with this primitive. Otherwise, nothing is displayed. And if the number of vertices specified is not a multiple of 4, the extra vertex positions are ignored.

Rearranging the vertex list in the previous quadrilateral code example and changing the primitive constant to \texttt{GL\_QUAD\_STRIP}, we can obtain the set of connected quadrilaterals shown in Figure 23(b):

```c
glBegin (GL\_QUAD\_STRIP);
glVertex2iv (p1);
glVertex2iv (p2);
glVertex2iv (p4);
glVertex2iv (p3);
glVertex2iv (p5);
glVertex2iv (p6);
glVertex2iv (p8);
glVertex2iv (p7);
glEnd ( );
```

A quadrilateral is set up for each pair of vertices specified after the first two vertices in the list, and we need to list the vertices so that we generate a correct counterclockwise vertex ordering for each polygon. For a list of \( N \) vertices, we obtain \( \frac{N}{2} - 1 \) quadrilaterals, providing that \( N \geq 4 \). If \( N \) is not a multiple of 4, any extra coordinate positions in the list are not used. We can enumerate these fill polygons and the vertices listed as \( n = 1, n = 2, \ldots, n = \frac{N}{2} - 1 \). Then polygon tables will list the vertices for quadrilateral \( n \) in the vertex order number \( 2n - 1, 2n, 2n + 2, 2n + 1 \). For this example, \( N = 8 \) and we have 3 quadrilaterals in the strip. Thus, our first quadrilateral \((n = 1)\) is listed as having a vertex ordering of \((p1, p2, p3, p4)\). The second quadrilateral \((n = 2)\) has the vertex ordering \((p4, p3, p6, p5)\), and the vertex ordering for the third quadrilateral \((n = 3)\) is \((p5, p6, p7, p8)\).

Most graphics packages display curved surfaces as a set of approximating plane facets. This is because plane equations are linear, and processing the linear equations is much quicker than processing quadric or other types of curve equations. So OpenGL and other packages provide polygon primitives to facilitate the approximation of a curved surface. Objects are modeled with polygon meshes, and a database of geometric and attribute information is set up to facilitate the processing of the polygon facets. In OpenGL, primitives that we can use for this purpose are the \texttt{triangle strip}, the \texttt{triangle fan}, and the \texttt{quad strip}. Fast hardware-implemented polygon renderers are incorporated into high-quality graphics systems with the capability for displaying millions of shaded polygons per second (usually triangles), including the application of surface texture and special lighting effects.

Although the OpenGL core library allows only convex polygons, the GLU library provides functions for dealing with concave polygons and other nonconvex objects with linear boundaries. A set of GLU \texttt{polygon tessellation} routines is available for converting such shapes into a set of triangles, triangle meshes, triangle fans, and straight-line segments. Once such objects have been decomposed, they can be processed with basic OpenGL functions.
## 9 OpenGL Vertex Arrays

Although our examples so far have contained relatively few coordinate positions, describing a scene containing several objects can get much more complicated. To illustrate, we first consider describing a single, very basic object: the unit cube shown in Figure 24, with coordinates given in integers to simplify our discussion. A straightforward method for defining the vertex coordinates is to use a double-subscripted array, such as

```c
GLint points [8][3] = { {0, 0, 0}, {0, 1, 0}, {1, 0, 0}, {1, 1, 0},
                      {0, 0, 1}, {0, 1, 1}, {1, 0, 1}, {1, 1, 1} };
```

Alternatively, we could first define a data type for a three-dimensional vertex position and then give the coordinates for each vertex position as an element of a single-subscripted array as, for example,

```c
typedef GLint vertex3 [3];
vertex3 pt [8] = { {0, 0, 0}, {0, 1, 0}, {1, 0, 0}, {1, 1, 0},
                 {0, 0, 1}, {0, 1, 1}, {1, 0, 1}, {1, 1, 1} };  
```

Next, we need to define each of the six faces of this object. For this, we could make six calls either to `glBegin (GL_POLYGON)` or to `glBegin (GL_QUADS)`. In either case, we must be sure to list the vertices for each face in a counterclockwise order when viewing that surface from the outside of the cube. In the following code segment, we specify each cube face as a quadrilateral and use a function call to pass array subscript values to the OpenGL primitive routines. Figure 25 shows the subscript values for array `pt` corresponding to the cube vertex positions.

![Figure 24](image1.png)  
**Figure 24**  
A cube with an edge length of 1.

![Figure 25](image2.png)  
**Figure 25**  
Subscript values for array `pt` corresponding to the vertex coordinates for the cube shown in Figure 24.
void quad (GLint n1, GLint n2, GLint n3, GLint n4)
{
    glBegin (GL_QUADS);
    glVertex3iv (pt [n1]);
    glVertex3iv (pt [n2]);
    glVertex3iv (pt [n3]);
    glVertex3iv (pt [n4]);
    glEnd ( );
}

void cube ( )
{
    quad (6, 2, 3, 7);
    quad (5, 1, 0, 4);
    quad (7, 3, 1, 5);
    quad (4, 0, 2, 6);
    quad (2, 0, 1, 3);
    quad (7, 5, 4, 6);
}

Thus, the specification for each face requires six OpenGL functions, and we have six faces to specify. When we add color specifications and other parameters, our display program for the cube could easily contain 100 or more OpenGL function calls. And scenes with many complex objects can require much more.

As we can see from the preceding cube example, a complete scene description could require hundreds or thousands of coordinate specifications. In addition, there are various attribute and viewing parameters that must be set for individual objects. Thus, object and scene descriptions could require an enormous number of function calls, which puts a demand on system resources and can slow execution of the graphics programs. A further problem with complex displays is that object surfaces (such as the cube in Figure 24) usually have shared vertex coordinates. Using the methods we have discussed up to now, these shared positions may need to be specified multiple times.

To alleviate these problems, OpenGL provides a mechanism for reducing the number of function calls needed in processing coordinate information. Using a vertex array, we can arrange the information for describing a scene so that we need only a very few function calls. The steps involved are

1. Invoke the function glEnableClientState (GL_VERTEX_ARRAY) to activate the vertex-array feature of OpenGL.
2. Use the function glVertexPointer to specify the location and data format for the vertex coordinates.
3. Display the scene using a routine such as glDrawElements, which can process multiple primitives with very few function calls.

Using the pt array previously defined for the cube, we implement these three steps in the following code example:

    glEnableClientState (GL_VERTEX_ARRAY);
    glVertexPointer (3, GL_INT, 0, pt);
    
    GLubyte vertIndex [] = (6, 2, 3, 7, 5, 1, 0, 4, 7, 3, 1, 5,
                             4, 0, 2, 6, 2, 0, 1, 3, 7, 5, 4, 6);
    
    glDrawElements (GL_QUADS, 24, GL_UNSIGNED_BYTE, vertIndex);

Graphics Output Primitives
With the first command, `glEnableClientState (GL_VERTEX_ARRAY)`, we activate a capability (in this case, a vertex array) on the client side of a client-server system. Because the client (the machine that is running the main program) retains the data for a picture, the vertex array must be there also. The server (our workstation, for example) generates commands and displays the picture. Of course, a single machine can be both client and server. The vertex-array feature of OpenGL is deactivated with the command

```
glDisableClientState (GL_VERTEX_ARRAY);
```

We next give the location and format of the coordinates for the object vertices in the function `glVertexPointer`. The first parameter in `glVertexPointer` (3 in this example) specifies the number of coordinates used in each vertex description. Data type for the vertex coordinates is designated using an OpenGL symbolic constant as the second parameter in this function. For our example, the data type is `GL_INT`. Other data types are specified with the symbolic constants `GL_BYTE`, `GL_SHORT`, `GL_FLOAT`, and `GL_DOUBLE`. With the third parameter, we give the byte offset between consecutive vertices. The purpose of this argument is to allow various kinds of data, such as coordinates and colors, to be packed together in one array. Because we are giving only the coordinate data, we assign a value of 0 to the offset parameter. The last parameter in the `glVertexPointer` function references the vertex array, which contains the coordinate values.

All the indices for the cube vertices are stored in array `vertIndex`. Each of these indices is the subscript for array `pt` corresponding to the coordinate values for that vertex. This index list is referenced as the last parameter value in function `glDrawElements` and is then used by the primitive `GL_QUADS`, which is the first parameter, to display the set of quadrilateral surfaces for the cube. The second parameter specifies the number of elements in array `vertIndex`. Because a quadrilateral requires just 4 vertices and we specified 24, the `glDrawElements` function continues to display another cube face after each successive set of 4 vertices until all 24 have been processed. Thus, we accomplish the final display of all faces of the cube with this single function call. The third parameter in function `glDrawElements` gives the type for the index values. Because our indices are small integers, we specified a type of `GL_UNSIGNED_BYTE`. The two other index types that can be used are `GL_UNSIGNED_SHORT` and `GL_UNSIGNED_INT`.

Additional information can be combined with the coordinate values in the vertex arrays to facilitate the processing of a scene description. We can specify color values and other attributes for objects in arrays that can be referenced by the `glDrawElements` function. Also, we can interlace the various arrays for greater efficiency.

### 10 Pixel-Array Primitives

In addition to straight lines, polygons, circles, and other primitives, graphics packages often supply routines to display shapes that are defined with a rectangular array of color values. We can obtain the rectangular grid pattern by digitizing (scanning) a photograph or other picture or by generating a shape with a graphics program. Each color value in the array is then mapped to one or more screen pixel positions. A pixel array of color values is typically referred to as a *pixmap*. 
Parameters for a pixel array can include a pointer to the color matrix, the size of the matrix, and the position and size of the screen area to be affected by the color values. Figure 26 gives an example of mapping a pixel-color array onto a screen area.

Another method for implementing a pixel array is to assign either the bit value 0 or the bit value 1 to each element of the matrix. In this case, the array is simply a bitmap, which is sometimes called a mask, that indicates whether a pixel is to be assigned (or combined with) a preset color.

11 OpenGL Pixel-Array Functions

There are two functions in OpenGL that we can use to define a shape or pattern specified with a rectangular array. One function defines a bitmap pattern, and the other a pixmap pattern. Also, OpenGL provides several routines for saving, copying, and manipulating arrays of pixel values.

OpenGL Bitmap Function

A binary array pattern is defined with the function

\[
glBitmap (\text{width}, \text{height}, x_0, y_0, xoffset, yoffset, \text{bitShape});
\]

Parameters width and height in this function give the number of columns and number of rows, respectively, in the array bitShape. Each element of bitShape is assigned either a 1 or a 0. A value of 1 indicates that the corresponding pixel is to be displayed in a previously defined color. Otherwise, the pixel is unaffected by the bitmap. (As an option, we could use a value of 1 to indicate that a specified color is to be combined with the color value stored in the refresh buffer at that position.) Parameters \(x_0\) and \(y_0\) define the position that is to be considered the "origin" of the rectangular array. This origin position is specified relative to the lower-left corner of bitShape, and values for \(x_0\) and \(y_0\) can be positive or negative. In addition, we need to designate a location in the frame buffer where the pattern is to be applied. This location is called the current raster position, and the bitmap is displayed by positioning its origin, \((x_0, y_0)\), at the current position.
raster position. Values assigned to parameters \texttt{xOffset} and \texttt{yOffset} are used as coordinate offsets to update the frame-buffer current raster position after the bitmap is displayed.

Coordinate values for \texttt{x0}, \texttt{y0}, \texttt{xOffset}, and \texttt{yOffset}, as well as the current raster position, are maintained as floating-point values. Of course, bitmaps will be applied at integer pixel positions. But floating-point coordinates allow a set of bitmaps to be spaced at arbitrary intervals, which is useful in some applications, such as forming character strings with bitmap patterns.

We use the following routine to set the coordinates for the current raster position:

\begin{verbatim}
glRasterPos* ( )
\end{verbatim}

Parameters and suffix codes are the same as those for the \texttt{glVertex} function. Thus, a current raster position is given in world coordinates, and it is transformed to screen coordinates by the viewing transformations. For our two-dimensional examples, we can specify coordinates for the current raster position directly in integer screen coordinates. The default value for the current raster position is the world-coordinate origin (0, 0, 0).

The color for a bitmap is the color that is in effect at the time that the \texttt{glRasterPos} command is invoked. Any subsequent color changes do not affect the bitmap.

Each row of a rectangular bit array is stored in multiples of 8 bits, where the binary data is arranged as a set of 8-bit unsigned characters. But we can describe a shape using any convenient grid size. For example, Figure 27 shows a bit pattern defined on a 10-row by 9-column grid, where the binary data is specified with 16 bits for each row. When this pattern is applied to the pixels in the frame buffer, all bit values beyond the ninth column are ignored.

We apply the bit pattern of Figure 27 to a frame-buffer location with the following code section:

\begin{verbatim}
GLubyte bitShape [20] = {
    0x1c, 0x00, 0x1c, 0x00, 0x1c, 0x00, 0x1c, 0x00, 0x1c, 0x00,
    0xff, 0x80, 0x7f, 0x00, 0x3e, 0x00, 0x1c, 0x00, 0x08, 0x00};

glPixelStorei (GL_UNPACK_ALIGNMENT, 1); // Set pixel storage mode.

glRasterPos2i (30, 40);

glBitmap (9, 10, 0.0, 0.0, 20.0, 15.0, bitShape);
\end{verbatim}

\textbf{Figure 27}
A bit pattern, specified in an array with 10 rows and 9 columns, is stored in 8-bit blocks of 10 rows with 16 bit values per row.
Array values for `bitShape` are specified row by row, starting at the bottom of the rectangular-grid pattern. Next we set the storage mode for the bitmap with the OpenGL routine `glPixelStorei`. The parameter value of 1 in this function indicates that the data values are to be aligned on byte boundaries. With `glRasterPos`, we set the current raster position to (30, 40). Finally, function `glBitmap` specifies that the bit pattern is given in array `bitShape`, and that this array has 9 columns and 10 rows. The coordinates for the origin of this pattern are (0.0, 0.0), which is the lower-left corner of the grid. We illustrate a coordinate offset with the values (20.0, 15.0), although we do not use the offset in this example.

### OpenGL Pixmap Function

A pattern defined as an array of color values is applied to a block of frame-buffer pixel positions with the function

\[
\text{glDrawPixels} \ (\text{width}, \text{height}, \text{dataFormat}, \text{dataType}, \text{pixMap});
\]

Again, parameters `width` and `height` give the column and row dimensions, respectively, of the array `pixMap`. Parameter `dataFormat` is assigned an OpenGL constant that indicates how the values are specified for the array. For example, we could specify a single blue color for all pixels with the constant `GL_BLUE`, or we could specify three color components in the order blue, green, red with the constant `GL_BGR`. A number of other color specifications are possible. An OpenGL constant, such as `GL_BYTE`, `GL_INT`, or `GL_FLOAT`, is assigned to parameter `dataType` to designate the data type for the color values in the array. The lower-left corner of this color array is mapped to the current raster position, as set by the `glRasterPos` function. As an example, the following statement displays a pixmap defined in a 128 \times 128 array of RGB color values:

\[
\text{glDrawPixels} \ (128, \ 128, \ \text{GL_RGB}, \ \text{GL_UNSIGNED_BYTE}, \ \text{colorShape});
\]

Because OpenGL provides several buffers, we can paste an array of values into a particular buffer by selecting that buffer as the target of the `glDrawPixels` routine. Some buffers store color values and some store other kinds of pixel data. A depth buffer, for instance, is used to store object distances (depths) from the viewing position, and a stencil buffer is used to store boundary patterns for a scene. We select one of these two buffers by setting parameter `dataFormat` in the `glDrawPixels` routine to either `GL_DEPTH_COMPONENT` or `GL_STENCIL_INDEX`. For these buffers, we would need to set up the pixel array using either depth values or stencil information.

There are four color buffers available in OpenGL that can be used for screen refreshing. Two of the color buffers constitute a left-right scene pair for displaying stereoscopic views. For each of the stereoscopic buffers, there is a front-back pair for double-buffered animation displays. In a particular implementation of OpenGL, either stereoscopic viewing or double buffering, or both, might not be supported. If neither stereoscopic effects nor double buffering is supported, then there is only a single refresh buffer, which is designated as the front-left color buffer. This is the default refresh buffer when double buffering is not available or not in effect. If double buffering is in effect, the default is either the back-left and back-right buffers or only the back-left buffer, depending on the current state of stereoscopic viewing. Also, a number of user-defined, auxiliary color buffers are supported that can be used for any nonrefresh purpose, such as saving a picture that is to be copied later into a refresh buffer for display.
We select a single color or auxiliary buffer or a combination of color buffers for storing a pixmap with the following command:

```
glDrawBuffer (buffer);
```

A variety of OpenGL symbolic constants can be assigned to parameter `buffer` to designate one or more “draw” buffers. For instance, we can pick a single buffer with either `GL_FRONT_LEFT`, `GL_FRONT_RIGHT`, `GL_BACK_LEFT`, or `GL_BACK_RIGHT`. We can select both front buffers with `GL_FRONT`, and we can select both back buffers with `GL_BACK`. This is assuming that stereoscopic viewing is in effect. Otherwise, the previous two symbolic constants designate a single buffer. Similarly, we can designate either the left or right buffer pairs with `GL_LEFT` or `GL_RIGHT`, and we can select all the available color buffers with `GL_FRONT_AND_BACK`. An auxiliary buffer is chosen with the constant `GL_AUXk`, where `k` is an integer value from 0 to 3, although more than four auxiliary buffers may be available in some implementations of OpenGL.

**OpenGL Raster Operations**

In addition to storing an array of pixel values in a buffer, we can retrieve a block of values from a buffer or copy the block into another buffer area, and we can perform a variety of other operations on a pixel array. In general, the term **raster operation** or **raster op** is used to describe any function that processes a pixel array in some way. A raster operation that moves an array of pixel values from one place to another is also referred to as a **block transfer** of pixel values. On a bilevel system, these operations are called **bitblt transfers** (**bit-block transfers**), particularly when the functions are hardware-implemented. On a multilevel system, the term **pixblt** is sometimes used for block transfers.

We use the following function to select a rectangular block of pixel values in a designated set of buffers:

```
glReadPixels (xmin, ymin, width, height, dataFormat, dataType, array);
```

The lower-left corner of the rectangular block to be retrieved is at screen-coordinate position `(xmin, ymin)`. Parameters `width`, `height`, `dataFormat`, and `dataType` are the same as in the `glDrawPixels` routine. The type of data to be saved in parameter `array` depends on the selected buffer. We can choose either the depth buffer or the stencil buffer by assigning either the value `GL_DEPTH_COMPONENT` or the value `GL_STENCIL_INDEX` to parameter `dataFormat`.

A particular combination of color buffers or an auxiliary buffer is selected for the application of the `glReadPixels` routine with the function

```
glReadBuffer (buffer);
```

Symbolic constants for specifying one or more buffers are the same as in the `glDrawBuffer` routine except that we cannot select all four of the color buffers. The default buffer selection is the front left-right pair or just the front-left buffer, depending on the status of stereoscopic viewing.

We can also copy a block of pixel data from one location to another within the set of OpenGL buffers using the following routine:

```
glCopyPixels (xmin, ymin, width, height, pixelValues);
```
The lower-left corner of the block is at screen-coordinate location \((x_{\text{min}}, y_{\text{min}})\), and parameters \(width\) and \(height\) are assigned positive integer values to designate the number of columns and rows, respectively, that are to be copied. Parameter \(pixelValues\) is assigned either \(\text{GL}_{\text{COLOR}}\), \(\text{GL}_{\text{DEPTH}}\), or \(\text{GL}_{\text{STENCIL}}\) to indicate the kind of data we want to copy: color values, depth values, or stencil values. In addition, the block of pixel values is copied from a \(source\ \text{buffer}\) to a \(destination\ \text{buffer}\), with its lower-left corner mapped to the current raster position. We select the source buffer with the \text{glReadBuffer} command, and we select the destination buffer with the \text{glDrawBuffer} command. Both the region to be copied and the destination area should lie completely within the bounds of the screen coordinates.

To achieve different effects as a block of pixel values is placed into a buffer with \text{glDrawPixels} or \text{glCopyPixels}, we can combine the incoming values with the old buffer values in various ways. As an example, we could apply logical operations, such as \(\text{and}\), \(\text{or}\), and \(\text{exclusive or}\), to combine the two blocks of pixel values. In OpenGL, we select a bitwise, logical operation for combining incoming and destination pixel color values with the functions

\[
glEnable (\text{GL}_{\text{COLOR} \_\text{LOGIC\_OP}}); \\
glLogicOp (\text{logicOp});
\]

A variety of symbolic constants can be assigned to parameter \(\text{logicOp}\), including \(\text{GL} \_\text{AND}\), \(\text{GL} \_\text{OR}\), and \(\text{GL} \_\text{XOR}\). In addition, either the incoming bit values or the destination bit values can be inverted (interchanging 0 and 1 values). We use the constant \(\text{GL} \_\text{COPY\_INVERTED}\) to invert the incoming color bit values and then replace the destination values with the inverted incoming values; and we could simply invert the destination bit values without replacing them with the incoming values using \(\text{GL} \_\text{INVERT}\). The various invert operations can also be combined with the logical \(\text{and}\), \(\text{or}\), and \(\text{exclusive or}\) operations. Other options include clearing all the destination bits to the value 0 (\(\text{GL} \_\text{CLEAR}\)), or setting all the destination bits to the value 1 (\(\text{GL} \_\text{SET}\)). The default value for the \text{glLogicOp} routine is \(\text{GL} \_\text{COPY}\), which simply replaces the destination values with the incoming values.

Additional OpenGL routines are available for manipulating pixel arrays processed by the \text{glDrawPixels}, \text{glReadPixels}, and \text{glCopyPixels} functions. For example, the \text{glPixelTransfer} and \text{glPixelMap} routines can be used to shift or adjust color values, depth values, or stencil values. We return to pixel operations in later chapters as we explore other facets of computer-graphics packages.

### 12 Character Primitives

Graphics displays often include textural information, such as labels on graphs and charts, signs on buildings or vehicles, and general identifying information for simulation and visualization applications. Routines for generating character primitives are available in most graphics packages. Some systems provide an extensive set of character functions, while other systems offer only minimal support for character generation.

Letters, numbers, and other characters can be displayed in a variety of sizes and styles. The overall design style for a set (or family) of characters is called a \textit{typeface}. Today, there are thousands of typefaces available for computer applications. Examples of a few common typefaces are Courier, Helvetica, New York,
Palatino, and Zapf Chancery. Originally, the term **font** referred to a set of cast metal character forms in a particular size and format, such as 10-point Courier Italic or 12-point Palatino Bold. A 14-point font has a total character height of about 0.5 centimeter. In other words, 72 points is about the equivalent of 2.54 centimeters (1 inch). The terms **font** and **typeface** are now often used interchangeably, since most printing is no longer done with cast metal forms.

Fonts can be divided into two broad groups: **serif** and **sans serif**. Serif type has small lines or accents at the ends of the main character strokes, while sans-serif type does not have such accents. For example, this text is set in a serif font (Palatino). But this sentence is printed in a sans-serif font (Univers). Serif type is generally more **readable**; that is, it is easier to read in longer blocks of text. On the other hand, the individual characters in sans-serif type are easier to recognize. For this reason, sans-serif type is said to be more **legible**. Since sans-serif characters can be recognized quickly, this font is good for labeling and short headings.

Fonts are also classified according to whether they are **monospace** or **proportional**. Characters in a monospace font all have the same width. In a proportional font, character width varies.

Two different representations are used for storing computer fonts. A simple method for representing the character shapes in a particular typeface is to set up a pattern of binary values on a rectangular grid. The set of characters is then referred to as a **bitmap font** (or **bitmapped font**). A bitmapped character set is also sometimes referred to as a **raster font**. Another, more flexible, scheme is to describe character shapes using straight-line and curve sections, as in PostScript, for example. In this case, the set of characters is called an **outline font** or a **stroke font**. Figure 28 illustrates the two methods for character representation. When the pattern in Figure 28(a) is applied to an area of the frame buffer, the 1 bits designate which pixel positions are to be displayed in a specified color. To display the character shape in Figure 28(b), the interior of the character outline is treated as a fill area.

Bitmap fonts are the simplest to define and display: We just need to map the character grids to a frame-buffer position. In general, however, bitmap fonts require more storage space because each variation (size and format) must be saved in a **font cache**. It is possible to generate different sizes and other variations, such as bold and italic, from one bitmap font set, but this often does not produce good results. We can increase or decrease the size of a character bitmap only in integer multiples of the pixel size. To double the size of a character, we need to double the number of pixels in the bitmap. This just increases the ragged appearance of its edges.

In contrast to bitmap fonts, outline fonts can be increased in size without distorting the character shapes. And outline fonts require less storage because each variation does not require a distinct font cache. We can produce boldface,
italic, or different sizes by manipulating the curve definitions for the character outlines. But it does take more time to process the outline fonts because they must be scan-converted into the frame buffer.

There is a variety of possible functions for implementing character displays. Some graphics packages provide a function that accepts any character string and a frame-buffer starting position for the string. Another type of function displays a single character at one or more selected positions. Since this character routine is useful for showing markers in a network layout or in displaying a point plot of a discrete data set, the character displayed by this routine is sometimes referred to as a marker symbol or polymarker, in analogy with a polyline primitive. In addition to standard characters, special shapes such as dots, circles, and crosses are often available as marker symbols. Figure 29 shows a plot of a discrete data set using an asterisk as a marker symbol.

Geometric descriptions for characters are given in world coordinates, just as they are for other primitives, and this information is mapped to screen coordinates by the viewing transformations. A bitmap character is described with a rectangular grid of binary values and a grid reference position. This reference position is then mapped to a specified location in the frame buffer. An outline character is defined by a set of coordinate positions that are to be connected with a series of curves and straight-line segments and a reference position that is to be mapped to a given frame-buffer location. The reference position can be specified either for a single outline character or for a string of characters. In general, character routines can allow the construction of both two-dimensional and three-dimensional character displays.

13 OpenGL Character Functions

Only low-level support is provided by the basic OpenGL library for displaying individual characters and text strings. We can explicitly define any character as a bitmap, as in the example shape shown in Figure 27, and we can store a set of bitmap characters as a font list. A text string is then displayed by mapping a selected sequence of bitmaps from the font list into adjacent positions in the frame buffer.

However, some predefined character sets are available in the GLUT library, so we do not need to create our own fonts as bitmap shapes unless we want to display a font that is not available in GLUT. The GLUT library contains routines for displaying both bitmapped and outline fonts. Bitmapped GLUT fonts are rendered using the OpenGL glBitmap function, and the outline fonts are generated with polyline (GL_LINE_STRIP) boundaries.

We can display a bitmap GLUT character with

```c
glutBitmapCharacter (font, character);
```
where parameter \texttt{font} is assigned a symbolic GLUT constant identifying a particular set of typefaces, and parameter \texttt{character} is assigned either the ASCII code or the specific character we wish to display. Thus, to display the uppercase letter “A,” we can either use the ASCII value 65 or the designation ‘A’. Similarly, a code value of 66 is equivalent to ‘B’, code 97 corresponds to the lowercase letter ‘a’, code 98 corresponds to ‘b’, and so forth. Both fixed-width fonts and proportionally spaced fonts are available. We can select a fixed-width font by assigning either \texttt{GLUT_BITMAP\_8\_BY\_13} or \texttt{GLUT_BITMAP\_9\_BY\_15} to parameter \texttt{font}. And we can select a 10-point, proportionally spaced font with either \texttt{GLUT_BITMAP\_TIMES\_ROMAN\_10} or \texttt{GLUT_BITMAP\_HELVETICA\_10}. A 12-point Times-Roman font is also available, as well as 12-point and 18-point Helvetica fonts.

Each character generated by \texttt{glutBitmapCharacter} is displayed so that the origin (lower-left corner) of the bitmap is at the current raster position. After the character bitmap is loaded into the refresh buffer, an offset equal to the width of the character is added to the \texttt{x} coordinate for the current raster position. As an example, we could display a text string containing 36 bitmap characters with the following code:

```c
  glRasterPosition2i (x, y);
  for (k = 0; k < 36; k++)
    glutBitmapCharacter (GLUT_BITMAP\_9\_BY\_15, text [k]);
```

Characters are displayed in the color that was specified before the execution of the \texttt{glutBitmapCharacter} function.

An outline character is displayed with the following function call:

```c
  glutStrokeCharacter (font, character);
```

For this function, we can assign parameter \texttt{font} either the value \texttt{GLUT\_STROKE\_ROMAN}, which displays a proportionally spaced font, or the value \texttt{GLUT\_STROKE\_MONO\_ROMAN}, which displays a font with constant spacing. We control the size and position of these characters by specifying transformation operations before executing the \texttt{glutStrokeCharacter} routine. After each character is displayed, a coordinate offset is applied automatically so that the position for displaying the next character is to the right of the current character. Text strings generated with outline fonts are part of the geometric description for a twodimensional or three-dimensional scene because they are constructed with line segments. Thus, they can be viewed from various directions, and we can shrink or expand them without distortion, or transform them in other ways. But they are slower to render, compared to bitmapped fonts.

---

**14 Picture Partitioning**

Some graphics libraries include routines for describing a picture as a collection of named sections and for manipulating the individual sections of a picture. Using these functions, we can create, edit, delete, or move a part of a picture independently of the other picture components. In addition, we can use this feature of a graphics package for hierarchical modeling, in which an object description is given as a tree structure composed of a number of levels specifying the object subparts.

Various names are used for the subsections of a picture. Some graphics packages refer to them as \texttt{structures}, while other packages call them \texttt{segments}
or objects. Also, the allowable subsection operations vary greatly from one package to another. Modeling packages, for example, provide a wide range of operations that can be used to describe and manipulate picture elements. On the other hand, for any graphics library, we can always structure and manage the components of a picture using procedural elements available in a high-level language such as C++.

15 OpenGL Display Lists

Often it can be convenient or more efficient to store an object description (or any other set of OpenGL commands) as a named sequence of statements. We can do this in OpenGL using a structure called a display list. Once a display list has been created, we can reference the list multiple times with different display operations. On a network, a display list describing a scene is stored on the server machine, which eliminates the need to transmit the commands in the list each time the scene is to be displayed. We can also set up a display list so that it is saved for later execution, or we can specify that the commands in the list be executed immediately. And display lists are particularly useful for hierarchical modeling, where a complex object can be described with a set of simpler subparts.

Creating and Naming an OpenGL Display List

A set of OpenGL commands is formed into a display list by enclosing the commands within the `glNewList/glEndList` pair of functions. For example,

```c
     glNewList (listID, listMode);

     .

     .

     glEndList ( );
```

This structure forms a display list with a positive integer value assigned to parameter `listID` as the name for the list. Parameter `listMode` is assigned an OpenGL symbolic constant that can be either `GL_COMPILE` or `GL_COMPILE_AND_EXECUTE`. If we want to save the list for later execution, we use `GL_COMPILE`. Otherwise, the commands are executed as they are placed into the list, in addition to allowing us to execute the list again at a later time.

As a display list is created, expressions involving parameters such as coordinate positions and color components are evaluated so that only the parameter values are stored in the list. Any subsequent changes to these parameters have no effect on the list. Because display-list values cannot be changed, we cannot include certain OpenGL commands, such as vertex-list pointers, in a display list.

We can create any number of display lists, and we execute a particular list of commands with a call to its identifier. Further, one display list can be embedded within another display list. But if a list is assigned an identifier that has already been used, the new list replaces the previous list that had been assigned that identifier. Therefore, to avoid losing a list by accidentally reusing its identifier, we can let OpenGL generate an identifier for us, as follows:

```c
     listID = glGenLists (1);
```

This statement returns one (1) unused positive integer identifier to the variable `listID`. A range of unused integer list identifiers is obtained if we change the argument of `glGenLists` from the value 1 to some other positive integer. For
instance, if we invoke `glGenLists (6)`, then a sequence of six contiguous positive integer values is reserved and the first value in this list of identifiers is returned to the variable `listID`. A value of 0 is returned by the `glGenLists` function if an error occurs or if the system cannot generate the range of contiguous integers requested. Therefore, before using an identifier obtained from the `glGenLists` routine, we could check to be sure that it is not 0.

Although unused list identifiers can be generated with the `glGenList` function, we can independently query the system to determine whether a specific integer value has been used as a list name. The function to accomplish this is

```c
glIsList (listID);
```

A value of `GL_TRUE` is returned if the value of `listID` is an integer that has already been used as a display-list name. If the integer value has not been used as a list name, the `glIsList` function returns the value `GL_FALSE`.

### Executing OpenGL Display Lists

We execute a single display list with the statement

```c
glCallList (listID);
```

The following code segment illustrates the creation and execution of a display list. We first set up a display list that contains the description for a regular hexagon, defined in the \(xy\) plane using a set of six equally spaced vertices around the circumference of a circle, whose center coordinates are \((200, 200)\) and whose radius is 150. Then we issue a call to function `glCallList`, which displays the hexagon.

```c
const double TWO_PI = 6.2831853:

GLuint regHex;
GLdouble theta;
GLint x, y, k;

/* Set up a display list for a regular hexagon.
 * Vertices for the hexagon are six equally spaced
 * points around the circumference of a circle.
 */
regHex = glGenLists (1); // Get an identifier for the display list.
glNewList (regHex, GL_COMPILE);

```
Several display lists can be executed using the following two statements:

```c
glListBase (offsetValue);

glCallLists (nLists, arrayDataType, listIDArray);
```

The integer number of lists that we want to execute is assigned to parameter `nLists`, and parameter `listIDArray` is an array of display-list identifiers. In general, `listIDArray` can contain any number of elements, and invalid display-list identifiers are ignored. Also, the elements in `listIDArray` can be specified in a variety of data formats, and parameter `arrayDataType` is used to indicate a data type, such as `GL_BYTE`, `GL_INT`, `GL_FLOAT`, `GL_3_BYTES`, or `GL_4_BYTES`. A display-list identifier is calculated by adding the value in an element of `listIDArray` to the integer value of `offsetValue` that is given in the `glListBase` function. The default value for `offsetValue` is 0.

This mechanism for specifying a sequence of display lists that are to be executed allows us to set up groups of related display lists, whose identifiers are formed from symbolic names or codes. A typical example is a font set where each display-list identifier is the ASCII value of a character. When several font sets are defined, we use parameter `offsetValue` in the `glListBase` function to obtain a particular font described within the array `listIDArray`.

**Deleting OpenGL Display Lists**

We eliminate a contiguous set of display lists with the function call

```c
glDeleteLists (startID, nLists);
```

Parameter `startID` gives the initial display-list identifier, and parameter `nLists` specifies the number of lists that are to be deleted. For example, the statement

```c
glDeleteLists (5, 4);
```

eliminates the four display lists with identifiers 5, 6, 7, and 8. An identifier value that references a nonexistent display list is ignored.

---

**16 OpenGL Display-Window Reshape Function**

After the generation of our picture, we often want to use the mouse pointer to drag the display window to another screen location or to change its size. Changing the size of a display window could change its aspect ratio and cause objects to be distorted from their original shapes.

To allow us to compensate for a change in display-window dimensions, the GLUT library provides the following routine:

```c
glutReshapeFunc (winReshapeFcn);
```

We can include this function in the `main` procedure in our program, along with the other GLUT routines, and it will be activated whenever the display-window size is altered. The argument for this GLUT function is the name of a procedure that
is to receive the new display-window width and height. We can then use the new dimensions to reset the projection parameters and perform any other operations, such as changing the display-window color. In addition, we could save the new width and height values so that they could be used by other procedures in our program.

As an example, the following program illustrates how we might structure the `winReshapeFcn` procedure. The `glLoadIdentity` command is included in the reshape function so that any previous values for the projection parameters will not affect the new projection settings. This program displays the regular hexagon discussed in Section 15. Although the hexagon center (at the position of the circle center) in this example is specified in terms of the display-window parameters, the position of the hexagon is unaffected by any changes in the size of the display window. This is because the hexagon is defined within a display list, and only the original center coordinates are stored in the list. If we want the position of the hexagon to change when the display window is resized, we need to define the hexagon in another way or alter the coordinate reference for the display window. The output from this program is shown in Figure 30.

```c
#include <GL/glut.h>
#include <math.h>
#include <stdlib.h>

const double TWO_PI = 6.2831853;

/* Initial display-window size. */
GLsizei winWidth = 400, winHeight = 400;
GLuint regHex;

class screenPt
{
private:
GLint x, y;

Graphics Output Primitives
```
public:
   /* Default Constructor: initializes coordinate position to (0, 0). */
   screenPt ( ) {
      x = y = 0;
   }

   void setCoords (GLint xCoord, GLint yCoord) {
      x = xCoord;
      y = yCoord;
   }

   GLint getx ( ) const {
      return x;
   }

   GLint gety ( ) const {
      return y;
   }
};

static void init (void)
{
   screenPt hexVertex, circCtr;
   GLdouble theta;
   GLint k;

   /* Set circle center coordinates. */
   circCtr.setCoords (winWidth / 2, winHeight / 2);

   glClearColor (1.0, 1.0, 1.0, 0.0); // Display-window color = white.

   /* Set up a display list for a red regular hexagon.
   * Vertices for the hexagon are six equally spaced
   * points around the circumference of a circle.
   */
   regHex = glGenLists (1); // Get an identifier for the display list.
   glNewList (regHex, GL_COMPILE);
   glColor3f (1.0, 0.0, 0.0); // Set fill color for hexagon to red.
   glBegin (GL_POLYGON);
   for (k = 0; k < 6; k++) {
      theta = TWO_PI * k / 6.0;
      hexVertex.setCoords (circCtr.getx ( ) + 150 * cos (theta),
                           circCtr.gety ( ) + 150 * sin (theta));
      glVertex2i (hexVertex.getx ( ), hexVertex.gety ( ));
   }
   glEnd ( );
   glEndList ( );
}

void regHexagon (void)
{
   glClear (GL_COLOR_BUFFER_BIT);
   glCallList (regHex);
   glFlush ( );
}
void winReshapeFcn (int newWidth, int newHeight)
{
    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ( );
    gluOrtho2D (0.0, (GLdouble) newWidth, 0.0, (GLdouble) newHeight);

    glClear (GL_COLOR_BUFFER_BIT);
}

void main (int argc, char** argv)
{
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowPosition (100, 100);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Reshape-Function & Display-List Example");

    init ( );
    glutDisplayFunc (regHexagon);
    glutReshapeFunc (winReshapeFcn);

    glutMainLoop ( );
}

17 Summary

The output primitives discussed in this chapter provide the basic tools for constructing pictures with individual points, straight lines, curves, filled color areas, array patterns, and text. We specify primitives by giving their geometric descriptions in a Cartesian, world-coordinate reference system.

A fill area is a planar region that is to be displayed in a solid color or color pattern. Fill-area primitives in most graphics packages are polygons. But, in general, we could specify a fill region with any boundary. Often, graphics systems allow only convex polygon fill areas. In that case, a concave-polygon fill area can be displayed by dividing it into a set of convex polygons. Triangles are the easiest polygons to fill, because each scan line crossing a triangle intersects exactly two polygon edges (assuming that the scan line does not pass through any vertices).

The odd-even rule can be used to locate the interior points of a planar region. Other methods for defining object interiors are also useful, particularly with irregular, self-intersecting objects. A common example is the nonzero winding-number rule. This rule is more flexible than the odd-even rule for handling objects defined with multiple boundaries. We can also use variations of the winding-number rule to combine plane areas using Boolean operations.

Each polygon has a front face and a back face, which determines the spatial orientation of the polygon plane. This spatial orientation can be determined from the normal vector, which is perpendicular to the polygon plane and points...
in the direction from the back face to the front face. We can determine the components of the normal vector from the polygon plane equation or by forming a vector cross-product using three points in the plane, where the three points are taken in a counterclockwise order and the angle formed by the three points is less than 180°. All coordinate values, spatial orientations, and other geometric data for a scene are entered into three tables: vertex, edge, and surface-facet tables.

Additional primitives available in graphics packages include pattern arrays and character strings. Pattern arrays can be used to specify two-dimensional shapes, including a character set, using either a rectangular set of binary values or a set of color values. Character strings are used to provide picture and graph labeling.

Using the primitive functions available in the basic OpenGL library, we can generate points, straight-line segments, convex polygon fill areas, and either bitmap or pixmap pattern arrays. Routines for displaying character strings are available in GLUT. Other types of primitives, such as circles, ellipses, and concave-polygon fill areas, can be constructed or approximated with these functions, or they can be generated using routines in GLU and GLUT. All coordinate values are expressed in absolute coordinates within a right-handed Cartesian-coordinate reference system. Coordinate positions describing a scene can be given in either a two-dimensional or a three-dimensional reference frame. We can use integer or floating-point values to give a coordinate position, and we can also reference a position with a pointer to an array of coordinate values. A scene description is then transformed by viewing functions into a two-dimensional display on an output device, such as a video monitor. Except for the glRect function, each coordinate position for a set of points, lines, or polygons is specified in a glVertex function. And the set of glVertex functions defining each primitive is included between a glBegin/glEnd pair of statements, where the primitive type is identified with a symbolic constant as the argument for the glBegin function. When describing a scene containing many polygon fill surfaces, we can generate the display efficiently using OpenGL vertex arrays to specify geometric and other data.

In Table 1, we list the basic functions for generating output primitives in OpenGL. Some related routines are also listed in this table.

---

**Example Programs**

Here, we present a few example OpenGL programs illustrating the use of output primitives. Each program uses one or more of the functions listed in Table 1. A display window is set up for the output from each program.

The first program illustrates the use of a polyline, a set of polymarkers, and bit-mapped character labels to generate a line graph for monthly data over a period of one year. A proportionally spaced font is demonstrated, although a fixed-width font is usually easier to align with graph positions. Because the bitmaps are referenced at the lower-left corner by the raster-position function, we must shift the reference position to align the center of a text string with a plotted data position. Figure 31 shows the output of the line-graph program.
### Table 1

Summary of OpenGL Output Primitive Functions and Related Routines

<table>
<thead>
<tr>
<th>Function Description</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specifies a two-dimensional world-coordinate reference.</td>
<td>gluOrtho2D</td>
</tr>
<tr>
<td>Selects a coordinate position. This function must be placed within a glBegin/glEnd pair.</td>
<td>glVertex*</td>
</tr>
<tr>
<td>Plots one or more point positions, each specified in a glVertex function. The list of positions is then closed with a glEnd statement.</td>
<td>glBegin (GL_POINTS);</td>
</tr>
<tr>
<td>Displays a set of straight-line segments, whose endpoint coordinates are specified in glVertex functions. The list of endpoints is then closed with a glEnd statement.</td>
<td>glBegin (GL_LINES);</td>
</tr>
<tr>
<td>Displays a polyline, specified using the same structure as GL_LINES.</td>
<td>glBegin (GL_LINE_STRIP);</td>
</tr>
<tr>
<td>Displays a closed polyline, specified using the same structure as GL_LINES.</td>
<td>glBegin (GL_LINE_LOOP);</td>
</tr>
<tr>
<td>Displays a fill rectangle in the xy plane.</td>
<td>glRect*</td>
</tr>
<tr>
<td>Displays a fill polygon, whose vertices are given in glVertex functions and terminated with a glEnd statement.</td>
<td>glBegin (GL_POLYGON);</td>
</tr>
<tr>
<td>Displays a set of fill triangles using the same structure as GL_POLYGON.</td>
<td>glBegin (GL_TRIANGLES);</td>
</tr>
<tr>
<td>Displays a fill-triangle mesh, specified using the same structure as GL_POLYGON.</td>
<td>glBegin (GL_TRIANGLE_STRIP);</td>
</tr>
<tr>
<td>Displays a fill-triangle mesh in a fan shape with all triangles connected to the first vertex, specified with same structure as GL_POLYGON.</td>
<td>glBegin (GL_TRIANGLE_FAN);</td>
</tr>
<tr>
<td>Displays a set of fill quadrilaterals, specified with the same structure as GL_POLYGON.</td>
<td>glBegin (GL_QUADS);</td>
</tr>
<tr>
<td>Displays a fill-quadrilateral mesh, specified with the same structure as GL_POLYGON.</td>
<td>glBegin (GL_QUAD_STRIP);</td>
</tr>
<tr>
<td>Activates vertex-array features of OpenGL.</td>
<td>glEnableClientState (GL_VERTEX_ARRAY);</td>
</tr>
<tr>
<td>Specifies an array of coordinate values.</td>
<td>glVertexPointer (size, type, stride, array);</td>
</tr>
<tr>
<td>Displays a specified primitive type from array data.</td>
<td>glDrawElements (prim, num, type, array);</td>
</tr>
</tbody>
</table>
### Graphics Output Primitives

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glNewList (listID, listMode)</td>
<td>Defines a set of commands as a display list, terminated with a glEndList statement.</td>
</tr>
<tr>
<td>glGenLists</td>
<td>Generates one or more display-list identifiers.</td>
</tr>
<tr>
<td>glIsList</td>
<td>Queries OpenGL to determine whether a display-list identifier is in use.</td>
</tr>
<tr>
<td>glCallList</td>
<td>Executes a single display list.</td>
</tr>
<tr>
<td>glListBase</td>
<td>Specifies an offset value for an array of display-list identifiers.</td>
</tr>
<tr>
<td>glCallLists</td>
<td>Executes multiple display lists.</td>
</tr>
<tr>
<td>glDeleteLists</td>
<td>Eliminates a specified sequence of display lists.</td>
</tr>
<tr>
<td>glRasterPos*</td>
<td>Specifies a two-dimensional or three-dimensional current position for the frame buffer. This position is used as a reference for bitmap and pixmap patterns.</td>
</tr>
<tr>
<td>glBitmap (w, h, x0, y0, xShift, yShift, pattern);</td>
<td>Specifies a binary pattern that is to be mapped to pixel positions relative to the current position.</td>
</tr>
<tr>
<td>glDrawPixels (w, h, type, format, pattern);</td>
<td>Specifies a color pattern that is to be mapped to pixel positions relative to the current position.</td>
</tr>
<tr>
<td>glDrawBuffer</td>
<td>Selects one or more buffers for storing a pixmap.</td>
</tr>
<tr>
<td>glReadPixels</td>
<td>Saves a block of pixels in a selected array.</td>
</tr>
<tr>
<td>glCopyPixels</td>
<td>Copies a block of pixels from one buffer position to another.</td>
</tr>
<tr>
<td>glLogicOp</td>
<td>Selects a logical operation for combining two pixel arrays, after enabling with the constant GL_COLOR_LOGIC_OP.</td>
</tr>
<tr>
<td>glutBitmapCharacter (font, char);</td>
<td>Specifies a font and a bitmap character for display.</td>
</tr>
<tr>
<td>glutStrokeCharacter (font, char);</td>
<td>Specifies a font and an outline character for display.</td>
</tr>
<tr>
<td>glutReshapeFunc</td>
<td>Specifies actions to be taken when display-window dimensions are changed.</td>
</tr>
</tbody>
</table>
FIGURE 31
A polyline and polymarker plot of data points output by the lineGraph routine.

#include <GL/glut.h>

GLsizei winWidth = 600, winHeight = 500; // Initial display window size.
GLint xRaster = 25, yRaster = 150; // Initialize raster position.
GLubyte label [36] = {'J', 'a', 'n', 'F', 'e', 'b', 'M', 'a', 'r',
'A', 'p', 'r', 'M', 'a', 'y', 'J', 'u', 'n',
'J', 'u', 'l', 'A', 'u', 'g', 'S', 'e', 'p',
'O', 'c', 't', 'N', 'o', 'v', 'D', 'e', 'c'};

GLint dataValue [12] = {420, 342, 324, 310, 262, 185,
190, 196, 217, 240, 312, 438};

void init (void)
{
    glClearColor (1.0, 1.0, 1.0, 1.0); // White display window.
    glMatrixMode (GL_PROJECTION);
    gluOrtho2D (0.0, 600.0, 0.0, 500.0);
}

void lineGraph (void)
{
    GLint month, k;
    GLint x = 30; // Initialize x position for chart.
    glClear (GL_COLOR_BUFFER_BIT); // Clear display window.
    glColor3f (0.0, 0.0, 1.0); // Set line color to blue.
We use the same data set in the second program to produce the bar chart in Figure 32. This program illustrates an application of rectangular fill areas, as well as bitmapped character labels.
Pie charts are used to show the percentage contribution of individual parts to the whole. The next program constructs a pie chart, using the midpoint routine for generating a circle. Example values are used for the number and relative sizes of the slices, and the output from this program appears in Figure 33.
#include <GL/glut.h>
#include <stdlib.h>
#include <math.h>

const GLdouble twoPi = 6.283185;

class scrPt {
public:
    GLint x, y;
};

 GLsizei winWidth = 400, winHeight = 300; // Initial display window size.

void init (void) {
    glClearColor (1.0, 1.0, 1.0, 1.0);
    glMatrixMode (GL_PROJECTION);
    gluOrtho2D (0.0, 200.0, 0.0, 150.0);
}

// Midpoint routines for displaying a circle.

void pieChart (void) {
    scrPt circCtr, piePt;
    GLint radius = winWidth / 4; // Circle radius.
    GLint k, nSlices = 12; // Number of slices.
    GLfloat dataValues[12] = {10.0, 7.0, 13.0, 5.0, 13.0, 14.0, 3.0, 16.0, 5.0, 3.0, 17.0, 8.0};
    GLfloat dataSum = 0.0;
    circCtr.x = winWidth / 2; // Circle center position.
    circCtr.y = winHeight / 2;
    circleMidpoint (circCtr, radius); // Call a midpoint circle-plot routine.
    for (k = 0; k < nSlices; k++)
        dataSum += dataValues[k];
    for (k = 0; k < nSlices; k++) {
        sliceAngle = twoPi * dataValues[k] / dataSum + previousSliceAngle;
        piePt.x = circCtr.x + radius * cos (sliceAngle);
        piePt.y = circCtr.y + radius * sin (sliceAngle);
        glBegin (GL_LINES);
        glVertex2i (circCtr.x, circCtr.y);
        glVertex2i (piePt.x, piePt.y);
        glEnd ( );
        previousSliceAngle = sliceAngle;
    }
}
void displayFcn (void)
{
    glClear (GL_COLOR_BUFFER_BIT); // Clear display window.
    glColor3f (0.0, 0.0, 1.0); // Set circle color to blue.
    pieChart ( );
    glFlush ( );
}

void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ( );
    gluOrtho2D (0.0, GLdouble (newWidth), 0.0, GLdouble (newHeight));
    glClear (GL_COLOR_BUFFER_BIT);
    /* Reset display-window size parameters. */
    winWidth = newWidth;
    winHeight = newHeight;
}

void main (int argc, char** argv)
{
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowPosition (100, 100);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Pie Chart");
    init ( );
    glutDisplayFunc (displayFcn);
    glutReshapeFunc (winReshapeFcn);
    glutMainLoop ( );
}

Some variations on the circle equations are displayed by our last example program, which uses the parametric polar equations (6-28) to compute points along the curve paths. These points are then used as the endpoint positions for straight-line sections, displaying the curves as approximating polylines. The curves shown in Figure 34 are generated by varying the radius \( r \) of a circle. Depending on how we vary \( r \), we can produce a limaçon, cardioid, spiral, or other similar figure.

![Curved figures displayed by the drawCurve procedure: (a) limaçon, (b) cardioid, (c) three-leaf curve, (d) four-leaf curve, and (e) spiral.](image_url)

**Figure 34**
Curved figures displayed by the `drawCurve` procedure: (a) limaçon, (b) cardioid, (c) three-leaf curve, (d) four-leaf curve, and (e) spiral.
```c
#include <GL/glut.h>
#include <stdlib.h>
#include <math.h>
#include <iostream.h>

struct screenPt
{
    GLint x;
    GLint y;
};

typedef enum { limacon = 1, cardioid, threeLeaf, fourLeaf, spiral } curveName;

GLsizei winWidth = 600, winHeight = 500; // Initial display window size.

void init (void)
{
    glClearColor (1.0, 1.0, 1.0, 1.0);
    glMatrixMode (GL_PROJECTION);
    gluOrtho2D (0.0, 200.0, 0.0, 150.0);
}

void lineSegment (screenPt pt1, screenPt pt2)
{
    glBegin (GL_LINES);
    glVertex2i (pt1.x, pt1.y);
    glVertex2i (pt2.x, pt2.y);
    glEnd ( );
}

void drawCurve (GLint curveNum)
{
    /* The limacon of Pascal is a modification of the circle equation
       * with the radius varying as r = a * cos (theta) + b, where a
       * and b are constants. A cardioid is a limacon with a = b.
       * Three-leaf and four-leaf curves are generated when
       * r = a * cos (n * theta), with n = 3 and n = 2, respectively.
       * A spiral is displayed when r is a multiple of theta.
       */

    const GLdouble twoPi = 6.283185;
    const GLint a = 175, b = 60;

    GLfloat r, theta, dtheta = 1.0 / float (a);
    GLint x0 = 200, y0 = 250; // Set an initial screen position.
    screenPt curvePt[2];

    glColor3f (0.0, 0.0, 0.0); // Set curve color to black.
    curvePt[0].x = x0; // Initialize curve position.
    curvePt[0].y = y0;
```
switch (curveNum) {
    case limacon:  curvePt[0].x += a + b;  break;
    case cardioid: curvePt[0].x += a + a;  break;
    case threeLeaf: curvePt[0].x += a;  break;
    case fourLeaf:  curvePt[0].x += a;  break;
    case spiral: break;
    default: break;
}

theta = dtheta;
while (theta < two_Pi) {
    switch (curveNum) {
        case limacon:
            r = a * cos (theta) + b;  break;
        case cardioid:
            r = a * (1 + cos (theta));  break;
        case threeLeaf:
            r = a * cos (3 * theta);  break;
        case fourLeaf:
            r = a * cos (2 * theta);  break;
        case spiral:
            r = (a / 4.0) * theta;  break;
        default: break;
    }
    curvePt[1].x = x0 + r * cos (theta);
    curvePt[1].y = y0 + r * sin (theta);
    lineSegment (curvePt[0], curvePt[1]);
    curvePt[0].x = curvePt[1].x;
    curvePt[0].y = curvePt[1].y;
    theta += dtheta;
}

void displayFcn (void)
{
    GLint curveNum;
    glClear (GL_COLOR_BUFFER_BIT);  // Clear display window.
    cout << \n\nEnter the integer value corresponding to\n\none of the following curve names.\nPress any other key to exit.\n\n1-limacon, 2-cardioid, 3-threeLeaf, 4-fourLeaf, 5-spiral:  
    cin >> curveNum;
    if (curveNum == 1 || curveNum == 2 || curveNum == 3 || curveNum == 4
        || curveNum == 5)
        drawCurve (curveNum);
    else
        exit (0);
    glFlush ( );
}
void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ();
    gluOrtho2D (0.0, (GLdouble) newWidth, 0.0, (GLdouble) newHeight);
    glClear (GL_COLOR_BUFFER_BIT);
}

void main (int argc, char** argv)
{
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowPosition (100, 100);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Draw Curves");
    init ( );
    glutDisplayFunc (displayFcn);
    glutReshapeFunc (winReshapeFcn);
    glutMainLoop ( );
}

REFERENCES


EXERCISES
1 Set up geometric data tables as in Figure 16 for a square pyramid (a square base with four triangular sides that meet at a pinnacle).
2 Set up geometric data tables for a square pyramid using just a vertex table and a surface-facet table, then store the same information using just the surface-facet table. Compare the two methods for representing the unit cube with a representation using the three tables in the previous exercise. Estimate the storage requirements for each.
3 Set up a procedure for establishing the geometric data tables for any input set of points defining the polygon facets for the surface of a three-dimensional object.
4 Devise routines for checking the three geometric data tables in Figure 16 to ensure consistency and completeness.
5 Calculate the plane parameters \( A, B, C, \) and \( D \) for each face of a unit cube centered at the world coordinate origin.
6 Write a program for calculating parameters \( A, B, C, \) and \( D \) for an input mesh of polygon-surface facets.
7 Write a procedure to determine whether an input coordinate position is in front of a polygon surface or behind it, given the plane parameters \( A, B, C, \) and \( D \) for the polygon.
8 Write a procedure to determine whether a given point is inside or outside of a cube with a given set of coordinates.
9 If the coordinate reference for a scene is changed from a right-handed system to a left-handed system, what changes could we make in the values of surface plane parameters \( A, B, C, \) and \( D \) to ensure that the orientation of the plane is correctly described?
10 Given that the first three vertices, \( V_1, V_2, \) and \( V_3, \) of a pentagon have been used to calculate plane parameters \( A = 15, B = 21, C = 9, D = 0, \) determine from the final two vertices \( V_4 = (2, -1, -1) \) and \( V_5 = (1, -2, 2) \) whether the pentagon is planar or non-planar.
11 Develop a procedure for identifying a nonplanar vertex list for a quadrilateral.

12 Extend the algorithm of the previous exercise to identify a nonplanar vertex list that contains more than four coordinate positions.

13 Write a procedure to split a set of four polygon vertex positions into a set of triangles.

14 Split the octagon given by the list of vertices \( V_1, V_2, V_3, V_4, V_5, V_6, V_7, V_8 \) into a set of triangles and give the vertices that make up each triangle.

15 Devise an algorithm for splitting a set of \( n \) polygon vertex positions, with \( n > 4 \), into a set of triangles.

16 Set up an algorithm for identifying a degenerate polygon vertex list that may contain repeated vertices or collinear vertices.

17 Devise an algorithm for identifying a polygon vertex list that contains intersecting edges.

18 Write a routine to identify concave polygons by calculating cross-products of pairs of edge vectors.

19 Write a routine to split a concave polygon, using the vector method.

20 Write a routine to split a concave polygon, using the rotational method.

21 Devise an algorithm for determining interior regions for any input set of vertices using the nonzero winding-number rule and cross-product calculations to identify the direction for edge crossings.

22 Devise an algorithm for determining interior regions for any input set of vertices using the nonzero winding-number rule and dot-product calculations to identify the direction for edge crossings.

23 What regions of the self-intersecting polyline shown in Figure 12 have a positive winding number? What are the regions that have a negative winding number? What regions have a winding number greater than 1?

24 Write a routine to implement a text-string function that has two parameters: one parameter specifies a world-coordinate position and the other parameter specifies a text string.

25 Write a routine to implement a polynumber function that has two parameters: one parameter is the character that is to be displayed and the other parameter is a list of world-coordinate positions.

26 Modify the example program in Section 7 so that the displayed hexagon is always at the center of the display window, regardless of how the display window may be resized.

27 Write a complete program for displaying a bar chart. Input to the program is to include the data points and the labeling required for the \( x \) and \( y \) axes. The data points are to be scaled by the program so that the graph is displayed across the full area of a display window.

28 Write a program to display a bar chart in any selected area of a display window.

29 Write a procedure to display a line graph for any input set of data points in any selected area of the screen, with the input data set scaled to fit the selected screen area. Data points are to be displayed as asterisks joined with straight-line segments, and the \( x \) and \( y \) axes are to be labeled according to input specifications. (Instead of asterisks, small circles or some other symbols could be used to plot the data points.)

30 Using a circle function, write a routine to display a pie chart with appropriate labeling. Input to the routine is to include a data set giving the distribution of the data over some set of intervals, the name of the pie chart, and the names of the intervals. Each section label is to be displayed outside the boundary of the pie chart near the corresponding pie section.

IN MORE DEPTH

1 For this exercise, draw a rough sketch of what a single “snapshot” of your application might look like and write a program to display this snapshot. Choose a background color and default window size. Make sure the snapshot includes at least a few objects. Represent each object as a polygonal approximation to the true object. Use a different shape for each object type. Represent at least one of the objects as a concave polygon. Make each object its own color distinct from the background color. It is a good idea to write a separate function for each object (or each object type) in which you define the representation. Use display lists to create and display each object. Include a display window reshape function to redraw the scene appropriately if the window is if the window is resized.

2 Choose one of the concave polygons you generated in the previous exercise and set up the vertex, edge, and surface facet tables for the shape as described in Section 7. Now split the shape it into a set of convex polygons using the vector method given in the same section. Then split each of the resulting convex polygons into a set of triangles using the method described in Section 7 as well. Finally, set up the vertex, edge, and surface facet tables for the resulting set of triangles. Compare the two table sets and the amount of memory needed to store each.
In general, a parameter that affects the way a primitive is to be displayed is referred to as an **attribute parameter**. Some attribute parameters, such as color and size, determine the fundamental characteristics of a primitive. Other attributes specify how the primitive is to be displayed under special conditions. Examples of special-condition attributes are the options such as visibility or detectability within an interactive object-selection program. These special-condition attributes are explored in later chapters. Here, we treat only those attributes that control the basic display properties of graphics primitives, without regard for special situations. For example, lines can be dotted or dashed, fat or thin, and blue or orange. Areas might be filled with one color or with a multicolor pattern. Text can appear reading from left to right, slanted diagonally across the screen, or in vertical columns. Individual characters can be displayed in different fonts, colors, and sizes. And we can apply intensity variations at the edges of objects to smooth out the raster stair-step effect.
Attributes of Graphics Primitives

One way to incorporate attribute options into a graphics package is to extend the parameter list associated with each graphics-primitive function to include the appropriate attribute values. A line-drawing function, for example, could contain additional parameters to set the color, width, and other properties of a line. Another approach is to maintain a system list of current attribute values. Separate functions are then included in the graphics package for setting the current values in the attribute list. To generate a primitive, the system checks the relevant attributes and invokes the display routine for that primitive using the current attribute settings. Some graphics packages use a combination of methods for setting attribute values, and other libraries, including OpenGL, assign attributes using separate functions that update a system attribute list.

A graphics system that maintains a list for the current values of attributes and other parameters is referred to as a state system or state machine. Attributes of output primitives and some other parameters, such as the current frame-buffer position, are referred to as state variables or state parameters. When we assign a value to one or more state parameters, we put the system into a particular state, and that state remains in effect until we change the value of a state parameter.

1 OpenGL State Variables

Attribute values and other parameter settings are specified with separate functions that define the current OpenGL state. The state parameters in OpenGL include color and other primitive attributes, the current matrix mode, the elements of the model-view matrix, the current position for the frame buffer, and the parameters for the lighting effects in a scene. All OpenGL state parameters have default values, which remain in effect until new values are specified. At any time, we can query the system to determine the current value of a state parameter. In the following sections of this chapter, we discuss only the attribute settings for output primitives. Other state parameters are examined in later chapters.

All graphics primitives in OpenGL are displayed with the attributes in the current state list. Changing one or more of the attribute settings affects only those primitives that are specified after the OpenGL state is changed. Primitives that were defined before the state change retain their attributes. Thus, we can display a green line, change the current color to red, and define another line segment. Both the green line and the red line will then be displayed. Also, some OpenGL state values can be specified within glBegin/glEnd pairs, along with the coordinate values, so that parameter settings can vary from one coordinate position to another.

2 Color and Grayscale

A basic attribute for all primitives is color. Various color options can be made available to a user, depending on the capabilities and design objectives of a particular system. Color options can be specified numerically or selected from menus or displayed slider scales. For a video monitor, these color codes are then converted to intensity-level settings for the electron beams. With color plotters, the codes might control ink-jet deposits or pen selections.

RGB Color Components

In a color raster system, the number of color choices available depends on the amount of storage provided per pixel in the frame buffer. Also, color information
can be stored in the frame buffer in two ways: We can store red, green, and blue (RGB) color codes directly in the frame buffer, or we can put the color codes into a separate table and use the pixel locations to store index values referencing the color-table entries. With the direct storage scheme, whenever a particular color code is specified in an application program, that color information is placed in the frame buffer at the location of each component pixel in the output primitives to be displayed in that color. A minimum number of colors can be provided in this scheme with 3 bits of storage per pixel, as shown in Table 1. Each of the three bit positions is used to control the intensity level (either on or off, in this case) of the corresponding electron gun in an RGB monitor. The leftmost bit controls the red gun, the middle bit controls the green gun, and the rightmost bit controls the blue gun. Adding more bits per pixel to the frame buffer increases the number of color choices that we have. With 6 bits per pixel, 2 bits can be used for each gun. This allows four different intensity settings for each of the three color guns, and a total of 64 color options are available for each screen pixel. As more color options are provided, the storage required for the frame buffer also increases. With a resolution of 1024 × 1024, a full-color (24-bit per pixel) RGB system needs 3 MB of storage for the frame buffer.

Color tables are an alternate means for providing extended color capabilities to a user without requiring large frame buffers. At one time, this was an important consideration; but today, hardware costs have decreased dramatically and extended color capabilities are fairly common, even in low-end personal computer systems. So most of our examples will simply assume that RGB color codes are stored directly in the frame buffer.

**Color Tables**

Figure 1 illustrates a possible scheme for storing color values in a color lookup table (or color map). Sometimes a color table is referred to as a video lookup table. Values stored in the frame buffer are now used as indices into the color table. In this example, each pixel can reference any of the 256 table positions, and each entry in the table uses 24 bits to specify an RGB color. For the hexadecimal color code 0x0821, a combination green-blue color is displayed for pixel location (x, y). Systems employing this particular lookup table allow a user to select any

<table>
<thead>
<tr>
<th><strong>Color Code</strong></th>
<th><strong>RED</strong></th>
<th><strong>GREEN</strong></th>
<th><strong>BLUE</strong></th>
<th><strong>Displayed Color</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>Black</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>Blue</td>
</tr>
<tr>
<td>2</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>Green</td>
</tr>
<tr>
<td>3</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>Cyan</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>Red</td>
</tr>
<tr>
<td>5</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>Magenta</td>
</tr>
<tr>
<td>6</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>Yellow</td>
</tr>
<tr>
<td>7</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>White</td>
</tr>
</tbody>
</table>
256 colors for simultaneous display from a palette of nearly 17 million colors. Compared to a full-color system, this scheme reduces the number of simultaneous colors that can be displayed, but it also reduces the frame-buffer storage requirement to 1 MB. Multiple color tables are sometimes available for handling specialized rendering applications, such as antialiasing, and they are used with systems that contain more than one color output device.

A color table can be useful in a number of applications, and it can provide a “reasonable” number of simultaneous colors without requiring large frame buffers. For most applications, 256 or 512 different colors are sufficient for a single picture. Also, table entries can be changed at any time, allowing a user to be able to experiment easily with different color combinations in a design, scene, or graph without changing the attribute settings for the graphics data structure. When a color value is changed in the color table, all pixels with that color index immediately change to the new color. Without a color table, we can change the color of a pixel only by storing the new color at that frame-buffer location. Similarly, data-visualization applications can store values for some physical quantity, such as energy, in the frame buffer and use a lookup table to experiment with various color combinations without changing the pixel values. Also, in visualization and image-processing applications, color tables are a convenient means for setting color thresholds so that all pixel values above or below a specified threshold can be set to the same color. For these reasons, some systems provide both capabilities for storing color information. A user can then elect either to use color tables or to store color codes directly in the frame buffer.

Grayscale

Because color capabilities are now common in computer-graphics systems, we use RGB color functions to set shades of gray, or grayscale, in an application program. When an RGB color setting specifies an equal amount of red, green, and blue, the result is some shade of gray. Values close to 0 for the color components produce dark gray, and higher values near 1.0 produce light gray. Applications for grayscale display methods include enhancing black-and-white photographs and generating visualization effects.

**FIGURE 1**

A color lookup table with 24 bits per entry that is accessed from a frame buffer with 8 bits per pixel. A value of 196 stored at pixel position \((x, y)\) references the location in this table containing the hexadecimal value 0x0821 (a decimal value of 2081). Each 8-bit segment of this entry controls the intensity level of one of the three electron guns in an RGB monitor.
Other Color Parameters

In addition to an RGB specification, other three-component color representations are useful in computer-graphics applications. For example, color output on printers is described with cyan, magenta, and yellow color components, and color interfaces sometimes use parameters such as lightness and darkness to choose a color. Also, color, and light in general, are complex subjects, and many terms and concepts have been devised in the fields of optics, radiometry, and psychology to describe the various aspects of light sources and lighting effects. Physically, we can describe a color as electromagnetic radiation with a particular frequency range and energy distribution, but then there are also the characteristics of our perception of the color. Thus, we use the physical term intensity to quantify the amount of light energy radiating in a particular direction over a period of time, and we use the psychological term luminance to characterize the perceived brightness of the light. We discuss these terms and other color concepts in greater detail when we consider methods for modeling lighting effects and the various models for describing color.

3 OpenGL Color Functions

In an OpenGL color routines use one function to set the color for the display window, and use another function to specify a color for the straight-line segment. Set the color display mode to RGB with the statement

```c
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
```

The first constant in the argument list states that we are using a single buffer for the frame buffer, and the second constant puts us into the RGB mode, which is the default color mode. If we wanted to specify colors by an index into a color table, we would replace the OpenGL constant GLUT_RGB with GLUT_INDEX. When we specify a particular set of color values for primitives, we define the color state of OpenGL. The current color is applied to all subsequently defined primitives until we change the color settings. A new color specification affects only the objects we define after the color change.

The OpenGL RGB and RGBA Color Modes

Most color settings for OpenGL primitives are made in the RGB mode. In addition to red, green, and blue color coefficients, there is a fourth component called the alpha coefficient which is used to control color blending. The four-dimensional color specification is called RGBA mode, and we can select it using the OpenGL constant GLUT_RGBA when we call glutInitDisplayMode. This fourth color parameter can be used to control color blending for overlapping primitives. An important application of color blending is in the simulation of transparency effects. For these calculations, the value of alpha corresponds to a transparency (or opacity) setting. The alpha value is optional; the only difference between the RGB and RGBA modes is whether we are employing it for color blending.

In the RGB (or RGBA) mode, we select the current color components with the function

```c
    glColor* (colorComponents);
```
Suffix codes are similar to those for the `glVertex` function. We use a code of either 3 or 4 to specify the RGB or RGBA mode along with the numerical data-type code and an optional vector suffix. The suffix codes for the numerical data types are `b` (byte), `i` (integer), `s` (short), `f` (float), and `d` (double), as well as unsigned numerical values. Floating-point values for the color components are in the range from 0.0 to 1.0, and the default color components for `glColor`, including the alpha value, are (1.0, 1.0, 1.0, 1.0), which sets the RGB color to white and the alpha value to 1.0. If we select the current color using an RGB specification (i.e., we use `glColor3` instead of `glColor4`), the alpha component will be automatically set to 1.0 to indicate that we do not want color blending. As an example, the following statement uses floating-point values in RGB mode to set the current color for primitives to cyan (a combination of the highest intensities for green and blue):

```
setColor3f (0.0, 1.0, 1.0);
```

Using an array specification for the three color components, we could set the color in this example as

```
setColor3fv (colorArray);
```

An OpenGL color selection can be assigned to individual point positions within `glBegin`/`glEnd` pairs.

Internally, OpenGL represents color information in floating-point format. We can specify colors using integer values, but they will be converted automatically to floating-point. The conversion is based on the data type we choose and the range of values that we can specify in that type. For unsigned types, the minimum value will be converted to a floating-point 0.0, and the maximum value to 1.0; for signed values, the minimum will be converted to \(-1.0\) and the maximum to 1.0. For example, unsigned byte values (suffix code `ub`) have a range of 0 to 255, which corresponds to the color specification system used by some windowing systems. We could specify the cyan color used in our previous example this way:

```
setColor3ub (0, 255, 255);
```

However, if we were to use unsigned 32-bit integers (suffix code `ui`), the range is 0 to 4,294,967,295! At this scale, small changes in color component values are essentially invisible; to make a one-percent change in the intensity of a single color component, for instance, we would need to change that component’s value by 42,949,673. For that reason, the most commonly used data types are floating-point and small integer types.

### OpenGL Color-Index Mode

Color specifications in OpenGL can also be given in the color-index mode, which references values in a color table. Using this mode, we set the current color by specifying an index into a color table as follows:

```
setColor (colorIndex);
```

Parameter `colorIndex` is assigned a nonnegative integer value. This index value is then stored in the frame-buffer positions for subsequently specified primitives. We can specify the color index in any of the following data types: unsigned byte,
integer, or floating point. The data type for parameter colorIndex is indicated with a suffix code of ub, s, i, d, or f, and the number of index positions in a color table is always a power of 2, such as 256 or 1024. The number of bits available at each table position depends on the hardware features of the system. As an example of specifying a color in index mode, the following statement sets the current color index to the value 196:

```c
glIndexi (196);
```

All primitives defined after this statement will be assigned the color stored at that position in the color table until the current color is changed.

There are no functions provided in the core OpenGL library for loading values into a color-lookup table because table-processing routines are part of a window system. Also, some window systems support multiple color tables and full color, while other systems may have only one color table and limited color choices. However, we do have a GLUT routine that interacts with a window system to set color specifications into a table at a given index position as follows:

```c
glutSetColor (index, red, green, blue);
```

Color parameters red, green, and blue are assigned floating-point values in the range from 0.0 to 1.0. This color is then loaded into the table at the position specified by the value of parameter index.

Routines for processing three other color tables are provided as extensions to the OpenGL core library. These routines are part of the Imaging Subset of OpenGL. Color values stored in these tables can be used to modify pixel values as they are processed through various buffers. Some examples of using these tables are setting camera focusing effects, filtering out certain colors from an image, enhancing certain intensities or making brightness adjustments, converting a grayscale photograph to color, and antialiasing a display. In addition, we can use these tables to change color models; that is, we can change RGB colors to another specification using three other “primary” colors (such as cyan, magenta, and yellow).

A particular color table in the Imaging Subset of OpenGL is activated with the glEnable function using one of the table names: GL_COLOR_TABLE, GL_POST_CONVOLUTION_COLOR_TABLE, or GL_POST_COLOR_MATRIX_COLOR_TABLE. We can then use routines in the Imaging Subset to select a particular color table, set color-table values, copy table values, or specify which component of a pixel’s color we want to change and how we want to change it.

**OpenGL Color Blending**

In many applications, it is convenient to be able to combine the colors of overlapping objects or to blend an object with the background. Some examples are simulating a paintbrush effect, forming a composite image of two or more pictures, modeling transparency effects, and antialiasing the objects in a scene. Most graphics packages provide methods for producing various color-mixing effects, and these procedures are called color-blending functions or image-compositing functions. In OpenGL, the colors of two objects can be blended by first loading one object into the frame buffer, then combining the color of the second object with the frame-buffer color. The current frame-buffer color is referred to as the OpenGL destination color and the color of the second object is the OpenGL source color. Blending methods can be performed only in RGB or RGBA mode. To apply
color blending in an application, we first need to activate this OpenGL feature using the following function:

```c
glEnable (GL_BLEND);
```

We turn off the color-blending routines in OpenGL with

```c
glDisable (GL_BLEND);
```

If color blending is not activated, an object’s color simply replaces the frame-buffer contents at the object’s location.

Colors can be blended in a number of different ways, depending on the effects that we want to achieve, and we generate different color effects by specifying two sets of **blending factors**. One set of blending factors is for the current object in the frame buffer (the “destination object”), and the other set of blending factors is for the incoming (“source”) object. The new, blended color that is then loaded into the frame buffer is calculated as

\[
(S_R s, S_G s, S_B s, S_A s, D_R d, D_G d, D_B d, D_A d)
\]

where the RGBA source color components are \((S_R s, S_G s, S_B s, S_A s)\), the destination color components are \((D_R d, D_G d, D_B d, D_A d)\), the source blending factors are \((S_R, S_G, S_B, S_A)\), and the destination blending factors are \((D_R, D_G, D_B, D_A)\). Computed values for the combined color components are clamped to the range from 0.0 to 1.0. That is, any sum greater than 1.0 is set to the value 1.0, and any sum less than 0.0 is set to 0.0.

We select the blending-factor values with the OpenGL function

```c
glBlendFunc (sFactor, dFactor);
```

Parameters `sFactor` and `dFactor`, the source and destination factors, are each assigned an OpenGL symbolic constant specifying a predefined set of four blending coefficients. For example, the constant `GL_ZERO` yields the blending factors \((0.0, 0.0, 0.0, 0.0)\) and `GL_ONE` gives us the set \((1.0, 1.0, 1.0, 1.0)\). We could set all four blending factors either to the destination alpha value or to the source alpha value using `GL_DST_ALPHA` or `GL_SRC_ALPHA`. Other OpenGL constants that are available for setting the blending factors include `GL_ONE_MINUS_DST_ALPHA`, `GL_ONE_MINUS_SRC_ALPHA`, `GL_DST_COLOR`, and `GL_SRC_COLOR`. These blending factors are often used for simulating transparency, and they are discussed in greater detail in Section 18-4. The default value for parameter `sFactor` is `GL_ONE`, and the default value for parameter `dFactor` is `GL_ZERO`. Hence, the default values for the blending factors result in the incoming color values replacing the current values in the frame buffer.

**OpenGL Color Arrays**

We can also specify color values for a scene in combination with the coordinate values in a vertex array. This can be done either in RGB mode or in color-index mode. As with vertex arrays, we must first activate the color-array features of OpenGL as follows:

```c
glEnableClientState (GL_COLOR_ARRAY);
```

Then, for RGB color mode, we specify the location and format of the color components with

```c
glColorPointer (nColorComponents, dataType,
               offset, colorArray);
```
Parameter \( nColorComponents \) is assigned a value of either 3 or 4, depending on whether we are listing RGB or RGBA color components in the array \( \text{colorArray} \). An OpenGL symbolic constant such as GL_INT or GL_FLOAT is assigned to parameter \( \text{dataType} \) to indicate the data type for the color values. For a separate color array, we can assign the value 0 to parameter \( \text{offset} \). However, if we combine color data with vertex data in the same array, the \( \text{offset} \) value is the number of bytes between each set of color components in the array.

As an example of using color arrays, we can modify a vertex-array to include a color array. The following code fragment sets the color of all vertices on the front face of the cube to blue, and all vertices of the back face are assigned the color red:

```c
typedef GLint vertex3 [3], color3 [3];

vertex3 pt [8] = { {0, 0, 0}, {0, 1, 0}, {1, 0, 0},
                  {1, 1, 0}, {0, 0, 1}, {0, 1, 1}, {1, 0, 1}, {1, 1, 1} };
color3 hue [8] = { {1, 0, 0}, {1, 0, 0}, {0, 0, 1},
                  {0, 0, 1}, {1, 0, 0}, {1, 0, 0}, {0, 0, 1}, {0, 0, 1} };

glEnableClientState (GL_VERTEX_ARRAY);
glEnableClientState (GL_COLOR_ARRAY);

glVertexPointer (3, GL_INT, 0, pt);
glColorPointer (3, GL_INT, 0, hue);
```

We can even stuff both the colors and the vertex coordinates into one interlaced array. Each of the pointers would then reference the single interlaced array with an appropriate \( \text{offset} \) value. For example,

```c
static GLint hueAndPt [ ] =
    {1, 0, 0, 0, 0, 1, 0, 0, 0, 1, 0, 1, 1, 0, 0, 1, 1, 0, 0, 1, 1, 0, 1, 1, 0, 0, 1, 1, 1, 1};

glVertexPointer (3, GL_INT, 6*sizeof(GLint), hueAndPt[3]);
glColorPointer (3, GL_INT, 6*sizeof(GLint), hueAndPt[0]);
```

The first three elements of this array specify an RGB color value, the next three elements specify a set of \((x, y, z)\) vertex coordinates, and this pattern continues to the last color-vertex specification. We set the \( \text{offset} \) parameter to the number of bytes between successive color, or vertex, values, which is \( 6 \times \text{sizeof} \) (GLint) for both. Color values start at the first element of the interlaced array, which is \( \text{hueAndPt [0]} \), and vertex values start at the fourth element, which is \( \text{hueAndPt [3]} \).

Because a scene generally contains several objects, each with multiple planar surfaces, OpenGL provides a function in which we can specify all the vertex and color arrays at once, as well as other types of information. If we change the color and vertex values in this example to floating-point, we use this function in the form

```c
glInterleavedArrays (GL_C3F_V3F, 0, hueAndPt);
```
Attributes of Graphics Primitives

The first parameter is an OpenGL constant that indicates three-element floating-point specifications for both color \((C)\) and vertex coordinates \((V)\). The elements of array \text{hueAndPt} are to be interlaced with the color for each vertex listed before the coordinates. This function also automatically enables both vertex and color arrays.

In color-index mode, we define an array of color indices with

\[ \text{glIndexPointer} \left( \text{type}, \text{stride}, \text{colorIndex} \right); \]

Color indices are listed in the array \text{colorIndex} and the \text{type} and \text{stride} parameters are the same as in \text{glColorPointer}. No \text{size} parameter is needed because color-table indices are specified with a single value.

Other OpenGL Color Functions

The following function selects RGB color components for a display window:

\[ \text{glClearColor} \left( \text{red}, \text{green}, \text{blue}, \text{alpha} \right); \]

Each color component in the designation (red, green, and blue), as well as the \text{alpha} parameter, is assigned a floating-point value in the range from 0.0 to 1.0. The default value for all four parameters is 0.0, which produces the color black. If each color component is set to 1.0, the clear color is white. Shades of gray are obtained with identical values for the color components between 0.0 and 1.0. The fourth parameter, \text{alpha}, provides an option for blending the previous color with the current color. This can occur only if we activate the blending feature of OpenGL; color blending cannot be performed with values specified in a color table.

There are several color buffers in OpenGL that can be used as the current refresh buffer for displaying a scene, and the \text{glClearColor} function specifies the color for all the color buffers. We then apply the clear color to the color buffers with the command

\[ \text{glClear} \left( \text{GL\_COLOR\_BUFFER\_BIT} \right); \]

We can also use the \text{glClear} function to set initial values for other buffers that are available in OpenGL. These are the accumulation buffer, which stores blended-color information, the depth buffer, which stores depth values (distances from the viewing position) for objects in a scene, and the stencil buffer, which stores information to define the limits of a picture.

In color-index mode, we use the following function (instead of \text{glClearColor}) to set the display-window color:

\[ \text{glClearIndex} \left( \text{index} \right); \]

The window background color is then assigned the color that is stored at position \text{index} in the color table; and the window is displayed in this color when we issue the \text{glClear} \left( \text{GL\_COLOR\_BUFFER\_BIT} \right) function.

Many other color functions are available in the OpenGL library for dealing with a variety of tasks, such as changing color models, setting lighting effects for a scene, specifying camera effects, and rendering the surfaces of an object. We examine other color functions as we explore each of the component processes in a computer-graphics system. For now, we limit our discussion to those functions relating to color specifications for graphics primitives.
4 Point Attributes

Basically, we can set two attributes for points: color and size. In a state system, the displayed color and size of a point is determined by the current values stored in the attribute list. Color components are set with RGB values or an index into a color table. For a raster system, point size is an integer multiple of the pixel size, so that a large point is displayed as a square block of pixels.

5 OpenGL Point-Attribute Functions

The displayed color of a designated point position is controlled by the current color values in the state list. Also, a color is specified with either the glColor function or the glIndex function.

We set the size for an OpenGL point with

\[ \text{glPointSize} \ (\text{size}); \]

and the point is then displayed as a square block of pixels. Parameter size is assigned a positive floating-point value, which is rounded to an integer (unless the point is to be anti-aliased). The number of horizontal and vertical pixels in the display of the point is determined by parameter size. Thus, a point size of 1.0 displays a single pixel, and a point size of 2.0 displays a 2 × 2 pixel array. If we activate the anti-aliasing features of OpenGL, the size of a displayed block of pixels will be modified to smooth the edges. The default value for point size is 1.0.

Attribute functions may be listed inside or outside of a glBegin/glEnd pair. For example, the following code segment plots three points in varying colors and sizes. The first is a standard-size red point, the second is a double-size green point, and the third is a triple-size blue point:

\[
\text{glColor3f} \ (1.0, \ 0.0, \ 0.0); \\
\text{glBegin} \ (\text{GL_POINTS}); \\
\text{glVertex2i} \ (50, \ 100); \\
\text{glPointSize} \ (2.0); \\
\text{glColor3f} \ (0.0, \ 1.0, \ 0.0); \\
\text{glVertex2i} \ (75, \ 150); \\
\text{glPointSize} \ (3.0); \\
\text{glColor3f} \ (0.0, \ 0.0, \ 1.0); \\
\text{glVertex2i} \ (100, \ 200); \\
\text{glEnd} \ ( );
\]

6 Line Attributes

A straight-line segment can be displayed with three basic attributes: color, width, and style. Line color is typically set with the same function for all graphics primitives, while line width and line style are selected with separate line functions. In addition, lines may be generated with other effects, such as pen and brush strokes.

Line Width

Implementation of line-width options depends on the capabilities of the output device. A heavy line could be displayed on a video monitor as adjacent parallel lines, while a pen plotter might require pen changes to draw a thick line.
Attributes of Graphics Primitives

For raster implementations, a standard-width line is generated with single pixels at each sample position, as in the Bresenham algorithm. Thicker lines are displayed as positive integer multiples of the standard line by plotting additional pixels along adjacent parallel line paths.

Line Style
Possible selections for the line-style attribute include solid lines, dashed lines, and dotted lines. We modify a line-drawing algorithm to generate such lines by setting the length and spacing of displayed solid sections along the line path. With many graphics packages, we can select the length of both the dashes and the inter-dash spacing.

Pen and Brush Options
With some packages, particularly painting and drawing systems, we can select different pen and brush styles directly. Options in this category include shape, size, and pattern for the pen or brush. Some example pen and brush shapes are given in Figure 2.

![Custom Document Brushes](image)

**Figure 2**
Pen and brush shapes for line display.
7 OpenGL Line-Attribute Functions

We can control the appearance of a straight-line segment in OpenGL with three attribute settings: line color, line width, and line style. We have already seen how to make a color selection, and OpenGL provides a function for setting the width of a line and another function for specifying a line style, such as a dashed or dotted line.

OpenGL Line-Width Function

Line width is set in OpenGL with the function

```
    glLineWidth (width);
```

We assign a floating-point value to parameter `width`, and this value is rounded to the nearest nonnegative integer. If the input value rounds to 0.0, the line is displayed with a standard width of 1.0, which is the default width. However, when antialiasing is applied to the line, its edges are smoothed to reduce the raster stair-step appearance and fractional widths are possible. Some implementations of the line-width function might support only a limited number of widths, and some might not support widths other than 1.0.

The magnitude of the horizontal and vertical separations of the line endpoints, \( \Delta x \) and \( \Delta y \), are compared to determine whether to generate a thick line using vertical pixel spans or horizontal pixel spans.

OpenGL Line-Style Function

By default, a straight-line segment is displayed as a solid line. However, we can also display dashed lines, dotted lines, or a line with a combination of dashes and dots, and we can vary the length of the dashes and the spacing between dashes or dots. We set a current display style for lines with the OpenGL function

```
    glLineStipple (repeatFactor, pattern);
```

Parameter `pattern` is used to reference a 16-bit integer that describes how the line should be displayed. A 1 bit in the pattern denotes an “on” pixel position, and a 0 bit indicates an “off” pixel position. The pattern is applied to the pixels along the line path starting with the low-order bits in the pattern. The default pattern is 0xFFFF (each bit position has a value of 1), which produces a solid line. Integer parameter `repeatFactor` specifies how many times each bit in the pattern is to be repeated before the next bit in the pattern is applied. The default repeat value is 1.

With a polyline, a specified line-style pattern is not restarted at the beginning of each segment. It is applied continuously across all the segments, starting at the first endpoint of the polyline and ending at the final endpoint for the last segment in the series.

As an example of specifying a line style, suppose that parameter `pattern` is assigned the hexadecimal representation 0x00FF and the repeat factor is 1. This would display a dashed line with eight pixels in each dash and eight pixel positions that are “off” (an eight-pixel space) between two dashes. Also, because low-order bits are applied first, a line begins with an eight-pixel dash starting at the first endpoint. This dash is followed by an eight-pixel space, then another eight-pixel dash, and so forth, until the second endpoint position is reached.
Attributes of Graphics Primitives

Before a line can be displayed in the current line-style pattern, we must activate the line-style feature of OpenGL. We accomplish this with the following function:

```c
glEnable (GL_LINE_STIPPLE);
```

If we forget to include this enable function, solid lines are displayed; that is, the default pattern 0xFFFF is used to display line segments. At any time, we can turn off the line-pattern feature with

```c
glDisable (GL_LINE_STIPPLE);
```

This replaces the current line-style pattern with the default pattern (solid lines).

In the following program outline, we illustrate use of the OpenGL line-attribute functions by plotting three line graphs in different styles and widths. Figure 3 shows the data plots that could be generated by this program.

```c
/* Define a two-dimensional world-coordinate data type. */
typedef struct { float x, y; } wcPt2D;

wcPt2D dataPts [5];

void linePlot (wcPt2D dataPts [5])
{
    int k;

    glBegin (GL_LINE_STRIP);
    for (k = 0; k < 5; k++)
    {
        glVertex2f (dataPts [k].x, dataPts [k].y);
    }
    glFlush ( );
    glEnd ( );

    /* Invoke a procedure here to draw coordinate axes. */
    glEnable (GL_LINE_STIPPLE);

    /* Input first set of (x, y) data values. */
    glLineStipple (1, 0x1C47); // Plot a dash-dot, standard-width polyline.
    linePlot (dataPts);

    /* Input second set of (x, y) data values. */
    glLineStipple (1, 0x00FF); // Plot a dashed, double-width polyline.
    glLineWidth (2.0);
    linePlot (dataPts);

    /* Input third set of (x, y) data values. */
    glLineStipple (1, 0x0101); // Plot a dotted, triple-width polyline.
    glLineWidth (3.0);
    linePlot (dataPts);
    glDisable (GL_LINE_STIPPLE);
```
Other OpenGL Line Effects

In addition to specifying width, style, and a solid color, we can display lines with color gradations. For example, we can vary the color along the path of a solid line by assigning a different color to each line endpoint as we define the line. In the following code segment, we illustrate this by assigning a blue color to one endpoint of a line and a red color to the other endpoint. The solid line is then displayed as a linear interpolation of the colors at the two endpoints:

```gl
void main()
{
  glShadeModel (GL_SMOOTH);
  glBegin (GL_LINES);
  glColor3f (0.0, 0.0, 1.0);
  glVertex2i (50, 50);
  glColor3f (1.0, 0.0, 0.0);
  glVertex2i (250, 250);
  glEnd ( );
}
```

Function `glShadeModel` can also be given the argument `GL_FLAT`. In that case, the line segment would have been displayed in a single color: the color of the second endpoint, (250, 250). That is, we would have generated a red line. Actually, `GL_SMOOTH` is the default, so we would generate a smoothly interpolated color line segment even if we did not include this function in our code.

We can produce other effects by displaying adjacent lines that have different colors and patterns. In addition, we can use the color-blending features of OpenGL by superimposing lines or other objects with varying alpha values. A brush stroke and other painting effects can be simulated with a pixmap and color blending. The pixmap can then be moved interactively to generate line segments. Individual pixels in the pixmap can be assigned different alpha values to display lines as brush or pen strokes.

8 Curve Attributes

Parameters for curve attributes are the same as those for straight-line segments. We can display curves with varying colors, widths, dot-dash patterns, and available pen or brush options. For adapting curve-drawing algorithms to accommodate attribute selections are similar to those for line drawing.
Painting and drawing programs allow pictures to be constructed interactively by using a pointing device, such as a stylus and a graphics tablet, to sketch various curve shapes. Some examples of such curve patterns are shown in Figure 4. An additional pattern option that can be provided in a paint package is the display of simulated brush strokes.

Strictly speaking, OpenGL does not consider curves to be drawing primitives in the same way that it considers points and lines to be primitives. Curves can be drawn in several ways in OpenGL. Perhaps the simplest approach is to approximate the shape of the curve using short line segments. Alternatively, curved segments can be drawn using splines. These can be drawn using OpenGL evaluator functions, or by using functions from the OpenGL Utility (GLU) library which draw splines.

### 9 Fill-Area Attributes

Most graphics packages limit fill areas to polygons because they are described with linear equations. A further restriction requires fill areas to be convex polygons, so that scan lines do not intersect more than two boundary edges. However, in general, we can fill any specified regions, including circles, ellipses, and other objects with curved boundaries. Also, application systems, such as paint programs, provide fill options for arbitrarily shaped regions.

#### Fill Styles

A basic fill-area attribute provided by a general graphics library is the display style of the interior. We can display a region with a single color, a specified fill pattern, or in a “hollow” style by showing only the boundary of the region. These three fill styles are illustrated in Figure 5. We can also fill selected regions of a scene using various brush styles, color-blending combinations, or textures. Other options include specifications for the display of the boundaries of a fill area. For polygons, we could show the edges in different colors, widths, and styles; and we can select different display attributes for the front and back faces of a region.

Fill patterns can be defined in rectangular color arrays that list different colors for different positions in the array. Alternatively, a fill pattern could be specified as a bit array that indicates which relative positions are to be displayed in a single selected color. An array specifying a fill pattern is a mask that is to be applied to the display area. Some graphics systems provide an option for selecting an arbitrary initial position for overlaying the mask. From this starting position, the mask is replicated in the horizontal and vertical directions until the display area is filled with nonoverlapping copies of the pattern. Where the pattern overlaps specified fill areas, the array pattern indicates which pixels should be displayed in a particular color. This process of filling an area with a rectangular pattern is called tiling, and a rectangular fill pattern is sometimes referred to as a tiling pattern. Sometimes, predefined fill patterns are available in a system, such as the hatch fill patterns shown in Figure 6.
Color-Blended Fill Regions

It is also possible to combine a fill pattern with background colors in various ways. A pattern could be combined with background colors using a transparency factor that determines how much of the background should be mixed with the object color.

Some fill methods using blended colors have been referred to as soft-fill or tint-fill algorithms. One use for these fill methods is to soften the fill colors at object borders that have been blurred to antialias the edges. Another application of a soft-fill algorithm is to allow repainting of a color area that was originally filled with a semitransparent brush, where the current color is then a mixture of the brush color and the background colors “behind” the area. In either case, we want the new fill color to have the same variations over the area as the current fill color.

10 OpenGL Fill-Area Attribute Functions

In the OpenGL graphics package, fill-area routines are available for convex polygons only. We generate displays of filled convex polygons in four steps:

1. Define a fill pattern.
2. Invoke the polygon-fill routine.
3. Activate the polygon-fill feature of OpenGL.
4. Describe the polygons to be filled.

A polygon fill pattern is displayed up to and including the polygon edges. Thus, there are no boundary lines around the fill region unless we specifically add them to the display.

In addition to specifying a fill pattern for a polygon interior, there are a number of other options available. One option is to display a hollow polygon, where no interior color or pattern is applied and only the edges are generated. A hollow polygon is equivalent to the display of a closed polyline primitive. Another option is to show the polygon vertices, with no interior fill and no edges. Also, we designate different attributes for the front and back faces of a polygon fill area.

OpenGL Fill-Pattern Function

By default, a convex polygon is displayed as a solid-color region, using the current color setting. To fill the polygon with a pattern in OpenGL, we use a \(32 \times 32\) bit mask. A value of 1 in the mask indicates that the corresponding pixel is to be set to the current color, and a 0 leaves the value of that frame-buffer position unchanged. The fill pattern is specified in unsigned bytes using the OpenGL data type `GLubyte`, just as we did with the `glBitmap` function. We define a bit pattern with hexadecimal values as, for example,

```c
GLubyte fillPattern[ ] = { 0xff, 0x00, 0xff, 0x00, ... };
```

The bits must be specified starting with the bottom row of the pattern, and continuing up to the topmost row (32) of the pattern. This pattern is replicated...
Once we have set a mask, we can establish it as the current fill pattern with the function

\[
glPolygonStipple(\text{fillPattern});
\]

Next, we need to enable the fill routines before we specify the vertices for the polygons that are to be filled with the current pattern. We do this with the statement

\[
glEnable(\text{GL_POLYGON_STIPPLE});
\]

Similarly, we turn off pattern filling with

\[
glDisable(\text{GL_POLYGON_STIPPLE});
\]

Figure 8 illustrates how a $3 \times 3$ bit pattern, repeated over a $32 \times 32$ bit mask, might be applied to fill a parallelogram.

OpenGL Texture and Interpolation Patterns

Another method for filling polygons is to use texture patterns. This can produce fill patterns that simulate the surface appearance of wood, brick, brushed steel, or some other material. Also, we can obtain an interpolation coloring of a polygon interior just as we did with the line primitive. To do this, we assign different colors to polygon vertices. Interpolation fill of a polygon interior is used to produce realistic displays of shaded surfaces under various lighting conditions.
As an example of an interpolation fill, the following code segment assigns either a blue, red, or green color to each of the three vertices of a triangle. The polygon fill is then a linear interpolation of the colors at the vertices:

```c
glShadeModel (GL_SMOOTH);

glBegin (GL_TRIANGLES);
    glColor3f (0.0, 0.0, 1.0);
    glVertex2i (50, 50);
    glColor3f (1.0, 0.0, 0.0);
    glVertex2i (150, 50);
    glColor3f (0.0, 1.0, 0.0);
    glVertex2i (75, 150);

glEnd ( );
```

Of course, if a single color is set for the triangle as a whole, the polygon is filled with that one color; and if we change the argument in the `glShadeModel` function to `GL_FLAT` in this example, the polygon is filled with the last color specified (green). The value `GL_SMOOTH` is the default shading, but we can include that specification to remind us that the polygon is to be filled as an interpolation of the vertex colors.

### OpenGL Wire-Frame Methods

We can also choose to show only polygon edges. This produces a wire-frame or hollow display of the polygon; or we could display a polygon by plotting a set of points only at the vertex positions. These options are selected with the function

```c
glPolygonMode (face, displayMode);
```

We use parameter `face` to designate which face of the polygon that we want to show as edges only or vertices only. This parameter is then assigned either `GL_FRONT`, `GL_BACK`, or `GL_FRONT_AND_BACK`. Then, if we want only the polygon edges displayed for our selection, we assign the constant `GL_LINE` to parameter `displayMode`. To plot only the polygon vertex points, we assign the constant `GL_POINT` to parameter `displayMode`. A third option is `GL_FILL`; but this is the default display mode, so we usually invoke only `glPolygonMode` when we want to set attributes for the polygon edges or vertices.

Another option is to display a polygon with both an interior fill and a different color or pattern for its edges (or for its vertices). This is accomplished by specifying the polygon twice: once with parameter `displayMode` set to `GL_FILL` and then again with `displayMode` set to `GL_LINE` (or `GL_POINT`). For example, the following code section fills a polygon interior with a green color, and then the edges are assigned a red color:

```c
    glColor3f (0.0, 1.0, 0.0);
    /* Invoke polygon-generating routine. */

    glColor3f (1.0, 0.0, 0.0);
    glPolygonMode (GL_FRONT, GL_LINE);
    /* Invoke polygon-generating routine again. */
```

For a three-dimensional polygon (one that does not have all vertices in the xy plane), this method for displaying the edges of a filled polygon may produce gaps along the edges. This effect, sometimes referred to as **stitching**, is caused by...
Attributes of Graphics Primitives

differences between calculations in the scan-line fill algorithm and calculations in the edge line-drawing algorithm. As the interior of a three-dimensional polygon is filled, the depth value (distance from the xy plane) is calculated for each (x, y) position. However, this depth value at an edge of the polygon is often not exactly the same as the depth value calculated by the line-drawing algorithm for the same (x, y) position. Therefore, when visibility tests are made, the interior fill color could be used instead of an edge color to display some points along the boundary of a polygon.

One way to eliminate the gaps along displayed edges of a three-dimensional polygon is to shift the depth values calculated by the fill routine so that they do not overlap with the edge depth values for that polygon. We do this with the following two OpenGL functions:

```c
 glEnable (GL_POLYGON_OFFSET_FILL);
 glPolygonOffset (factor1, factor2);
```

The first function activates the offset routine for scan-line filling, and the second function is used to set a couple of floating-point values factor1 and factor2 that are used to calculate the amount of depth offset. The calculation for this depth offset is

\[
\text{depthOffset} = \text{factor1} \cdot \text{maxSlope} + \text{factor2} \cdot \text{const}
\]

where maxSlope is the maximum slope of the polygon and const is an implementation constant. For a polygon in the xy plane, the slope is 0. Otherwise, the maximum slope is calculated as the change in depth of the polygon divided by either the change in x or the change in y. A typical value for the two factors is either 0.75 or 1.0, although some experimentation with the factor values is often necessary to produce good results. As an example of assigning values to offset factors, we can modify the previous code segment as follows:

```c
 glColor3f (0.0, 1.0, 0.0);
 glEnable (GL_POLYGON_OFFSET_FILL);
 glPolygonOffset (1.0, 1.0); /* Invoke polygon-generating routine. */
 glDisable (GL_POLYGON_OFFSET_FILL);
 glColor3f (1.0, 0.0, 0.0);
 glPolygonMode (GL_FRONT, GL_LINE); /* Invoke polygon-generating routine again. */
```

Now the interior fill of the polygon is pushed a little farther away in depth, so that it does not interfere with the depth values of its edges. It is also possible to implement this method by applying the offset to the line-drawing algorithm, by changing the argument of the glEnable function to GL_POLYGON_OFFSET_LINE. In this case, we want to use negative factors to bring the edge depth values closer. Also, if we just wanted to display different color points at the vertex positions, instead of highlighted edges, the argument in the glEnable function would be GL_POLYGON_OFFSET_POINT.

Another method for eliminating the stitching effect along polygon edges is to use the OpenGL stencil buffer to limit the polygon interior filling so that it does not overlap the edges. However, this approach is more complicated and generally slower, so the polygon depth-offset method is preferred.
To display a concave polygon using OpenGL routines, we must first split it into a set of convex polygons. We typically divide a concave polygon into a set of triangles. Then we could display the concave polygon as a fill region by filling the triangles. Similarly, if we want to show only the polygon vertices, we plot the triangle vertices. To display the original concave polygon in a wire-frame form, however, we cannot just set the display mode to GL_LINE because that would show all the triangle edges that are interior to the original concave polygon (Figure 9).

Fortunately, OpenGL provides a mechanism that allows us to eliminate selected edges from a wire-frame display. Each polygon vertex is stored with a one-bit flag that indicates whether or not that vertex is connected to the next vertex by a boundary edge. So all we need do is set that bit flag to “off” and the edge following that vertex will not be displayed. We set this flag for an edge with the following function:

```c
glEdgeFlag (flag);
```

To indicate that a vertex does not precede a boundary edge, we assign the OpenGL constant GL_FALSE to parameter flag. This applies to all subsequently specified vertices until the next call to glEdgeFlag is made. The OpenGL constant GL_TRUE turns the edge flag on again, which is the default. Function glEdgeFlag can be placed between glBegin/glEnd pairs. As an illustration of the use of an edge flag, the following code displays only two edges of the defined triangle (Figure 10):

```c
g1PolygonMode (GL_FRONT_AND_BACK, GL_LINE);

g1Begin (GL_POLYGON);
    g1Vertex3fv (v1);
    glEdgeFlag (GL_FALSE);
    g1Vertex3fv (v2);
    glEdgeFlag (GL_TRUE);
    g1Vertex3fv (v3);
    glEnd ()
```
Polygon edge flags can also be specified in an array that could be combined or associated with a vertex array (see Section 3). The statements for creating an array of edge flags are

```c
glEnableClientState (GL_EDGE_FLAG_ARRAY);

glEdgeFlagPointer (offset, edgeFlagArray);
```

Parameter `offset` indicates the number of bytes between the values for the edge flags in the array `edgeFlagArray`. The default value for parameter `offset` is 0.

**OpenGL Front-Face Function**

Although, by default, the ordering of polygon vertices controls the identification of front and back faces, we can label selected surfaces in a scene independently as front or back with the function

```c
glFrontFace (vertexOrder);
```

If we set parameter `vertexOrder` to the OpenGL constant `GL_CW`, then a subsequently defined polygon with a clockwise ordering for its vertices is considered to be front-facing. This OpenGL feature can be used to swap faces of a polygon for which we have specified vertices in a clockwise order. The constant `GL_CCW` labels a counterclockwise ordering of polygon vertices as front-facing, which is the default ordering.

## 11 Character Attributes

We control the appearance of displayed characters with attributes such as font, size, color, and orientation. In many packages, attributes can be set both for entire character strings (text) and for individual characters that can be used for special purposes such as plotting a data graph.

There are a great many possible text-display options. First, there is the choice of font (or typeface), which is a set of characters with a particular design style such as New York, Courier, Helvetica, London, Times Roman, and various special symbol groups. The characters in a selected font can also be displayed with assorted underlining styles (solid, dotted, double), in boldface, in italic, and in OUTLINE or shadow styles.

Color settings for displayed text can be stored in the system attribute list and used by the procedures that generate character definitions in the frame buffer. When a character string is to be displayed, the current color is used to set pixel values in the frame buffer corresponding to the character shapes and positions.

We could adjust text size by scaling the overall dimensions (height and width) of characters or by scaling only the height or the width. Character size (height) is specified by printers and compositors in points, where 1 point is about 0.035146 centimeters (or 0.013837 inch, which is approximately \( \frac{1}{72} \) inch). For example, the characters in this book are set in a 10-point font. Point measurements specify the size of the body of a character (Figure 11), but different fonts with the same point specifications can have different character sizes, depending on the design of the typeface. The distance between the bottomline and the topline of the character body is the same for all characters in a particular size and typeface, but the body width may vary. Proportionally spaced fonts assign a smaller body width to narrow characters such as \( i, j, l \), and \( f \) compared to broad characters.
such as \( W \) or \( M \). Character height is defined as the distance between the baseline and the capline of characters. Kerned characters, such as \( f \) and \( j \) in Figure 11, typically extend beyond the character body limits, and letters with descenders (\( g, j, p, q, y \)) extend below the baseline. Each character is positioned within the character body by a font designer in such a way that suitable spacing is attained along and between print lines when text is displayed with character bodies touching.

Sometimes, text size is adjusted without changing the width-to-height ratio of characters. Figure 12 shows a character string displayed with three different character heights, while maintaining the ratio of width to height. Examples of text displayed with a constant height and varying widths are given in Figure 13.

Spacing between characters is another attribute that can often be assigned to a character string. Figure 14 shows a character string displayed with three different settings for the intercharacter spacing.

The orientation for a character string can be set according to the direction of a character up vector. Text is then displayed so that the orientation of characters from baseline to capline is in the direction of the up vector. For example, with the direction of the up vector at 45°, text would be displayed as shown in Figure 15. A procedure for orienting text could rotate characters so that the sides of character bodies, from baseline to capline, are aligned with the up vector. The rotated character shapes are then scan converted into the frame buffer.

It is useful in many applications to be able to arrange character strings vertically or horizontally. Examples of this are given in Figure 16. We could also arrange the characters in a text string so that the string is displayed forward or backward. Examples of text displayed with these options are shown in Figure 17. A procedure for implementing text-path orientation adjusts the position of the individual characters in the frame buffer according to the option selected.

Character strings could also be oriented using a combination of up-vector and text-path specifications to produce slanted text. Figure 18 shows the directions...
Attributes of Graphics Primitives

**FIGURE 15**
Direction of the up vector (a) controls the orientation of displayed text (b).

**FIGURE 16**
Text-path attributes can be set to produce horizontal or vertical arrangements of character strings.

**FIGURE 17**
A text string displayed with the four text-path options: left, right, up, and down.

**FIGURE 18**
An up-vector specification (a) and associated directions for the text path (b).

**FIGURE 19**
The $45^\circ$ up vector in Figure 18 produces the display (a) for a down path and the display (b) for a right path.

of character strings generated by various text path settings for a $45^\circ$ up vector. Examples of character strings generated for text-path values down and right with this up vector are illustrated in Figure 19.

Another possible attribute for character strings is alignment. This attribute specifies how text is to be displayed with respect to a reference position. For example, individual characters could be aligned according to the base lines or the character centers. Figure 20 illustrates typical character positions for horizontal and vertical alignments. String alignments are also possible, and Figure 21 shows common alignment positions for horizontal and vertical text labels.

In some graphics packages, a text-precision attribute is also available. This parameter specifies the amount of detail and the particular processing options that are to be used with a text string. For a low-precision text string, many attribute selections, such as text path, are ignored, and faster procedures are used for processing the characters through the viewing pipeline.

Finally, a library of text-processing routines often supplies a set of special characters, such as a small circle or cross, which are useful in various applications. Most
often these characters are used as marker symbols in network layouts or in graphing data sets. The attributes for these marker symbols are typically color and size.

### 12 OpenGL Character-Attribute Functions

We have two methods for displaying characters with the OpenGL package. Either we can design a font set using the bitmap functions in the core library, or we can invoke the GLUT character-generation routines. The GLUT library contains functions for displaying predefined bitmap and stroke character sets. Therefore, the character attributes we can set are those that apply to either bitmaps or line segments.

For either bitmap or outline fonts, the display color is determined by the current color state. In general, the spacing and size of characters is determined by the font designation, such as GLUT_BITMAP_9_BY_15 and GLUT_STROKE_MONO_ROMAN. However, we can also set the line width and line type for the outline fonts. We specify the width for a line with the `glLineWidth` function, and we select a line type with the `glLineStipple` function. The GLUT stroke fonts will then be displayed using the current values we specified for the OpenGL line-width and line-type attributes.

We can accomplish some other text-display characteristics using transformation functions. The transformation routines allow us to scale, position, and rotate the GLUT stroke characters in either two-dimensional space or three-dimensional space. In addition, the three-dimensional viewing transformations can be used to generate other display effects.

### 13 OpenGL Antialiasing Functions

Line segments and other graphics primitives generated by raster algorithms have a jagged, or stair-step, appearance because the sampling process digitizes coordinate points on an object to discrete integer pixel positions. This distortion of information due to low-frequency sampling (undersampling) is called aliasing. We can improve the appearance of displayed raster lines by applying antialiasing methods that compensate for the undersampling process.

OpenGL provides antialiasing support for three types of primitives. We activate the antialiasing routines with the function:

```c
glEnable (primitiveType);
```

where parameter `primitiveType` is assigned one of the symbolic constant values GL_POINT_SMOOTH, GL_LINE_SMOOTH, or GL_POLYGON_SMOOTH. Assuming that we are specifying color values using the RGBA mode, we also need to activate the OpenGL color-blending operations as follows:

```c
glEnable (GL_BLEND);
```

Next, we apply the color-blending method described in Section 3 using the function

```c
glBlendFunc (GL_SRC_ALPHA, GL_ONE_MINUS_SRC_ALPHA);
```

The smoothing operations are more effective if we use large alpha values in the color specifications for the objects.
Antialiasing can also be applied when we use color tables. However, in this color mode, we must create a color ramp, which is a table of color graduations from the background color to the object color. This color ramp is then used to antialias object boundaries.

### 14 OpenGL Query Functions

We can retrieve current values for any of the state parameters, including attribute settings, using OpenGL query functions. These functions copy specified state values into an array, which we can save for later reuse or to check the current state of the system if an error occurs.

For current attribute values we use an appropriate “glGet” function, such as

```
glGetBooleanv ( )    glGetFloatv ( )
glGetIntegerv ( )    glGetDoublev ( )
```

In each of the preceding functions, we specify two arguments. The first argument is an OpenGL symbolic constant that identifies an attribute or other state parameter. The second argument is a pointer to an array of the data type indicated by the function name. For instance, we can retrieve the current RGBA floating-point color settings with

```
glGetFloatv (GL_CURRENT_COLOR, colorValues);
```

The current color components are then passed to the array colorValues. To obtain the integer values for the current color components, we invoke the glGetIntegerv function. In some cases, a type conversion may be necessary to return the specified data type.

Other OpenGL constants, such as GL_POINT_SIZE, GL_LINE_WIDTH, and GL_CURRENT_RASTER_POSITION, can be used in these functions to return current state values. Also, we could check the range of point sizes or line widths that are supported using the constants GL_POINT_SIZE_RANGE and GL_LINE_WIDTH_RANGE.

Although we can retrieve and reuse settings for a single attribute with the glGet functions, OpenGL provides other functions for saving groups of attributes and reusing their values. We consider the use of these functions for saving current attribute settings in the next section.

There are many other state and system parameters that are often useful to query. For instance, to determine how many bits per pixel are provided in the frame buffer on a particular system, we can ask the system how many bits are available for each individual color component, such as

```
glGetIntegerv (GL_RED_BITS, redBitSize);
```

Here, array redBitSize is assigned the number of red bits available in each of the buffers (frame buffer, depth buffer, accumulation buffer, and stencil buffer). Similarly, we can make an inquiry for the other color bits using GL_GREEN_BITS, GL_BLUE_BITS, GL_ALPHA_BITS, or GL_INDEX_BITS.

We can also find out whether edge flags have been set, whether a polygon face was tagged as a front face or a back face, and whether the system supports double buffering. In addition, we can inquire whether certain routines, such as color blending, line stippling or antialiasing, have been enabled or disabled.
15 OpenGL Attribute Groups

Attributes and other OpenGL state parameters are arranged in attribute groups. Each group contains a set of related state parameters. For instance, the point-attribute group contains the size and point-smooth (antialiasing) parameters, and the line-attribute group contains the width, stipple status, stipple pattern, stipple repeat counter, and line-smooth status. Similarly, the polygon-attribute group contains eleven polygon parameters, such as fill pattern, front-face flag, and polygon-smooth status. Because color is an attribute for all primitives, it has its own attribute group; and some parameters are included in more than one group.

About twenty different attribute groups are available in OpenGL, and all parameters in one or more groups can be saved or reset with a single function. We save all parameters within a specified group using the following command:

```c
glPushAttrib (attrGroup);
```

Parameter `attrGroup` is assigned an OpenGL symbolic constant that identifies an attribute group, such as `GL_POINT_BIT`, `GL_LINE_BIT`, or `GL_POLYGON_BIT`. To save color parameters, we use the symbolic constant `GL_CURRENT_BIT`, and we can save all state parameters in all attribute groups with the constant `GL_ALL_ATTRIB_BITS`. The `glPushAttrib` function places all parameters within the specified group onto an attribute stack.

We can also save parameters within two or more groups by combining their symbolic constants with a logical OR operation. The following statement places all parameters for points, lines, and polygons on the attribute stack:

```c
glPushAttrib (GL_POINT_BIT | GL_LINE_BIT | GL_POLYGON_BIT);
```

Once we have saved a group of state parameters, we can reinstate all values on the attribute stack with this function:

```c
glPopAttrib ( );
```

No arguments are used in the `glPopAttrib` function because it resets the current state of OpenGL using all values on the stack.

These commands for saving and resetting state parameters use a server attribute stack. There is also a client attribute stack available in OpenGL for saving and resetting client state parameters. The functions for accessing this stack are `glPushClientAttrib` and `glPopClientAttrib`. Only two client attribute groups are available: one for pixel-storage modes and the other for vertex arrays. Pixel-storage parameters include information such as byte alignment and the type of arrays used to store subimages of a display. Vertex-array parameters give information about the current vertex-array state, such as the enable/disable state of various arrays.

16 Summary

Attributes control the display characteristics of graphics primitives. In many graphics systems, attribute values are stored as state variables and primitives are generated using the current attribute values. When we change the value of a state variable, it affects only those primitives defined after the change.

A common attribute for all primitives is color, which is most often specified in terms of RGB (or RGBA) components. The red, green, and blue color values are
Attributes of Graphics Primitives

stored in the frame buffer, and they are used to control the intensity of the three electron guns in an RGB monitor. Color selections can also be made using color-lookup tables. In this case, a color in the frame buffer is indicated as a table index, and the table location at that index stores a particular set of RGB color values. Color tables are useful in data-visualization and image-processing applications, and they can also be used to provide a wide range of colors without requiring a large frame buffer. Often, computer-graphics packages provide options for using either color tables or storing color values directly in the frame buffer.

The basic point attributes are color and size. Line attributes are color, width, and style. Specifications for line width are given in terms of multiples of a standard, one-pixel-wide line. The line-style attributes include solid, dashed, and dotted lines, as well as various brush or pen styles. These attributes can be applied to both straight lines and curves.

Fill-area attributes include a solid-color fill, a fill pattern, and a hollow display that shows only the area boundaries. Various pattern fills can be specified in color arrays, which are then mapped to the interior of the region. Scan-line methods are commonly used to fill polygons, circles, and ellipses.

Areas can also be filled using color blending. This type of fill has applications in antialiasing and in painting packages. Soft-fill procedures provide a new fill color for a region that has the same variations as the previous fill color.

Characters can be displayed in different styles (fonts), colors, sizes, spacing, and orientations. To set the orientation of a character string, we can specify a direction for the character up vector and a direction for the text path. In addition, we can set the alignment of a text string in relation to the start coordinate position. Individual characters, called marker symbols, can be used for applications such as plotting data graphs. Marker symbols can be displayed in various sizes and colors using standard characters or special symbols.

Because scan conversion is a digitizing process on raster systems, displayed primitives have a jagged appearance. This is due to the undersampling of information, which rounds coordinate values to pixel positions. We can improve the appearance of raster primitives by applying antialiasing procedures that adjust pixel intensities.

In OpenGL, attribute values for the primitives are maintained as state variables. An attribute setting remains in effect for all subsequently defined primitives until that attribute value is changed. Changing an attribute value does not affect previously displayed primitives. We can specify colors in OpenGL using either the RGB (or RGBA) color mode or the color-index mode, which uses color-table indices to select colors. Also, we can blend color values using the alpha color component, and we can specify values in color arrays that are to be used in conjunction with vertex arrays. In addition to color, OpenGL provides functions for selecting point size, line width, line style, and convex-polygon fill style, as well as providing functions for the display of polygon fill areas as either a set of edges or a set of vertex points. We can also eliminate selected polygon edges from a display, and we can reverse the specification of front and back faces. We can generate text strings in OpenGL using bitmaps or routines that are available in GLUT. Attributes that can be set for the display of GLUT characters include color, font, size, spacing, line width, and line type. The OpenGL library also provides functions to antialias the display of output primitives. We can use query functions to obtain the current value for state variables, and we can also obtain all values within an OpenGL attribute group using a single function.

Table 2 summarizes the OpenGL attribute functions discussed in this chapter. In addition, the table lists some attribute-related functions.
### Summary of OpenGL Attribute Functions

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glutInitDisplayMode</td>
<td>Selects the color mode, which can be either GLUT_RGB or GLUT_INDEX.</td>
</tr>
<tr>
<td>glColor*</td>
<td>Specifies an RGB or RGBA color.</td>
</tr>
<tr>
<td>glIndex*</td>
<td>Specifies a color using a color-table index.</td>
</tr>
<tr>
<td>glutSetColor (index, r, g, b);</td>
<td>Loads a color into a color-table position.</td>
</tr>
<tr>
<td>glEnable (GL_BLEND);</td>
<td>Activates color blending.</td>
</tr>
<tr>
<td>blendFunc (sFact, dFact);</td>
<td>Specifies factors for color blending.</td>
</tr>
<tr>
<td>glEnableClientState (GL_COLOR_ARRAY);</td>
<td>Activates color-array features of OpenGL.</td>
</tr>
<tr>
<td>glColorPointer (size, type, stride, array);</td>
<td>Specifies an RGB color array.</td>
</tr>
<tr>
<td>glIndexPointer (type, stride, array);</td>
<td>Specifies a color array using color-index mode.</td>
</tr>
<tr>
<td>glPointSize (size)</td>
<td>Specifies a point size.</td>
</tr>
<tr>
<td>glLineWidth (width);</td>
<td>Specifies a line width.</td>
</tr>
<tr>
<td>glEnable (GL_LINE_STIPPLE);</td>
<td>Activates line style.</td>
</tr>
<tr>
<td>glEnable (GL_POLYGON_STIPPLE);</td>
<td>Activates fill style.</td>
</tr>
<tr>
<td>glLineStipple (repeat, pattern);</td>
<td>Specifies a line-style pattern.</td>
</tr>
<tr>
<td>glPolygonStipple (pattern);</td>
<td>Specifies a fill-style pattern.</td>
</tr>
<tr>
<td>glPolygonMode</td>
<td>Displays front or back face as either a set of edges or a set of vertices.</td>
</tr>
<tr>
<td>glEdgeFlag</td>
<td>Sets fill-polygon edge flag to GL_TRUE or GL_FALSE to determine display status for an edge.</td>
</tr>
<tr>
<td>glFrontFace</td>
<td>Specifies front-face vertex order as either GL_CCW or GL_CW.</td>
</tr>
<tr>
<td>glEnable</td>
<td>Activates antialiasing with GL_POINT_SMOOTH, GL_LINE_SMOOTH, or GL_POLYGON_SMOOTH. (Also need to activate color blending.)</td>
</tr>
<tr>
<td>glGet**</td>
<td>Queries OpenGL to retrieve an attribute value of a specific data type, identified by the symbolic name of the attribute, placing the result in an array parameter.</td>
</tr>
<tr>
<td>glPushAttrib</td>
<td>Saves all state parameters within a specified attribute group.</td>
</tr>
<tr>
<td>glPopAttrib ( )</td>
<td>Reinstates all state parameter values that were last saved.</td>
</tr>
</tbody>
</table>
REFERENCES


EXERCISES
1 Use the glutSetColor function to set up a color table for an input set of color values.
2 Using vertex and color arrays, set up the description for a scene containing at least six two-dimensional objects.
3 Write a program to display the two-dimensional scene description in the previous exercise.
4 Using vertex and color arrays, set up the description for a scene containing at least four three-dimensional objects.
5 Write a program to display a two-dimensional, grayscale “target” scene, where the target is made up of a small, filled central circle and two concentric rings around the circle spaced as far apart as their thickness, which should be equal to the radius of the inner circle. The circle and rings are to be described as point patterns on a white background. The rings/circle should “fade in” from their outer edges so that the inner portion of the shape is darker than the outer portion. This can be achieved by varying the sizes and inter-point spacing of the points that make up the rings/circle. For example, the edges of a ring can be modeled with small, widely spaced, light-gray points, while the inner portion can be modeled with larger, more closely spaced, dark-gray points.
6 Modify the program in the previous exercise to display the circle and rings in various shades of red instead of gray.
7 Modify the code segments in Section 7 for displaying data line plots, so that the line-width parameter is passed to procedure linePlot.
8 Modify the code segments in Section 7 for displaying data line plots, so that the line-style parameter is passed to procedure linePlot.
9 Complete the program in Section 7 for displaying line plots using input values from a data file.
10 Complete the program in Section 7 for displaying line plots using input values from a data file. In addition, the program should provide labeling for the axes and the coordinates for the display area on the screen. The data sets are to be scaled to fit the coordinate range of the display window, and each plotted line is to be displayed in a different line style, width, and color.
11 Write a routine to display a bar graph in any specified screen area. Input is to include the data set, labeling for the coordinate axes, and the coordinates for the screen area. The data set is to be scaled to fit the designated screen area, and the bars are to be displayed in designated colors or patterns.
12 Write a procedure to display two data sets defined over the same x-coordinate range, with the data values scaled to fit a specified region of the display screen. The bars for one of the data sets are to be displaced horizontally to produce an overlapping bar pattern for easy comparison of the two sets of data. Use a different color or a different fill pattern for the two sets of bars.
13 Devise an algorithm for implementing a color lookup table.
14 Suppose you have a system with an 10 inch by 14 inch video screen that can display 120 pixels per inch. If a color lookup table with 256 positions is used with this system, what is the smallest possible size (in bytes) for the frame buffer?
15 Consider an RGB raster system that has a 1024-by-786 frame buffer with 16 bits per pixel and a color lookup table with 24 bits per pixel. (a) How many distinct gray levels can be displayed with this system? (b) How many distinct colors (including gray levels) can be displayed? (c) How many colors can be displayed at any one time? (d) What is the total memory size? (e) Explain two methods for reducing memory size while maintaining the same color capabilities.
16 Write a program to output a grayscale scatter plot of two data sets defined over the same x- and y-coordinate ranges. Inputs to the program are the two sets of data. The data sets are to be scaled to fit within a defined coordinate range for a display window. Each data set is to be plotted using points in a distinct shade of gray.
17 Modify the program in the previous exercise to plot the two data sets in different colors instead of shades of gray. Also, add a legend somewhere on the plot bordered by a solid black line. The legend should display the name of each data set (given as input) in the color associated with that data set.
IN MORE DEPTH

1. Develop an application and experiment with different methods of shading the simple shapes. Using the OpenGL functions for hollow, solid color, and pattern fills of polygons, assign a fill type to each shape in the scene and apply these fills. At least one of the objects should have a hollow fill, one should be filled with a solid color, and one should be filled with a bit pattern that you specify yourself. Don’t worry if the fill patterns do not necessarily make sense for the objects in the scene. The goal here is to experiment with the different fill attributes available in OpenGL. In addition, experiment with different line drawing attributes to draw the boundaries of the shapes in your snapshot. Employ the use of solid boundary lines as well as dotted ones, each of varying thickness. Add the ability to turn anti-aliasing on and off, and examine the visual differences between the two cases.

2. Set up a small color table that serves as a color palette for your scene and draw the scene as it exists after the previous exercise using this color table instead of the standard OpenGL color functions as before. Once you produce your color table, compare its memory requirements and rendering capabilities with the standard color assignment method on your system. How many different colors can be displayed simultaneously by using the table? How much memory is saved when representing the frame buffer by using the color table instead of directly assigning colors to pixels? How small can you make the color table without noticing a significant difference in the rendering of the scene? Discuss the advantages and disadvantages to using the color table versus using direct color assignment.
In this chapter, we discuss the device-level algorithms for implementing OpenGL primitives. Exploring the implementation algorithms for a graphics library will give us valuable insight into the capabilities of these packages. It will also provide us with an understanding of how the functions work, perhaps how they could be improved, and how we might implement graphics routines ourselves for some special application. Research in computer graphics is continually discovering new and improved implementation techniques to provide us with methods for special applications, such as Internet graphics, and for developing faster and more realistic graphics displays in general.
1 Line-Drawing Algorithms

A straight-line segment in a scene is defined by the coordinate positions for the endpoints of the segment. To display the line on a raster monitor, the graphics system must first project the endpoints to integer screen coordinates and determine the nearest pixel positions along the line path between the two endpoints. Then the line color is loaded into the frame buffer at the corresponding pixel coordinates. Reading from the frame buffer, the video controller plots the screen pixels. This process digitizes the line into a set of discrete integer positions that, in general, only approximates the actual line path. A computed line position of (10.48, 20.51), for example, is converted to pixel position (10, 21). This rounding of coordinate values to integers causes all but horizontal and vertical lines to be displayed with a stair-step appearance (known as “the jaggies”), as represented in Figure 1. The characteristic stair-step shape of raster lines is particularly noticeable on systems with low resolution, and we can improve their appearance somewhat by displaying them on high-resolution systems. More effective techniques for smoothing a raster line are based on adjusting pixel intensities along the line path (see Section 15 for details).

Line Equations

We determine pixel positions along a straight-line path from the geometric properties of the line. The Cartesian slope-intercept equation for a straight line is

\[ y = m \cdot x + b \]  

with \( m \) as the slope of the line and \( b \) as the \( y \) intercept. Given that the two endpoints of a line segment are specified at positions \((x_0, y_0)\) and \((x_{\text{end}}, y_{\text{end}})\), as shown in Figure 2, we can determine values for the slope \( m \) and \( y \) intercept \( b \) with the following calculations:

\[ m = \frac{y_{\text{end}} - y_0}{x_{\text{end}} - x_0} \]  

(2)

\[ b = y_0 - m \cdot x_0 \]  

(3)

Algorithms for displaying straight lines are based on Equation 1 and the calculations given in Equations 2 and 3.

For any given \( x \) interval \( \delta x \) along a line, we can compute the corresponding \( y \) interval, \( \delta y \), from Equation 2 as

\[ \delta y = m \cdot \delta x \]  

(4)

Similarly, we can obtain the \( x \) interval \( \delta x \) corresponding to a specified \( \delta y \) as

\[ \delta x = \frac{\delta y}{m} \]  

(5)

These equations form the basis for determining deflection voltages in analog displays, such as a vector-scan system, where arbitrarily small changes in deflection voltage are possible. For lines with slope magnitudes \(|m| < 1\), \( \delta x \) can be set proportional to a small horizontal deflection voltage, and the corresponding vertical deflection is then set proportional to \( \delta y \) as calculated from Equation 4. For lines
whose slopes have magnitudes \(|m| > 1\), \(\delta y\) can be set proportional to a small vertical deflection voltage with the corresponding horizontal deflection voltage set proportional to \(\delta x\), calculated from Equation 5. For lines with \(m = 1\), \(\delta x = \delta y\) and the horizontal and vertical deflections voltages are equal. In each case, a smooth line with slope \(m\) is generated between the specified endpoints.

On raster systems, lines are plotted with pixels, and step sizes in the horizontal and vertical directions are constrained by pixel separations. That is, we must “sample” a line at discrete positions and determine the nearest pixel to the line at each sampled position. This scan-conversion process for straight lines is illustrated in Figure 3 with discrete sample positions along the \(x\) axis.

**DDA Algorithm**

The digital differential analyzer (DDA) is a scan-conversion line algorithm based on calculating either \(\delta y\) or \(\delta x\), using Equation 4 or Equation 5. A line is sampled at unit intervals in one coordinate and the corresponding integer values nearest the line path are determined for the other coordinate.

We consider first a line with positive slope, as shown in Figure 2. If the slope is less than or equal to 1, we sample at unit \(x\) intervals (\(\delta x = 1\)) and compute successive \(y\) values as

\[
y_{k+1} = y_k + m
\]

(6)

Subscript \(k\) takes integer values starting from 0, for the first point, and increases by 1 until the final endpoint is reached. Because \(m\) can be any real number between 0.0 and 1.0, each calculated \(y\) value must be rounded to the nearest integer corresponding to a screen pixel position in the \(x\) column that we are processing.

For lines with a positive slope greater than 1.0, we reverse the roles of \(x\) and \(y\). That is, we sample at unit \(y\) intervals (\(\delta y = 1\)) and compute consecutive \(x\) values as

\[
x_{k+1} = x_k + \frac{1}{m}
\]

(7)

In this case, each computed \(x\) value is rounded to the nearest pixel position along the current \(y\) scan line.

Equations 6 and 7 are based on the assumption that lines are to be processed from the left endpoint to the right endpoint (Figure 2). If this processing is reversed, so that the starting endpoint is at the right, then either we have \(\delta x = -1\) and

\[
y_{k+1} = y_k - m
\]

(8)

or (when the slope is greater than 1) we have \(\delta y = -1\) with

\[
x_{k+1} = x_k - \frac{1}{m}
\]

(9)

Similar calculations are carried out using Equations 6 through 9 to determine pixel positions along a line with negative slope. Thus, if the absolute value of the slope is less than 1 and the starting endpoint is at the left, we set \(\delta x = 1\) and calculate \(y\) values with Equation 6. When the starting endpoint is at the right (for the same slope), we set \(\delta x = -1\) and obtain \(y\) positions using Equation 8. For a negative slope with absolute value greater than 1, we use \(\delta y = -1\) and Equation 9, or we use \(\delta y = 1\) and Equation 7.

This algorithm is summarized in the following procedure, which accepts as input two integer screen positions for the endpoints of a line segment. Horizontal and vertical differences between the endpoint positions are assigned to parameters \(\delta x\) and \(\delta y\). The difference with the greater magnitude determines the value of parameter \(\text{step}\). This value is the number of pixels that must be drawn beyond the starting pixel; from it, we calculate the \(x\) and \(y\) increments needed to generate...
the next pixel position at each step along the line path. We draw the starting pixel at position \((x_0, y_0)\), and then draw the remaining pixels iteratively, adjusting \(x\) and \(y\) at each step to obtain the next pixel’s position before drawing it. If the magnitude of \(dx\) is greater than the magnitude of \(dy\) and \(x_0\) is less than \(x_{End}\), the values for the increments in the \(x\) and \(y\) directions are 1 and \(m\), respectively. If the greater change is in the \(x\) direction, but \(x_0\) is greater than \(x_{End}\), then the decrements \(-1\) and \(-m\) are used to generate each new point on the line. Otherwise, we use a unit increment (or decrement) in the \(y\) direction and an \(x\) increment (or decrement) of \(\frac{1}{m}\).

```c
#include <stdlib.h>
#include <math.h>

inline int round (const float a) { return int (a + 0.5); }

void lineDDA (int x0, int y0, int xEnd, int yEnd)
{
    int dx = xEnd - x0, dy = yEnd - y0, steps, k;
    float xIncrement, yIncrement, x = x0, y = y0;
    if (fabs (dx) > fabs (dy))
        steps = fabs (dx);
    else
        steps = fabs (dy);
    xIncrement = float (dx) / float (steps);
    yIncrement = float (dy) / float (steps);
    setPixel (round (x), round (y));
    for (k = 0; k < steps; k++) {
        x += xIncrement;
        y += yIncrement;
        setPixel (round (x), round (y));
    }
}
```

The DDA algorithm is a faster method for calculating pixel positions than one that directly implements Equation 1. It eliminates the multiplication in Equation 1 by using raster characteristics, so that appropriate increments are applied in the \(x\) or \(y\) directions to step from one pixel position to another along the line path. The accumulation of round-off error in successive additions of the floating-point increment, however, can cause the calculated pixel positions to drift away from the true line path for long line segments. Furthermore, the rounding operations and floating-point arithmetic in this procedure are still time-consuming. We can improve the performance of the DDA algorithm by separating the increments \(m\) and \(\frac{1}{m}\) into integer and fractional parts so that all calculations are reduced to integer operations. A method for calculating \(\frac{1}{m}\) increments in integer steps is discussed in Section 10. In the next section, we consider a more general scanline approach that can be applied to both lines and curves.

**Bresenham’s Line Algorithm**

In this section, we introduce an accurate and efficient raster line-generating algorithm, developed by Bresenham, that uses only incremental integer calculations. In addition, Bresenham’s line algorithm can be adapted to display circles and other curves. Figures 4 and 5 illustrate sections of a display screen where
straight-line segments are to be drawn. The vertical axes show scan-line positions, and the horizontal axes identify pixel columns. Sampling at unit \( x \) intervals in these examples, we need to decide which of two possible pixel positions is closer to the line path at each sample step. Starting from the left endpoint shown in Figure 4, we need to determine at the next sample position whether to plot the pixel at position \((11, 11)\) or the one at \((11, 12)\). Similarly, Figure 5 shows a negative-slope line path starting from the left endpoint at pixel position \((50, 50)\). In this one, do we select the next pixel position as \((51, 50)\) or as \((51, 49)\)? These questions are answered with Bresenham’s line algorithm by testing the sign of an integer parameter whose value is proportional to the difference between the vertical separations of the two pixel positions from the actual line path.

To illustrate Bresenham’s approach, we first consider the scan-conversion process for lines with positive slope less than 1.0. Pixel positions along a line path are then determined by sampling at unit \( x \) intervals. Starting from the left endpoint \((x_0, y_0)\) of a given line, we step to each successive column \( (x) \) position and plot the pixel whose scan-line \( y \) value is closest to the line path. Figure 6 demonstrates the \( k \)th step in this process. Assuming that we have determined that the pixel at \((x_k, y_k)\) is to be displayed, we next need to decide which pixel to plot in column \( x_{k+1} = x_k + 1 \). Our choices are the pixels at positions \((x_k + 1, y_k)\) and \((x_k + 1, y_k + 1)\).

At sampling position \( x_k + 1 \), we label vertical pixel separations from the mathematical line path as \( d_{\text{lower}} \) and \( d_{\text{upper}} \) (Figure 7). The \( y \) coordinate on the mathematical line at pixel column position \( x_k + 1 \) is calculated as

\[
y = mx_k + b
\]

Then

\[
d_{\text{lower}} = y - y_k = m(x_k + 1) + b - y_k
\]

and

\[
d_{\text{upper}} = (y_k + 1) - y = y_k + 1 - m(x_k + 1) - b
\]

To determine which of the two pixels is closest to the line path, we can set up an efficient test that is based on the difference between the two pixel separations as follows:

\[
d_{\text{lower}} - d_{\text{upper}} = 2m(x_k + 1) - 2y_k + 2b - 1
\]

A decision parameter \( p_k \) for the \( k \)th step in the line algorithm can be obtained by rearranging Equation 13 so that it involves only integer calculations. We accomplish this by substituting \( m = \Delta y / \Delta x \), where \( \Delta y \) and \( \Delta x \) are the vertical and horizontal separations of the endpoint positions, and defining the decision parameter as

\[
p_k = \Delta x(d_{\text{lower}} - d_{\text{upper}}) = 2\Delta y \cdot x_k - 2\Delta x \cdot y_k + c
\]

The sign of \( p_k \) is the same as the sign of \( d_{\text{lower}} - d_{\text{upper}} \), because \( \Delta x > 0 \) for our example. Parameter \( c \) is constant and has the value \( 2\Delta y + \Delta x(2b - 1) \), which is independent of the pixel position and will be eliminated in the recursive calculations for \( p_k \). If the pixel at \( y_k \) is “closer” to the line path than the pixel at \( y_k + 1 \) (that is, \( d_{\text{lower}} < d_{\text{upper}} \)), then decision parameter \( p_k \) is negative. In that case, we plot the lower pixel; otherwise, we plot the upper pixel.

\[
\begin{align*}
y_{k+1} &= y_k + 1
\end{align*}
\]

\[
\begin{align*}
y_{k+2} &= y_k + 1
\end{align*}
\]

\[
\begin{align*}
y_{k+3} &= y_k + 1
\end{align*}
\]

\[
\begin{align*}
x_k &< x_{k+1} < x_{k+2} < x_{k+3}
\end{align*}
\]
Coordinate changes along the line occur in unit steps in either the x or y direction. Therefore, we can obtain the values of successive decision parameters using incremental integer calculations. At step \( k + 1 \), the decision parameter is evaluated from Equation 14 as

\[
p_{k+1} = 2\Delta y \cdot x_{k+1} - 2\Delta x \cdot y_{k+1} + c
\]

Subtracting Equation 14 from the preceding equation, we have

\[
p_{k+1} - p_k = 2\Delta y(x_{k+1} - x_k) - 2\Delta x(y_{k+1} - y_k)
\]

However, \( x_{k+1} = x_k + 1 \), so that

\[
p_{k+1} = p_k + 2\Delta y - 2\Delta x(y_{k+1} - y_k)
\]

where the term \( y_{k+1} - y_k \) is either 0 or 1, depending on the sign of parameter \( p_k \).

This recursive calculation of decision parameters is performed at each integer position, starting at the left coordinate endpoint of the line. The first parameter, \( p_0 \), is evaluated from Equation 14 at the starting pixel position \((x_0, y_0)\) and with \( m \) evaluated as \( \Delta y/\Delta x \) as follows:

\[
p_0 = 2\Delta y - \Delta x
\]

We summarize Bresenham line drawing for a line with a positive slope less than 1 in the following outline of the algorithm. The constants \( 2\Delta y \) and \( 2\Delta y - 2\Delta x \) are calculated once for each line to be scan-converted, so the arithmetic involves only integer addition and subtraction of these two constants. Step 4 of the algorithm will be performed a total of \( \Delta x \) times.

**Bresenham’s Line-Drawing Algorithm for \(|m| < 1.0\)**

1. Input the two line endpoints and store the left endpoint in \((x_0, y_0)\).
2. Set the color for frame-buffer position \((x_0, y_0)\); i.e., plot the first point.
3. Calculate the constants \( \Delta x, \Delta y, 2\Delta y, \) and \( 2\Delta y - 2\Delta x \), and obtain the starting value for the decision parameter as
   \[
p_0 = 2\Delta y - \Delta x
   \]
4. At each \( x_k \) along the line, starting at \( k = 0 \), perform the following test:
   If \( p_k < 0 \), the next point to plot is \((x_k + 1, y_k)\) and
   \[
p_{k+1} = p_k + 2\Delta y
   \]
   Otherwise, the next point to plot is \((x_k + 1, y_k + 1)\) and
   \[
p_{k+1} = p_k + 2\Delta y - 2\Delta x
   \]
5. Repeat step 4 \( \Delta x - 1 \) more times.

**Example 1 Bresenham Line Drawing**

To illustrate the algorithm, we digitize the line with endpoints \((20, 10)\) and \((30, 18)\). This line has a slope of 0.8, with

\[
\Delta x = 10, \quad \Delta y = 8
\]

The initial decision parameter has the value

\[
p_0 = 2\Delta y - \Delta x = 6
\]
and the increments for calculating successive decision parameters are
\[ \frac{2}{\Delta y} = 16, \quad \frac{2}{\Delta y} - \frac{2}{\Delta x} = -4 \]

We plot the initial point \((x_0, y_0) = (20, 10)\), and determine successive pixel positions along the line path from the decision parameter as follows:

<table>
<thead>
<tr>
<th>(k)</th>
<th>(p_k)</th>
<th>((x_{k+1}, y_{k+1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>6</td>
<td>(21, 11)</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>(22, 12)</td>
</tr>
<tr>
<td>2</td>
<td>−2</td>
<td>(23, 12)</td>
</tr>
<tr>
<td>3</td>
<td>14</td>
<td>(24, 13)</td>
</tr>
<tr>
<td>4</td>
<td>10</td>
<td>(25, 14)</td>
</tr>
<tr>
<td>5</td>
<td>6</td>
<td>(26, 15)</td>
</tr>
<tr>
<td>6</td>
<td>2</td>
<td>(27, 16)</td>
</tr>
<tr>
<td>7</td>
<td>−2</td>
<td>(28, 16)</td>
</tr>
<tr>
<td>8</td>
<td>14</td>
<td>(29, 17)</td>
</tr>
<tr>
<td>9</td>
<td>10</td>
<td>(30, 18)</td>
</tr>
</tbody>
</table>

A plot of the pixels generated along this line path is shown in Figure 8.

An implementation of Bresenham line drawing for slopes in the range \(0 < m < 1.0\) is given in the following procedure. Endpoint pixel positions for the line are passed to this procedure, and pixels are plotted from the left endpoint to the right endpoint.

```c
#include <stdlib.h>
#include <math.h>

/* Bresenham line-drawing procedure for \(|m| < 1.0\). */
void lineBres (int x0, int y0, int xEnd, int yEnd) {
    int dx = fabs (xEnd - x0), dy = fabs(yEnd - y0);
    int p = 2 * dy - dx;
    int twoDy = 2 * dy, twoDyMinusDx = 2 * (dy - dx);
    int x, y;

    /* Determine which endpoint to use as start position. */
    if (x0 > xEnd) {
        x = xEnd;
        y = yEnd;
        xEnd = x0;
    }
```
Bresenham’s algorithm is generalized to lines with arbitrary slope by considering the symmetry between the various octants and quadrants of the $xy$ plane. For a line with positive slope greater than 1.0, we interchange the roles of the $x$ and $y$ directions. That is, we step along the $y$ direction in unit steps and calculate successive $x$ values nearest the line path. Also, we could revise the program to plot pixels starting from either endpoint. If the initial position for a line with positive slope is the right endpoint, both $x$ and $y$ decrease as we step from right to left. To ensure that the same pixels are plotted regardless of the starting endpoint, we always choose the upper (or the lower) of the two candidate pixels whenever the two vertical separations from the line path are equal ($d_{\text{lower}} = d_{\text{upper}}$). For negative slopes, the procedures are similar, except that now one coordinate decreases as the other increases. Finally, special cases can be handled separately: Horizontal lines ($\Delta y = 0$), vertical lines ($\Delta x = 0$), and diagonal lines ($|\Delta x| = |\Delta y|$) can each be loaded directly into the frame buffer without processing them through the line-plotting algorithm.

Displaying Polylines
Implementation of a polyline procedure is accomplished by invoking a line-drawing routine $n - 1$ times to display the lines connecting the $n$ endpoints. Each successive call passes the coordinate pair needed to plot the next line section, where the first endpoint of each coordinate pair is the last endpoint of the previous section. Once the color values for pixel positions along the first line segment have been set in the frame buffer, we process subsequent line segments starting with the next pixel position following the first endpoint for that segment. In this way, we can avoid setting the color of some endpoints twice. We discuss methods for avoiding the overlap of displayed objects in more detail in Section 8.

2 Parallel Line Algorithms
The line-generating algorithms we have discussed so far determine pixel positions sequentially. Using parallel processing, we can calculate multiple pixel positions along a line path simultaneously by partitioning the computations

```c
else {
    x = x0;
y = y0;
}
setPixel (x, y);
while (x < xEnd) {
x++;
    if (p < 0)
p += twoDy;
else {
y++;
p += ...
}
setPixel (x, y);
}
```
among the various processors available. One approach to the partitioning problem is to adapt an existing sequential algorithm to take advantage of multiple processors. Alternatively, we can look for other ways to set up the processing so that pixel positions can be calculated efficiently in parallel. An important consideration in devising a parallel algorithm is to balance the processing load among the available processors.

Given \( n_p \) processors, we can set up a parallel Bresenham line algorithm by subdividing the line path into \( n_p \) partitions and simultaneously generating line segments in each of the subintervals. For a line with slope \( 0 < m < 1.0 \) and left endpoint coordinate position \( (x_0, y_0) \), we partition the line along the positive \( x \) direction. The distance between beginning \( x \) positions of adjacent partitions can be calculated as

\[
\Delta x_p = \frac{\Delta x + n_p - 1}{n_p}
\]  

(17)

where \( \Delta x \) is the width of the line, and the value for partition width \( \Delta x_p \) is computed using integer division. Numbering the partitions, and the processors, as 0, 1, 2, up to \( n_p - 1 \), we calculate the starting \( x \) coordinate for the \( k \)th partition as

\[
x_k = x_0 + k \Delta x_p
\]  

(18)

For example, if we have \( n_p = 4 \) processors, with \( \Delta x = 15 \), the width of the partitions is 4 and the starting \( x \) values for the partitions are \( x_0, x_0 + 4, x_0 + 8 \), and \( x_0 + 12 \). With this partitioning scheme, the width of the last (rightmost) subinterval will be smaller than the others in some cases. In addition, if the line endpoints are not integers, truncation errors can result in variable-width partitions along the length of the line.

To apply Bresenham’s algorithm over the partitions, we need the initial value for the \( y \) coordinate and the initial value for the decision parameter in each partition. The change \( \Delta y_p \) in the \( y \) direction over each partition is calculated from the line slope \( m \) and partition width \( \Delta x_p \):

\[
\Delta y_p = m \Delta x_p
\]  

(19)

At the \( k \)th partition, the starting \( y \) coordinate is then

\[
y_k = y_0 + \text{round}(k \Delta y_p)
\]  

(20)

The initial decision parameter for Bresenham’s algorithm at the start of the \( k \)th subinterval is obtained from Equation 14:

\[
p_k = (k \Delta x_p)(2 \Delta y) - \text{round}(k \Delta y_p)(2 \Delta x) + 2 \Delta y - \Delta x
\]  

(21)

Each processor then calculates pixel positions over its assigned subinterval using the preceding starting decision parameter value and the starting coordinates \( (x_k, y_k) \). Floating-point calculations can be reduced to integer arithmetic in the computations for starting values \( y_k \) and \( p_k \) by substituting \( m = \Delta y/\Delta x \) and rearranging terms. We can extend the parallel Bresenham algorithm to a line with slope greater than 1.0 by partitioning the line in the \( y \) direction and calculating beginning \( x \) values for the partitions. For negative slopes, we increment coordinate values in one direction and decrement in the other.

Another way to set up parallel algorithms on raster systems is to assign each processor to a particular group of screen pixels. With a sufficient number of processors, we can assign each processor to one pixel within some screen region. This
approach can be adapted to a line display by assigning one processor to each of
the pixels within the limits of the coordinate extents of the line and calculating
pixel distances from the line path. The number of pixels within the bounding box
of a line is \( \Delta x \cdot \Delta y \) (as illustrated in Figure 9). Perpendicular distance \( d \) from
the line in Figure 9 to a pixel with coordinates \((x, y)\) is obtained with the calculation
\[
d = Ax + By + C
\]
where
\[
A = -\frac{\Delta y}{\text{linelength}}
B = \frac{\Delta x}{\text{linelength}}
C = \frac{x_0 \Delta y - y_0 \Delta x}{\text{linelength}}
\]
with
\[
\text{linelength} = \sqrt{\Delta x^2 + \Delta y^2}
\]
Once the constants \( A, B, \) and \( C \) have been evaluated for the line, each processor
must perform two multiplications and two additions to compute the pixel distance \( d \). A pixel is plotted if \( d \) is less than a specified line thickness parameter.

Instead of partitioning the screen into single pixels, we can assign to each
processor either a scan line or a column of pixels depending on the line slope. Each
processor then calculates the intersection of the line with the horizontal row or
vertical column of pixels assigned to that processor. For a line with slope \(|m| < 1.0
\), each processor simply solves the line equation for \( y \), given an \( x \) column value.
For a line with slope magnitude greater than 1.0, the line equation is solved for \( x \)
by each processor, given a scan line \( y \) value. Such direct methods, although slow
on sequential machines, can be performed efficiently using multiple processors.

\section{3 Setting Frame-Buffer Values}
A final stage in the implementation procedures for line segments and other objects
is to set the frame-buffer color values. Because scan-conversion algorithms generate
pixel positions at successive unit intervals, incremental operations can also be used to access the frame buffer efficiently at each step of the scan-conversion
process.

As a specific example, suppose the frame buffer array is addressed in row-
major order and that pixel positions are labeled from \((0, 0)\) at the lower-left corner
to \((x_{\text{max}}, y_{\text{max}})\) at the top-right corner (Figure 10) of the screen. For a bilevel system (one bit per pixel), the frame-buffer bit address for pixel position \((x, y)\) is
calculated as
\[
\text{addr}(x, y) = \text{addr}(0, 0) + y(x_{\text{max}} + 1) + x
\]
Moving across a scan line, we can calculate the frame-buffer address for the pixel
at \((x + 1, y)\) as the following offset from the address for position \((x, y)\):
\[
\text{addr}(x + 1, y) = \text{addr}(x, y) + 1
\]
Stepping diagonally up to the next scan line from \((x, y)\), we get to the frame-buffer address of \((x + 1, y + 1)\) with the calculation
\[
\text{addr}(x + 1, y + 1) = \text{addr}(x, y) + x_{\text{max}} + 2
\]
where the constant $x_{\text{max}} + 2$ is precomputed once for all line segments. Similar incremental calculations can be obtained from Equation 23 for unit steps in the negative $x$ and $y$ screen directions. Each of the address calculations involves only a single integer addition.

Methods for implementing these procedures depend on the capabilities of a particular system and the design requirements of the software package. With systems that can display a range of intensity values for each pixel, frame-buffer address calculations include pixel width (number of bits), as well as the pixel screen location.

4 Circle-Generating Algorithms

Because the circle is a frequently used component in pictures and graphs, a procedure for generating either full circles or circular arcs is included in many graphics packages. In addition, sometimes a general function is available in a graphics library for displaying various kinds of curves, including circles and ellipses.

Properties of Circles

A circle (Figure 11) is defined as the set of points that are all at a given distance $r$ from a center position $(x_c, y_c)$. For any circle point $(x, y)$, this distance relationship is expressed by the Pythagorean theorem in Cartesian coordinates as

$$\text{Equation 26}$$

$$\left( x - x_c \right)^2 + \left( y - y_c \right)^2 = r^2$$

We could use this equation to calculate the position of points on a circle circumference by stepping along the $x$ axis in unit steps from $x_c - r$ to $x_c + r$ and calculating the corresponding $y$ values at each position as

$$\text{Equation 27}$$

$$y = y_c \pm \sqrt{r^2 - (x_c - x)^2}$$

However, this is not the best method for generating a circle. One problem with this approach is that it involves considerable computation at each step. Moreover, the spacing between plotted pixel positions is not uniform, as demonstrated in Figure 12. We could adjust the spacing by interchanging $x$ and $y$ (stepping through $y$ values and calculating $x$ values) whenever the absolute value of the slope of the circle is greater than 1; but this simply increases the computation and processing required by the algorithm.

Another way to eliminate the unequal spacing shown in Figure 12 is to calculate points along the circular boundary using polar coordinates $r$ and $\theta$. 

![Figure 10](image1.png)

**Figure 10**

Pixel screen positions stored linearly in row-major order within the frame buffer.

![Figure 11](image2.png)

**Figure 11**

Circle with center coordinates $(x_c, y_c)$ and radius $r$.

![Figure 12](image3.png)

**Figure 12**

Upper half of a circle plotted with Equation 27 and with $(x_c, y_c) = (0, 0)$. 

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Expressing the circle equation in parametric polar form yields the pair of equations

\[ x = x_c + r \cos \theta \]
\[ y = y_c + r \sin \theta \]

When a display is generated with these equations using a fixed angular step size, a circle is plotted with equally spaced points along the circumference. To reduce calculations, we can use a large angular separation between points along the circumference and connect the points with straight-line segments to approximate the circular path. For a more continuous boundary on a raster display, we can set the angular step size at \( \frac{\pi}{2} \). This plots pixel positions that are approximately one unit apart. Although polar coordinates provide equal point spacing, the trigonometric calculations are still time-consuming.

For any of the previous circle-generating methods, we can reduce computations by considering the symmetry of circles. The shape of the circle is similar in each quadrant. Therefore, if we determine the curve positions in the first quadrant, we can generate the circle section in the second quadrant of the \( xy \) plane by noting that the two circle sections are symmetric with respect to the \( y \) axis. Also, circle sections in the third and fourth quadrants can be obtained from sections in the first and second quadrants by considering symmetry about the \( x \) axis. We can take this one step further and note that there is also symmetry between octants. Circle sections in adjacent octants within one quadrant are symmetric with respect to the 45\(^\circ\) line dividing the two octants. These symmetry conditions are illustrated in Figure 13, where a point at position \((x, y)\) on a one-eighth circle sector is mapped into the seven circle points in the other octants of the \( xy \) plane. Taking advantage of the circle symmetry in this way, we can generate all pixel positions around a circle by calculating only the points within the sector from \( x = 0 \) to \( x = y \). The slope of the curve in this octant has a magnitude less than or equal to 1.0. At \( x = 0 \), the circle slope is 0, and at \( x = y \), the slope is \(-1.0\).

Determining pixel positions along a circle circumference using symmetry and either Equation 26 or Equation 28 still requires a good deal of computation. The Cartesian equation 26 involves multiplications and square-root calculations, while the parametric equations contain multiplications and trigonometric calculations. More efficient circle algorithms are based on incremental calculation of decision parameters, as in the Bresenham line algorithm, which involves only simple integer operations.

Bresenham’s line algorithm for raster displays is adapted to circle generation by setting up decision parameters for finding the closest pixel to the circumference at each sampling step. The circle equation 26, however, is nonlinear, so that square-root evaluations would be required to compute pixel distances from a circular path. Bresenham’s circle algorithm avoids these square-root calculations by comparing the squares of the pixel separation distances.

However, it is possible to perform a direct distance comparison without a squaring operation. The basic idea in this approach is to test the halfway position between two pixels to determine if this midpoint is inside or outside the circle boundary. This method is applied more easily to other conics; and for an integer circle radius, the midpoint approach generates the same pixel positions as the Bresenham circle algorithm. For a straight-line segment, the midpoint method is equivalent to the Bresenham line algorithm. Also, the error involved in locating pixel positions along any conic section using the midpoint test is limited to half the pixel separation.
Midpoint Circle Algorithm

As in the raster line algorithm, we sample at unit intervals and determine the closest pixel position to the specified circle path at each step. For a given radius \( r \) and screen center position \((x_c, y_c)\), we can first set up our algorithm to calculate pixel positions around a circle path centered at the coordinate origin \((0, 0)\). Then each calculated position \((x, y)\) is moved to its proper screen position by adding \(x_c\) to \(x\) and \(y_c\) to \(y\). Along the circle section from \(x = 0\) to \(x = y\) in the first quadrant, the slope of the curve varies from 0 to \(-1.0\). Therefore, we can take unit steps in the positive \(x\) direction over this octant and use a decision parameter to determine which of the two possible pixel positions in any column is vertically closer to the circle path. Positions in the other seven octants are then obtained by symmetry.

To apply the midpoint method, we define a circle function as

\[
f_{\text{circ}}(x, y) = x^2 + y^2 - r^2
\]  

(29)

Any point \((x, y)\) on the boundary of the circle with radius \(r\) satisfies the equation \(f_{\text{circ}}(x, y) = 0\). If the point is in the interior of the circle, the circle function is negative; and if the point is outside the circle, the circle function is positive. To summarize, the relative position of any point \((x, y)\) can be determined by checking the sign of the circle function as follows:

\[
f_{\text{circ}}(x, y) \begin{cases} < 0, & \text{if } (x, y) \text{ is inside the circle boundary} \\ = 0, & \text{if } (x, y) \text{ is on the circle boundary} \\ > 0, & \text{if } (x, y) \text{ is outside the circle boundary} \end{cases}
\]

(30)

The tests in 30 are performed for the midpositions between pixels near the circle path at each sampling step. Thus, the circle function is the decision parameter in the midpoint algorithm, and we can set up incremental calculations for this function as we did in the line algorithm.

Figure 14 shows the midpoint between the two candidate pixels at sampling position \(x_k + 1\). Assuming that we have just plotted the pixel at \((x_k, y_k)\), we next need to determine whether the pixel at position \((x_k + 1, y_k)\) or the one at position \((x_k + 1, y_k - 1)\) is closer to the circle. Our decision parameter is the circle function 29 evaluated at the midpoint between these two pixels:

\[
p_k = f_{\text{circ}} \left( x_k + 1, y_k - \frac{1}{2} \right)
\]

(31)

If \(p_k < 0\), this midpoint is inside the circle and the pixel on scan line \(y_k\) is closer to the circle boundary. Otherwise, the midposition is outside or on the circle boundary, and we select the pixel on scan line \(y_k - 1\).

Successive decision parameters are obtained using incremental calculations. We obtain a recursive expression for the next decision parameter by evaluating the circle function at sampling position \(x_{k+1} + 1 = x_k + 2\):

\[
p_{k+1} = f_{\text{circ}} \left( x_{k+1} + 1, y_{k+1} - \frac{1}{2} \right)
\]

\[
= [(x_k + 1) + 1]^2 + \left( y_{k+1} - \frac{1}{2} \right)^2 - r^2
\]

or

\[
p_{k+1} = p_k + 2(x_k + 1) + \left( y_{k+1}^2 - y_k^2 \right) - (y_{k+1} - y_k) + 1
\]

(32)

where \(y_{k+1}\) is either \(y_k\) or \(y_k - 1\), depending on the sign of \(p_k\).
Increments for obtaining $p_{k+1}$ are either $2x_{k+1} + 1$ (if $p_k$ is negative) or $2x_{k+1} + 1 - 2y_{k+1}$. Evaluation of the terms $2x_{k+1}$ and $2y_{k+1}$ can also be done incrementally as

$$
2x_{k+1} = 2x_k + 2
$$

$$
2y_{k+1} = 2y_k - 2
$$

At the start position $(0, r)$, these two terms have the values 0 and $2r$, respectively. Each successive value for the $2x_{k+1}$ term is obtained by adding 2 to the previous value, and each successive value for the $2y_{k+1}$ term is obtained by subtracting 2 from the previous value.

The initial decision parameter is obtained by evaluating the circle function at the start position $(x_0, y_0) = (0, r)$:

$$
p_0 = f_{\text{circ}} \left( 1, r - \frac{1}{2} \right)
$$

$$
= 1 + \left( r - \frac{1}{2} \right)^2 - r^2
$$

or

$$
p_0 = \frac{5}{4} - r
$$

If the radius $r$ is specified as an integer, we can simply round $p_0$ to

$$
p_0 = 1 - r \quad \text{(for } r \text{ an integer)}
$$

because all increments are integers.

As in Bresenham’s line algorithm, the midpoint method calculates pixel positions along the circumference of a circle using integer additions and subtractions, assuming that the circle parameters are specified in integer screen coordinates. We can summarize the steps in the midpoint circle algorithm as follows:

**Midpoint Circle Algorithm**

1. Input radius $r$ and circle center $(x_c, y_c)$, then set the coordinates for the first point on the circumference of a circle centered on the origin as $(x_0, y_0) = (0, r)$.

2. Calculate the initial value of the decision parameter as

$$
p_0 = \frac{5}{4} - r
$$

3. At each $x_k$ position, starting at $k = 0$, perform the following test: If $p_k < 0$, the next point along the circle centered on $(0, 0)$ is $(x_{k+1}, y_k)$ and

$$
p_{k+1} = p_k + 2x_{k+1} + 1
$$

Otherwise, the next point along the circle is $(x_k + 1, y_k - 1)$ and

$$
p_{k+1} = p_k + 2x_{k+1} + 1 - 2y_{k+1}
$$

where $2x_{k+1} = 2x_k + 2$ and $2y_{k+1} = 2y_k - 2$.

4. Determine symmetry points in the other seven octants.

5. Move each calculated pixel position $(x, y)$ onto the circular path centered at $(x_c, y_c)$ and plot the coordinate values as follows:

$$
x = x + x_c, \quad y = y + y_c.
$$

6. Repeat steps 3 through 5 until $x \geq y$. 

---

*Implementation Algorithms for Graphics Primitives and Attributes*
EXAMPLE 2 Midpoint Circle Drawing

Given a circle radius \( r = 10 \), we demonstrate the midpoint circle algorithm by determining positions along the circle octant in the first quadrant from \( x = 0 \) to \( x = y \). The initial value of the decision parameter is

\[ p_0 = 1 - r = -9 \]

For the circle centered on the coordinate origin, the initial point is \((x_0, y_0) = (0, 10)\), and initial increment terms for calculating the decision parameters are

\[ 2x_0 = 0, \quad 2y_0 = 20 \]

Successive midpoint decision parameter values and the corresponding coordinate positions along the circle path are listed in the following table:

<table>
<thead>
<tr>
<th>( k )</th>
<th>( p_k )</th>
<th>((x_{k+1}, y_{k+1}))</th>
<th>( 2y_{k+1} )</th>
<th>( 2y_{k+1} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>-9</td>
<td>(1, 10)</td>
<td>2</td>
<td>20</td>
</tr>
<tr>
<td>1</td>
<td>-6</td>
<td>(2, 10)</td>
<td>4</td>
<td>20</td>
</tr>
<tr>
<td>2</td>
<td>-1</td>
<td>(3, 10)</td>
<td>6</td>
<td>20</td>
</tr>
<tr>
<td>3</td>
<td>6</td>
<td>(4, 9)</td>
<td>8</td>
<td>18</td>
</tr>
<tr>
<td>4</td>
<td>8</td>
<td>(5, 9)</td>
<td>10</td>
<td>18</td>
</tr>
<tr>
<td>5</td>
<td>5</td>
<td>(6, 8)</td>
<td>12</td>
<td>16</td>
</tr>
<tr>
<td>6</td>
<td>5</td>
<td>(7, 7)</td>
<td>14</td>
<td>14</td>
</tr>
</tbody>
</table>

A plot of the generated pixel positions in the first quadrant is shown in Figure 15.

The following code segment illustrates procedures that could be used to implement the midpoint circle algorithm. Values for a circle radius and for the center coordinates of the circle are passed to procedure `circleMidpoint`. A pixel position along the circular path in the first octant is then computed and passed to procedure `circlePlotPoints`. This procedure sets the circle color in the frame buffer for all circle symmetry positions with repeated calls to the `setPixel` routine, which is implemented with the OpenGL point-plotting functions.
#include <GL/glut.h>

class screenPt
{
    private:
        GLint x, y;

    public:
        /* Default Constructor: initializes coordinate position to (0, 0). */
        screenPt () {
            x = y = 0;
        }

        void setCoords (GLint xCoordValue, GLint yCoordValue) {
            x = xCoordValue;
            y = yCoordValue;
        }

        GLint getx ( ) const {
            return x;
        }

        GLint gety ( ) const {
            return y;
        }

        void incrementx () {
            x++;
        }

        void decrementy () {
            y--;
        }
};

void setPixel (GLint xCoord, GLint yCoord)
{
    glBegin (GL_POINTS);
        glVertex2i (xCoord, yCoord);
    glEnd ( );
}

void circleMidpoint (GLint xc, GLint yc, GLint radius)
{
    screenPt circPt;

    GLint p = 1 - radius;       // Initial value for midpoint parameter.

    circPt.setCoords (0, radius);  // Set coordinates for top point of circle.

    void circlePlotPoints (GLint, GLint, screenPt);
    /* Plot the initial point in each circle quadrant. */
    circlePlotPoints (xc, yc, circPt);
    /* Calculate next point and plot in each octant. */
while (circPt.getx() < circPt.gety()) {
    circPt.incrementx();
    if (p < 0)
        p += 2 * circPt.getx() + 1;
    else {
        circPt.decrementy();
        p += 2 * (circPt.getx() - circPt.gety()) + 1;
    }
    circlePlotPoints(xc, yc, circPt);
}

void circlePlotPoints(GLint xc, GLint yc, screenPt circPt) {
    setPixel(xc + circPt.getx(), yc + circPt.gety());
    setPixel(xc - circPt.getx(), yc + circPt.gety());
    setPixel(xc + circPt.getx(), yc - circPt.gety());
    setPixel(xc - circPt.getx(), yc - circPt.gety());
    setPixel(xc + circPt.gety(), yc + circPt.getx());
    setPixel(xc - circPt.gety(), yc + circPt.getx());
    setPixel(xc + circPt.gety(), yc - circPt.getx());
    setPixel(xc - circPt.gety(), yc - circPt.getx());
}

5 Ellipse-Generating Algorithms

Loosely stated, an ellipse is an elongated circle. We can also describe an ellipse as a modified circle whose radius varies from a maximum value in one direction to a minimum value in the perpendicular direction. The straight-line segments through the interior of the ellipse in these two perpendicular directions are referred to as the major and minor axes of the ellipse.

Properties of Ellipses

A precise definition of an ellipse can be given in terms of the distances from any point on the ellipse to two fixed positions, called the foci of the ellipse. The sum of these two distances is the same value for all points on the ellipse (Figure 16). If the distances to the two focus positions from any point \( P = (x, y) \) on the ellipse are labeled \( d_1 \) and \( d_2 \), then the general equation of an ellipse can be stated as

\[
d_1 + d_2 = \text{constant} \tag{34}\]

Expressing distances \( d_1 \) and \( d_2 \) in terms of the focal coordinates \( F_1 = (x_1, y_1) \) and \( F_2 = (x_2, y_2) \), we have

\[
\sqrt{(x - x_1)^2 + (y - y_1)^2} + \sqrt{(x - x_2)^2 + (y - y_2)^2} = \text{constant} \tag{35}\]

By squaring this equation, isolating the remaining radical, and squaring again, we can rewrite the general ellipse equation in the form

\[
Ax^2 + By^2 + Cxy + Dx + Ey + F = 0 \tag{36}\]
where the coefficients $A, B, C, D, E,$ and $F$ are evaluated in terms of the focal coordinates and the dimensions of the major and minor axes of the ellipse. The major axis is the straight-line segment extending from one side of the ellipse to the other through the foci. The minor axis spans the shorter dimension of the ellipse, perpendicularly bisecting the major axis at the halfway position (ellipse center) between the two foci.

An interactive method for specifying an ellipse in an arbitrary orientation is to input the two foci and a point on the ellipse boundary. With these three coordinate positions, we can evaluate the constant in Equation 35. Then, the values for the coefficients in Equation 36 can be computed and used to generate pixels along the elliptical path.

Ellipse equations are greatly simplified if the major and minor axes are oriented to align with the coordinate axes. In Figure 17, we show an ellipse in “standard position,” with major and minor axes oriented parallel to the $x$ and $y$ axes. Parameter $r_x$ for this example labels the semimajor axis, and parameter $r_y$ labels the semiminor axis. The equation for the ellipse shown in Figure 17 can be written in terms of the ellipse center coordinates and parameters $r_x$ and $r_y$ as

$$\left(\frac{x - x_c}{r_x}\right)^2 + \left(\frac{y - y_c}{r_y}\right)^2 = 1$$

(37)

Using polar coordinates $r$ and $\theta$, we can also describe the ellipse in standard position with the parametric equations

$$x = x_c + r \cos \theta$$

$$y = y_c + r \sin \theta$$

(38)

Angle $\theta$, called the eccentric angle of the ellipse, is measured around the perimeter of a bounding circle. If $r_x > r_y$, the radius of the bounding circle is $r = r_x$ (Figure 18). Otherwise, the bounding circle has radius $r = r_y$.

As with the circle algorithm, symmetry considerations can be used to reduce computations. An ellipse in standard position is symmetric between quadrants, but, unlike a circle, it is not symmetric between the two octants of a quadrant. Thus, we must calculate pixel positions along the elliptical arc throughout one quadrant, then use symmetry to obtain curve positions in the remaining three quadrants (Figure 19).

**Midpoint Ellipse Algorithm**

Our approach here is similar to that used in displaying a raster circle. Given parameters $r_x, r_y$, and $(x_c, y_c)$, we determine curve positions $(x, y)$ for an ellipse in standard position centered on the origin, then we shift all the points using a fixed offset so that the ellipse is centered at $(x_c, y_c)$. If we wish also to display the ellipse in nonstandard position, we could rotate the ellipse about its center coordinates to reorient the major and minor axes in the desired directions. For the present, we consider only the display of ellipses in standard position.

The midpoint ellipse method is applied throughout the first quadrant in two parts. Figure 20 shows the division of the first quadrant according to the slope of an ellipse with $r_x < r_y$. We process this quadrant by taking unit steps in the $x$ direction where the slope of the curve has a magnitude less than 1.0, and then we take unit steps in the $y$ direction where the slope has a magnitude greater than 1.0.
At the next sampling position \( x_{k+1} + 1 = x_k + 2 \), the decision parameter for region 1 is evaluated as

\[
p_{1k+1} = f_{\text{ellipse}} \left( x_{k+1} + 1, y_{k+1} - \frac{1}{2} \right)
\]

\[
= r_y^2\left[(x_k + 1) + 1\right]^2 + r_x^2\left(y_{k+1} - \frac{1}{2}\right)^2 - r_x^2r_y^2
\]

or

\[
p_{1k+1} = p_{1k} + 2r_y^2(x_k + 1) + r_x^2 + r_x^2\left[(y_{k+1} - \frac{1}{2}\right)^2 - \left(y_k - \frac{1}{2}\right)^2)
\]  

(44)

where \( y_{k+1} \) is either \( y_k \) or \( y_k - 1 \), depending on the sign of \( p_{1k} \).

Decision parameters are incremented by the following amounts:

\[
\text{increment} = \begin{cases} 
2r_y^2x_k + r_x^2, & \text{if } p_{1k} < 0 \\
2r_y^2x_k + r_x^2 - 2r_y^2y_k, & \text{if } p_{1k} \geq 0 
\end{cases}
\]

Increments for the decision parameters can be calculated using only addition and subtraction, as in the circle algorithm, because values for the terms \( 2r_y^2x \) and \( 2r_y^2y \) can be obtained incrementally. At the initial position \((0, r_y)\), these two terms evaluate to

\[
2r_y^2x = 0
\]

\[
2r_y^2y = 2r_x^2r_y
\]

(45)

(46)

As \( x \) and \( y \) are incremented, updated values are obtained by adding \( 2r_y^2 \) to the current value of the increment term in Equation 45 and subtracting \( 2r_y^2 \) from the current value of the increment term in Equation 46. The updated increment values are compared at each step, and we move from region 1 to region 2 when condition 42 is satisfied.

In region 1, the initial value of the decision parameter is obtained by evaluating the ellipse function at the start position \((x_0, y_0) = (0, r_y)\):

\[
p_{10} = f_{\text{ellipse}} \left( 1, r_y - \frac{1}{2} \right)
\]

\[
= r_y^2 + r_x^2\left(r_y - \frac{1}{2}\right)^2 - r_x^2r_y^2
\]

or

\[
p_{10} = r_y^2 - r_x^2r_y + \frac{1}{4}r_x^2
\]

(47)

Over region 2, we sample at unit intervals in the negative \( y \) direction, and the midpoint is now taken between horizontal pixels at each step (Figure 22). For this region, the decision parameter is evaluated as

\[
p_{2k} = f_{\text{ellipse}} \left( x_k + \frac{1}{2}, y_k - 1 \right)
\]

\[
= r_y^2\left(x_k + \frac{1}{2}\right)^2 + r_x^2(y_k - 1)^2 - r_x^2r_y^2
\]

(48)

If \( p_{2k} > 0 \), the midposition is outside the ellipse boundary, and we select the pixel at \( x_k \). If \( p_{2k} \leq 0 \), the midposition is inside or on the ellipse boundary, and we select pixel position \( x_k + 1 \).
Regions 1 and 2 (Figure 20) can be processed in various ways. We can start at position \((0, r_y)\) and step clockwise along the elliptical path in the first quadrant, shifting from unit steps in \(x\) to unit steps in \(y\) when the slope becomes less than \(-1.0\). Alternatively, we could start at \((r_x, 0)\) and select points in a counterclockwise order, shifting from unit steps in \(y\) to unit steps in \(x\) when the slope becomes greater than \(-1.0\). With parallel processors, we could calculate pixel positions in the two regions simultaneously. As an example of a sequential implementation of the midpoint algorithm, we take the start position at \((0, r_y)\) and step along the ellipse path in clockwise order throughout the first quadrant.

We define an ellipse function from Equation 37 with \((x, y) = (0, 0)\) as

\[
f_{\text{ellipse}}(x, y) = r_x^2 x^2 + r_y^2 y^2 - r_x^2 r_y^2
\]

which has the following properties:

\[
f_{\text{ellipse}}(x, y) \begin{cases} < 0, & \text{if } (x, y) \text{ is inside the ellipse boundary} \\ = 0, & \text{if } (x, y) \text{ is on the ellipse boundary} \\ > 0, & \text{if } (x, y) \text{ is outside the ellipse boundary} \end{cases}
\]

Thus, the ellipse function \(f_{\text{ellipse}}(x, y)\) serves as the decision parameter in the midpoint algorithm. At each sampling position, we select the next pixel along the ellipse path according to the sign of the ellipse function evaluated at the midpoint between the two candidate pixels.

Starting at \((0, r_y)\), we take unit steps in the \(x\) direction until we reach the boundary between region 1 and region 2 (Figure 20). Then we switch to unit steps in the \(y\) direction over the remainder of the curve in the first quadrant. At each step we need to test the value of the slope of the curve. The ellipse slope is calculated from Equation 39 as

\[
\frac{dy}{dx} = -\frac{2r_x^2 x}{2r_y^2 y}
\]

At the boundary between region 1 and region 2, \(dy/dx = -1.0\) and

\[2r_x^2 x = 2r_y^2 y\]

Therefore, we move out of region 1 whenever

\[2r_x^2 x \geq 2r_y^2 y\]

Figure 21 shows the midpoint between the two candidate pixels at sampling position \(x_k + 1\) in the first region. Assuming position \((x_k, y_k)\) has been selected in the previous step, we determine the next position along the ellipse path by evaluating the decision parameter (that is, the ellipse function 39) at this midpoint:

\[
p_{1k} = f_{\text{ellipse}}(x_k + 1, y_k - \frac{1}{2})
\]

\[= r_y^2 (x_k + 1)^2 + r_x^2 \left(y_k - \frac{1}{2}\right)^2 - r_x^2 r_y^2
\]

If \(p_{1k} < 0\), the midpoint is inside the ellipse and the pixel on scan line \(y_k\) is closer to the ellipse boundary. Otherwise, the midpoint is outside or on the ellipse boundary, and we select the pixel on scan line \(y_k - 1\).
To determine the relationship between successive decision parameters in region 2, we evaluate the ellipse function at the next sampling step $y_{k+1} - 1 = y_k - 2$:

$$p_{2k+1} = f_{\text{ellipse}}\left(x_{k+1} + \frac{1}{2}, y_{k+1} - 1\right)$$

$$= r_y^2 \left(x_{k+1} + \frac{1}{2}\right)^2 + r_x^2 \left[(y_k - 1) - 1\right]^2 - r_x^2 r_y^2$$

or

$$p_{2k+1} = p_{2k} - 2r_x^2(y_k - 1) + r_x^2 + r_y^2 \left[\left(x_{k+1} + \frac{1}{2}\right)^2 - \left(x_k + \frac{1}{2}\right)^2\right]$$

with $x_{k+1}$ set either to $x_k$ or to $x_k + 1$, depending on the sign of $p_{2k}$.

When we enter region 2, the initial position $(x_0, y_0)$ is taken as the last position selected in region 1 and the initial decision parameter in region 2 is then

$$p_{20} = f_{\text{ellipse}}\left(x_0 + \frac{1}{2}, y_0 - 1\right)$$

$$= r_y^2 \left(x_0 + \frac{1}{2}\right)^2 + r_x^2 (y_0 - 1)^2 - r_x^2 r_y^2$$

To simplify the calculation of $p_{20}$, we could select pixel positions in counterclockwise order starting at $(r_x, 0)$. Unit steps would then be taken in the positive $y$ direction up to the last position selected in region 1.

This midpoint algorithm can be adapted to generate an ellipse in nonstandard position using the ellipse function Equation 36 and calculating pixel positions over the entire elliptical path. Alternatively, we could reorient the ellipse axes to standard position, apply the midpoint ellipse algorithm to determine curve positions, and then convert calculated pixel positions to path positions along the original ellipse orientation.

Assuming $r_x$, $r_y$, and the ellipse center are given in integer screen coordinates, we need only incremental integer calculations to determine values for the decision parameters in the midpoint ellipse algorithm. The increments $r_x^2$, $r_y^2$, $2r_x^2$, and $2r_y^2$ are evaluated once at the beginning of the procedure. In the following summary, we list the steps for displaying an ellipse using the midpoint algorithm:

### Midpoint Ellipse Algorithm

1. Input $r_x$, $r_y$, and ellipse center $(x_c, y_c)$, and obtain the first point on an ellipse centered on the origin as
   
   $$(x_0, y_0) = (0, r_y)$$

2. Calculate the initial value of the decision parameter in region 1 as
   
   $$p_{1_0} = r_y^2 - r_x^2 r_y + \frac{1}{4} r_x^2$$

3. At each $x_k$ position in region 1, starting at $k = 0$, perform the following test: If $p_{1_k} < 0$, the next point along the ellipse centered on $(0, 0)$ is $(x_{k+1}, y_k)$ and
   
   $$p_{1_{k+1}} = p_{1_k} + 2r_y^2 x_{k+1} + r_y^2$$

   Otherwise, the next point along the ellipse is $(x_k + 1, y_k - 1)$ and
   
   $$p_{1_{k+1}} = p_{1_k} + 2r_y^2 x_{k+1} - 2r_x^2 y_{k+1} + r_y^2$$
with
\[ 2r_y^2 x_{k+1} = 2r_y^2 x_k + 2r_y^2, \quad 2r_x^2 y_{k+1} = 2r_x^2 y_k - 2r_x^2 \]
and continue until \( 2r_y^2 x \geq 2r_x^2 y \).

4. Calculate the initial value of the decision parameter in region 2 as
\[ p2_0 = r_y^2 \left( x_0 + \frac{1}{2} \right)^2 + r_x^2 (y_0 - 1)^2 - r_x^2 r_y^2 \]
where \((x_0, y_0)\) is the last position calculated in region 1.

5. At each \( y_k \) position in region 2, starting at \( k = 0 \), perform the following test: If \( p2_k > 0 \), the next point along the ellipse centered on \((0, 0)\) is \((x_k, y_k - 1)\) and
\[ p2_{k+1} = p2_k - 2r_x^2 y_{k+1} + r_x^2 \]
Otherwise, the next point along the ellipse is \((x_k + 1, y_k - 1)\) and
\[ p2_{k+1} = p2_k + 2r_x^2 x_{k+1} - 2r_x^2 y_{k+1} + r_x^2 \]
using the same incremental calculations for \( x \) and \( y \) as in region 1. Continue until \( y = 0 \).

6. For both regions, determine symmetry points in the other three quadrants.

7. Move each calculated pixel position \((x, y)\) onto the elliptical path centered on \((x_r, y_r)\) and plot these coordinate values:
\[ x = x + x_r, \quad y = y + y_r. \]

**EXAMPLE 3  Midpoint Ellipse Drawing**

Given input ellipse parameters \( r_x = 8 \) and \( r_y = 6 \), we illustrate the steps in the midpoint ellipse algorithm by determining raster positions along the ellipse path in the first quadrant. Initial values and increments for the decision parameter calculations are
\[ 2r_y^2 x = 0 \quad \text{(with increment } 2r_y^2 = 72) \]
\[ 2r_x^2 y = 2r_x^2 r_y \quad \text{(with increment } -2r_x^2 = -128) \]
For region 1, the initial point for the ellipse centered on the origin is \((x_0, y_0) = (0, 6)\), and the initial decision parameter value is
\[ p1_0 = r_y^2 - r_x^2 r_y + \frac{1}{4} r_x^2 = -332 \]
Successive midpoint decision-parameter values and the pixel positions along the ellipse are listed in the following table:

<table>
<thead>
<tr>
<th>( k )</th>
<th>( p1_k )</th>
<th>((x_{k+1}, y_{k+1}))</th>
<th>( 2r_y^2 x_{k+1} )</th>
<th>( 2r_x^2 y_{k+1} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>-332</td>
<td>(1, 6)</td>
<td>72</td>
<td>768</td>
</tr>
<tr>
<td>1</td>
<td>-224</td>
<td>(2, 6)</td>
<td>144</td>
<td>768</td>
</tr>
<tr>
<td>2</td>
<td>-44</td>
<td>(3, 6)</td>
<td>216</td>
<td>768</td>
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<td>3</td>
<td>208</td>
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<td>640</td>
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<tr>
<td>4</td>
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<td>(5, 5)</td>
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<td>640</td>
</tr>
<tr>
<td>5</td>
<td>288</td>
<td>(6, 4)</td>
<td>432</td>
<td>512</td>
</tr>
<tr>
<td>6</td>
<td>244</td>
<td>(7, 3)</td>
<td>504</td>
<td>384</td>
</tr>
</tbody>
</table>
We now move out of region 1 because $2r_x^2 > 2r_y^2$.

For region 2, the initial point is $(x_0, y_0) = (7, 3)$ and the initial decision parameter is

$$p_{20} = f_{\text{ellipse}} \left( 7 + \frac{1}{2}, 2 \right) = -151$$

The remaining positions along the ellipse path in the first quadrant are then calculated as

<table>
<thead>
<tr>
<th>$k$</th>
<th>$p_{1k}$</th>
<th>$\left( x_{k+1}, y_{k+1} \right)$</th>
<th>$2r_y^2x_{k+1}$</th>
<th>$2r_x^2y_{k+1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>$-151$</td>
<td>$\left( 8, 2 \right)$</td>
<td>576</td>
<td>256</td>
</tr>
<tr>
<td>1</td>
<td>233</td>
<td>$\left( 8, 1 \right)$</td>
<td>576</td>
<td>128</td>
</tr>
<tr>
<td>2</td>
<td>745</td>
<td>$\left( 8, 0 \right)$</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

A plot of the calculated positions for the ellipse within the first quadrant is shown in Figure 23.

In the following code segment, example procedures are given for implementing the midpoint ellipse algorithm. Values for the ellipse parameters $R_x$, $R_y$, $x_{\text{Center}}$, and $y_{\text{Center}}$ are input to procedure `ellipseMidpoint`. Positions along the curve in the first quadrant are then calculated and passed to procedure `ellipsePlotPoints`. Symmetry is used to obtain ellipse positions in the other three quadrants, and the `setPixel` routine sets the ellipse color in the framebuffer locations corresponding to these positions.

```c
inline int round (const float a) { return int (a + 0.5); }

/* The following procedure accepts values for an ellipse */
/* center position and its semimajor and semiminor axes, then */
/* calculates ellipse positions using the midpoint algorithm. */
void ellipseMidpoint (int xCenter, int yCenter, int Rx, int Ry)
{
    int Rx2 = Rx * Rx;
    int Ry2 = Ry * Ry;
    int twoRx2 = 2 * Rx2;
    int twoRy2 = 2 * Ry2;
    int p;
    int x = 0;
    int y = Ry;
    int px = 0;
    int py = twoRx2 * y;
    void ellipsePlotPoints (int, int, int, int):
```
/* Plot the initial point in each quadrant. */
ellipsePlotPoints (xCenter, yCenter, x, y);

/* Region 1 */
p = round (Ry2 * (x+0.5) * (x+0.5) + Rx2 * (y-1) * (y-1) - Rx2 * Ry2);
while (y > 0) {
    y--;
    py -= twoRx2;
    if (p > 0)
        p += Rx2 - py;
    else {
        x++;
        px += twoRy2;
    }
    ellipsePlotPoints (xCenter, yCenter, x, y);
}

/* Region 2 */
p = round (Ry2 - (Rx2 * Ry) + (0.25 * Rx2));
while (px < py) {
    x++;
    px += twoRy2;
    if (p < 0)
        p += Ry2 + px;
    else {
        y--;
        py -= twoRx2;
        p += Ry2 + px - py;
    }
    ellipsePlotPoints (xCenter, yCenter, x, y);
}

void ellipsePlotPoints (int xCenter, int yCenter, int x, int y):
{
    setPixel (xCenter + x, yCenter + y);
    setPixel (xCenter - x, yCenter + y);
    setPixel (xCenter + x, yCenter - y);
    setPixel (xCenter - x, yCenter - y);
}

6 Other Curves

Various curve functions are useful in object modeling, animation path specifications, data and function graphing, and other graphics applications. Commonly encountered curves include conics, trigonometric and exponential functions, probability distributions, general polynomials, and spline functions. Displays of these curves can be generated with methods similar to those discussed for the circle and ellipse functions. We can obtain positions along curve paths directly from explicit representations $y = f(x)$ or from parametric forms. Alternatively, we could apply the incremental midpoint method to plot curves described with implicit functions $f(x, y) = 0$. 
A simple method for displaying a curved line is to approximate it with straight-line segments. Parametric representations are often useful in this case for obtaining equally spaced positions along the curve path for the line endpoints. We can also generate equally spaced positions from an explicit representation by choosing the independent variable according to the slope of the curve. Where the slope of $y = f(x)$ has a magnitude less than 1, we choose $x$ as the independent variable and calculate $y$ values at equal $x$ increments. To obtain equal spacing where the slope has a magnitude greater than 1, we use the inverse function, $x = f^{-1}(y)$, and calculate values of $x$ at equal $y$ steps.

Straight-line or curve approximations are used to generate a line graph for a set of discrete data values. We could join the discrete points with straight-line segments, or we could use linear regression (least squares) to approximate the data set with a single straight line. A nonlinear least-squares approach is used to display the data set with some approximating function, usually a polynomial.

As with circles and ellipses, many functions possess symmetries that can be exploited to reduce the computation of coordinate positions along curve paths. For example, the normal probability distribution function is symmetric about a center position (the mean), and all points within one cycle of a sine curve can be generated from the points in a 90° interval.

**Conic Sections**

In general, we can describe a conic section (or conic) with the second-degree equation

$$Ax^2 + By^2 + Cxy + Dx + Ey + F = 0$$

where the values for parameters $A$, $B$, $C$, $D$, $E$, and $F$ determine the kind of curve that we are to display. Given this set of coefficients, we can determine the particular conic that will be generated by evaluating the discriminant $B^2 - 4AC$:

$$B^2 - 4AC < 0, \text{ generates an ellipse (or circle)}$$

$$B^2 - 4AC = 0, \text{ generates a parabola}$$

$$B^2 - 4AC > 0, \text{ generates a hyperbola}$$

Equation 52 also describes the “degenerate” conics: points and straight lines.

In some applications, circular and elliptical arcs are conveniently specified with the beginning and ending angular values for the arc, as illustrated in Figure 24. Such arcs are sometimes defined by their endpoint coordinate positions. For either case, we could generate the arc with a modified midpoint method, or we could display a set of approximating straight-line segments.

Ellipses, hyperbolas, and parabolas are particularly useful in certain animation applications. These curves describe orbital and other motions for objects subjected to gravitational, electromagnetic, or nuclear forces. Planetary orbits in the solar system, for example, are approximated with ellipses; and an object projected into a uniform gravitational field travels along a parabolic trajectory. Figure 25 shows a parabolic path in standard position for a gravitational field acting in the negative $y$ direction. The explicit equation for the parabolic trajectory of the object shown can be written as

$$y = y_0 + a(x - x_0)^2 + b(x - x_0)$$
construct a cubic polynomial curve section between each pair of specified points. Each curve section is then described in parametric form as

\[
\begin{align*}
x &= a_x0 + a_x1u + a_x2u^2 + a_x3u^3 \\
y &= a_y0 + a_y1u + a_y2u^2 + a_y3u^3
\end{align*}
\]  

(58)

where parameter \(u\) varies over the interval from 0 to 1.0. Values for the coefficients of \(u\) in the preceding equations are determined from boundary conditions on the curve sections. One boundary condition is that two adjacent curve sections have the same coordinate position at the boundary, and a second condition is to match the two curve slopes at the boundary so that we obtain one continuous, smooth curve (Figure 27). Continuous curves that are formed with polynomial pieces are called **spline curves**, or simply **splines**.

### 7 Parallel Curve Algorithms

Methods for exploiting parallelism in curve generation are similar to those used in displaying straight-line segments. We can either adapt a sequential algorithm by allocating processors according to curve partitions, or we could devise other methods and assign processors to screen partitions.

A parallel midpoint method for displaying circles is to divide the circular arc from 45° to 90° into equal subarcs and assign a separate processor to each subarc. As in the parallel Bresenham line algorithm, we then need to set up computations to determine the beginning \(y\) value and decision parameter \(p_k\) value for each processor. Pixel positions are calculated throughout each subarc, and positions in the other circle octants can be obtained by symmetry. Similarly, a parallel ellipse midpoint method divides the elliptical arc over the first quadrant into equal subarcs and parcels these out to separate processors. Again, pixel positions in the other quadrants are determined by symmetry. A screen-partitioning scheme for circles and ellipses is to assign each scan line that crosses the curve to a separate processor. In this case, each processor uses the circle or ellipse equation to calculate curve intersection coordinates.

For the display of elliptical arcs or other curves, we can simply use the scanline partitioning method. Each processor uses the curve equation to locate the intersection positions along its assigned scan line. With processors assigned to individual pixels, each processor would calculate the distance (or distance squared) from the curve to its assigned pixel. If the calculated distance is less than a predefined value, the pixel is plotted.

### 8 Pixel Addressing and Object Geometry

In discussing the raster algorithms for displaying graphics primitives, we assumed that frame-buffer coordinates referenced the center of a screen pixel position. We now consider the effects of different addressing schemes and an alternate pixel-addressing method used by some graphics packages, including OpenGL.

An object description that is input to a graphics program is given in terms of precise world-coordinate positions, which are infinitesimally small mathematical points. However, when the object is scan-converted into the frame buffer, the input description is transformed to pixel coordinates which reference finite screen
with constants $a$ and $b$ determined by the initial velocity $v_0$ of the object and the acceleration $g$ due to the uniform gravitational force. We can also describe such parabolic motions with parametric equations using a time parameter $t$, measured in seconds from the initial projection point:

$$
x = x_0 + v_{x0} t
$$
$$
y = y_0 + v_{y0} t - \frac{1}{2}gt^2
$$

(55)

Here, $v_{x0}$ and $v_{y0}$ are the initial velocity components, and the value of $g$ near the surface of the earth is approximately 980 cm/sec$^2$. Object positions along the parabolic path are then calculated at selected time steps.

Hyperbolic curves (Figure 26) are useful in various scientific-visualization applications. Motions of objects along hyperbolic paths occur in connection with the collision of charged particles and in certain gravitational problems. For example, comets or meteorites moving around the sun may travel along hyperbolic paths and escape to outer space, never to return. The particular branch (left or right, in Figure 26) describing the motion of an object depends on the forces involved in the problem. We can write the standard equation for the hyperbola centered on the origin in Figure 26 as

$$
\left( \frac{x}{r_x} \right)^2 - \left( \frac{y}{r_y} \right)^2 = 1
$$

(56)

with $x \leq -r_x$ for the left branch and $x \geq r_x$ for the right branch. Because this equation differs from the standard ellipse equation 39 only in the sign between the $x^2$ and $y^2$ terms, we can generate points along a hyperbolic path with a slightly modified ellipse algorithm.

Parabolas and hyperbolas possess a symmetry axis. For example, the parabola described by Equation 55 is symmetric about the axis

$$
x = x_0 + v_{x0}v_{y0}/g
$$

The methods used in the midpoint ellipse algorithm can be applied directly to obtain points along one side of the symmetry axis of hyperbolic and parabolic paths in the two regions: (1) where the magnitude of the curve slope is less than 1, and (2) where the magnitude of the slope is greater than 1. To do this, we first select the appropriate form of Equation 52 and then use the selected function to set up expressions for the decision parameters in the two regions.

**Polynomials and Spline Curves**

A polynomial function of $n$th degree in $x$ is defined as

$$
y = \sum_{k=0}^{n} a_k x^k
$$

$$
= a_0 + a_1 x + \cdots + a_{n-1} x^{n-1} + a_n x^n
$$

(57)

where $n$ is a nonnegative integer and the $a_k$ are constants, with $a_n \neq 0$. We obtain a quadratic curve when $n = 2$, a cubic polynomial when $n = 3$, a quartic curve when $n = 4$, and so forth. We have a straight line when $n = 1$. Polynomials are useful in a number of graphics applications, including the design of object shapes, the specification of animation paths, and the graphing of data trends in a discrete set of data points.

Designing object shapes or motion paths is typically accomplished by first specifying a few points to define the general curve contour, then the selected points are fitted with a polynomial. One way to accomplish the curve fitting is to
areas, and the displayed raster image may not correspond exactly with the relative dimensions of the input object. If it is important to preserve the specified geometry of world objects, we can compensate for the mapping of mathematical input points to finite pixel areas. One way to do this is simply to adjust the pixel dimensions of displayed objects so as to correspond to the dimensions given in the original mathematical description of the scene. For example, if a rectangle is specified as having a width of 40 cm, then we could adjust the screen display so that the rectangle has a width of 40 pixels, with the width of each pixel representing one centimeter. Another approach is to map world coordinates onto screen positions between pixels, so that we align object boundaries with pixel boundaries instead of pixel centers.

**Screen Grid Coordinates**

Figure 28 shows a screen section with grid lines marking pixel boundaries, one unit apart. In this scheme, a screen position is given as the pair of integer values identifying a grid-intersection position between two pixels. The address for any pixel is now at its lower-left corner, as illustrated in Figure 29. A straight-line path is now envisioned as between grid intersections. For example, the mathematical line path for a polyline with endpoint coordinates (0, 0), (5, 2), and (1, 4) would then be as shown in Figure 30.

Using screen grid coordinates, we now identify the area occupied by a pixel with screen coordinates \((x, y)\) as the unit square with diagonally opposite corners at \((x, y)\) and \((x + 1, y + 1)\). This pixel-addressing method has several advantages: it avoids half-integer pixel boundaries, it facilitates precise object representations, and it simplifies the processing involved in many scan-conversion algorithms and other raster procedures.

The algorithms for line drawing and curve generation discussed in the preceding sections are still valid when applied to input positions expressed as screen grid coordinates. Decision parameters in these algorithms would now be a measure of screen grid separation differences, rather than separation differences from pixel centers.

**Maintaining Geometric Properties of Displayed Objects**

When we convert geometric descriptions of objects into pixel representations, we transform mathematical points and lines into finite screen areas. If we are to maintain the original geometric measurements specified by the input coordinates for an object, we need to account for the finite size of pixels when we transform the object definition to a screen display.

Figure 31 shows the line plotted in the Bresenham line-algorithm example of Section 1. Interpreting the line endpoints \((20, 10)\) and \((30, 18)\) as precise grid-crossing positions, we see that the line should not extend past screen-grid position \((30, 18)\). If we were to plot the pixel with screen coordinates \((30, 18)\), as in the example given in Section 1, we would display a line that spans 11 horizontal units and 9 vertical units. For the mathematical line, however, \(\Delta x = 10\) and \(\Delta y = 8\). If we are addressing pixels by their center positions, we can adjust the length of the displayed line by omitting one of the endpoint pixels. But if we think of screen coordinates as addressing pixel boundaries, as shown in Figure 31, we plot a line using only those pixels that are “interior” to the line path; that is, only those pixels that are between the line endpoints. For our example, we would plot the leftmost pixel at \((20, 10)\) and the rightmost pixel at \((29, 17)\). This displays a
line that has the same geometric magnitudes as the mathematical line from (20, 10) to (30, 18).

For an enclosed area, input geometric properties are maintained by displaying the area using only those pixels that are interior to the object boundaries. The rectangle defined with the screen coordinate vertices shown in Figure 32(a), for example, is larger when we display it filled with pixels up to and including the border pixel lines joining the specified vertices [Figure 32(b)]. As defined, the area of the rectangle is 12 units, but as displayed in Figure 32(b), it has an area of 20 units. In Figure 32(c), the original rectangle measurements are maintained by displaying only the internal pixels. The right boundary of the input rectangle is at \( x = 4 \). To maintain the rectangle width in the display, we set the rightmost pixel grid coordinate for the rectangle at \( x = 3 \) because the pixels in this vertical column span the interval from \( x = 3 \) to \( x = 4 \). Similarly, the mathematical top boundary of the rectangle is at \( y = 3 \), so we set the top pixel row for the displayed rectangle at \( y = 2 \).

These compensations for finite pixel size can be applied to other objects, including those with curved boundaries, so that the raster display maintains the input object specifications. A circle with radius 5 and center position (10, 10), for instance, would be displayed as in Figure 33 by the midpoint circle algorithm.
using pixel centers as screen-coordinate positions. However, the plotted circle has a diameter of 11. To plot the circle with the defined diameter of 10, we can modify the circle algorithm to shorten each pixel scan line and each pixel column, as in Figure 34. One way to do this is to generate points clockwise along the circular arc in the third quadrant, starting at screen coordinates (10, 5). For each generated point, the other seven circle symmetry points are generated by decreasing the $x$ coordinate values by 1 along scan lines and decreasing the $y$ coordinate values by 1 along pixel columns. Similar methods are applied in ellipse algorithms to maintain the specified proportions in the display of an ellipse.

9 Attribute Implementations for Straight-Line Segments and Curves

Recall that line segment primitives can be displayed with three basic attributes: color, width, and style. Of these, line width and style are selected with separate line functions.

Line Width

Implementation of line-width options depends on the capabilities of the output device. For raster implementations, a standard-width line is generated with single pixels at each sample position, as in the Bresenham algorithm. Thicker lines are displayed as positive integer multiples of the standard line by plotting additional pixels along adjacent parallel line paths. If a line has slope magnitude less than or equal to 1.0, we can modify a line-drawing routine to display thick lines by plotting a vertical span of pixels in each column ($x$ position) along the line. The number of pixels to be displayed in each column is set equal to the integer value of the line width. In Figure 35, we display a double-width line by generating a parallel line above the original line path. At each $x$ sampling position, we calculate the corresponding $y$ coordinate and plot pixels at screen coordinates $(x, y)$ and $(x, y + 1)$. We could display lines with a width of 3 or greater by alternately plotting pixels above and below the single-width line path.
and add butt caps that are positioned half of the line width beyond the specified endpoints.

Other methods for producing thick lines include displaying the line as a filled rectangle or generating the line with a selected pen or brush pattern, as discussed in the next section. To obtain a rectangle representation for the line boundary, we calculate the position of the rectangle vertices along perpendiculars to the line path so that the rectangle vertex coordinates are displaced from the original line-endpoint positions by half the line width. The rectangular line then appears as in Figure 37(a). We could add round caps to the filled rectangle, or we could extend its length to display projecting square caps.

Generating thick polylines requires some additional considerations. In general, the methods that we have considered for displaying a single line segment will not produce a smoothly connected series of line segments. Displaying thick polylines using horizontal and vertical pixel spans, for example, leaves pixel gaps at the boundaries between line segments with different slopes where there is a shift from horizontal pixel spans to vertical spans. We can generate thick polylines that are smoothly joined at the cost of additional processing at the segment endpoints. Figure 38 shows three possible methods for smoothly joining two line segments. A miter join is accomplished by extending the outer boundaries of each of the two line segments until they meet. A round join is produced by capping the connection between the two segments with a circular boundary whose diameter is equal to the line width. A bevel join is generated by displaying the line segments with butt caps and filling in the triangular gap where the segments meet. If the angle between two connected line segments is very small, a miter join can generate a long spike that distorts the appearance of the polyline. A graphics package can avoid this effect by switching from a miter join to a bevel join when, for example, the angle between any two consecutive segments is small.

Line Style

Raster line algorithms display line-style attributes by plotting pixel spans. For dashed, dotted, and dot-dashed patterns, the line-drawing procedure outputs sections of contiguous pixels along the line path, skipping over a number of intervening pixels between the solid spans. Pixel counts for the span length and
With a line slope greater than 1.0 in magnitude, we can display thick lines using horizontal spans, alternately picking up pixels to the right and left of the line path. This scheme is demonstrated in Figure 36, where a line segment with a width of 4 is plotted using multiple pixels across each scan line. Similarly, a thick line with slope less than or equal to 1.0 can be displayed using vertical pixel spans. We can implement this procedure by comparing the magnitudes of the horizontal and vertical separations ($\Delta x$ and $\Delta y$) of the line endpoints. If $|\Delta x| \geq |\Delta y|$, pixels are replicated along columns. Otherwise, multiple pixels are plotted across rows.

Although thick lines are generated quickly by plotting horizontal or vertical pixel spans, the displayed width of a line (measured perpendicular to the line path) depends on its slope. A $45^\circ$ line will be displayed thinner by a factor of $1/\sqrt{2}$ compared to a horizontal or vertical line plotted with the same-length pixel spans.

Another problem with implementing width options using horizontal or vertical pixel spans is that the method produces lines whose ends are horizontal or vertical regardless of the slope of the line. This effect is more noticeable with very thick lines. We can adjust the shape of the line ends to give them a better appearance by adding line caps (Figure 37). One kind of line cap is the butt cap, which has square ends that are perpendicular to the line path. If the specified line has slope $m$, the square ends of the thick line have slope $-1/m$. Each of the component parallel lines is then displayed between the two perpendicular lines at each end of the specified line path. Another line cap is the round cap obtained by adding a filled semicircle to each butt cap. The circular arcs are centered at the middle of the thick line and have a diameter equal to the line thickness. A third type of line cap is the projecting square cap. Here, we simply extend the line
inter-span spacing can be specified in a **pixel mask**, which is a pattern of binary digits indicating which positions to plot along the line path. The linear mask 11110000, for instance, could be used to display a dashed line with a dash length of five pixels and an inter-dash spacing of three pixels. Pixel positions corresponding to the 1 bits are assigned the current color, and pixel positions corresponding to the 0 bits are displayed in the background color.

Plotting dashes with a fixed number of pixels results in unequal length dashes for different line orientations, as illustrated in Figure 39. Both dashes shown are plotted with four pixels, but the diagonal dash is longer by a factor of $\sqrt{2}$. For precision drawings, dash lengths should remain approximately constant for any line orientation. To accomplish this, we could adjust the pixel counts for the solid spans and inter-span spacing according to the line slope. In Figure 39, we can display approximately equal length dashes by reducing the diagonal dash to three pixels. Another method for maintaining dash length is to treat dashes as individual line segments. Endpoint coordinates for each dash are located and passed to the line routine, which then calculates pixel positions along the dash path.

**Pen and Brush Options**

Pen and brush shapes can be stored in a pixel mask that identifies the array of pixel positions that are to be set along the line path. For example, a rectangular pen could be implemented with the mask shown in Figure 40 by moving the center (or one corner) of the mask along the line path, as in Figure 41. To avoid setting pixels more than once in the frame buffer, we can simply accumulate the horizontal spans generated at each position of the mask and keep track of the beginning and ending $x$ positions for the spans across each scan line.

Lines generated with pen (or brush) shapes can be displayed in various widths by changing the size of the mask. For example, the rectangular pen line in Figure 41 could be narrowed with a $2 \times 2$ rectangular mask or widened with a $4 \times 4$ mask. Also, lines can be displayed with selected patterns by superimposing the pattern values onto the pen or brush mask.
Curve Attributes

Methods for adapting curve-drawing algorithms to accommodate attribute selections are similar to those for line drawing. Raster curves of various widths can be displayed using the method of horizontal or vertical pixel spans. Where the magnitude of the curve slope is less than or equal to 1.0, we plot vertical spans; where the slope magnitude is greater than 1.0, we plot horizontal spans. Figure 42 demonstrates this method for displaying a circular arc with a width of 4 in the first quadrant. Using circle symmetry, we generate the circle path with vertical spans in the octant from \( x = 0 \) to \( x = y \), and then reflect pixel positions about the line \( y = x \) to obtain the remainder of the curve shown. Circle sections in the other quadrants are obtained by reflecting pixel positions in the first quadrant about the coordinate axes. The thickness of curves displayed with this method is again a function of curve slope. Circles, ellipses, and other curves will appear thinnest where the slope has a magnitude of 1.

Another method for displaying thick curves is to fill in the area between two parallel curve paths, whose separation distance is equal to the desired width. We could do this using the specified curve path as one boundary and setting up the second boundary either inside or outside the original curve path. This approach, however, shifts the original curve path either inward or outward, depending on which direction we choose for the second boundary. We can maintain the original curve position by setting the two boundary curves at a distance of half the width on either side of the specified curve path. An example of this approach is shown in Figure 43 for a circle segment with a radius of 16 and a specified width of 4. The boundary arcs are then set at a separation distance of 2 on either side of the radius of 16. To maintain the proper dimensions of the circular arc, as discussed in Section 8, we can set the radii for the concentric boundary arcs at \( r = 14 \) and \( r = 17 \). Although this method is accurate for generating thick circles, it provides, in general, only an approximation to the true area of other thick curves. For example, the inner and outer boundaries of a fat ellipse generated with this method do not have the same foci.

The pixel masks discussed for implementing line-style options could also be used in raster curve algorithms to generate dashed or dotted patterns. For example, the mask 11100 produces the dashed circular arc shown in Figure 44.
FIGURE 44  A dashed circular arc displayed with a dash span of 3 pixels and an inter-dash spacing of 2 pixels.

FIGURE 45  A circular arc displayed with a rectangular pen.

We can generate the dashes in the various octants using circle symmetry, but we must shift the pixel positions to maintain the correct sequence of dashes and spaces as we move from one octant to the next. Also, as in straight-line algorithms, pixel masks display dashes and inter-dash spaces that vary in length according to the slope of the curve. If we want to display constant length dashes, we need to adjust the number of pixels plotted in each dash as we move around the circle circumference. Instead of applying a pixel mask with constant spans, we plot pixels along equal angular arcs to produce equal-length dashes.

Pen (or brush) displays of curves are generated using the same techniques discussed for straight-line segments. We replicate a pen shape along the line path, as illustrated in Figure 45 for a circular arc in the first quadrant. Here, the center of the rectangular pen is moved to successive curve positions to produce the curve shape shown. Curves displayed with a rectangular pen in this manner will be thicker where the magnitude of the curve slope is 1. A uniform curve thickness can be displayed by rotating the rectangular pen to align it with the slope direction as we move around the curve or by using a circular pen shape. Curves drawn with pen and brush shapes can be displayed in different sizes and with superimposed patterns or simulated brush strokes.

10 General Scan-Line Polygon-Fill Algorithm

A scan-line fill of a region is performed by first determining the intersection positions of the boundaries of the fill region with the screen scan lines. Then the fill colors are applied to each section of a scan line that lies within the interior of the fill region. The scan-line fill algorithm identifies the same interior regions as the odd-even rule. The simplest area to fill is a polygon because each scan-line intersection point with a polygon boundary is obtained by solving a pair of simultaneous linear equations, where the equation for the scan line is simply \( y = \text{constant} \).

Figure 46 illustrates the basic scan-line procedure for a solid-color fill of a polygon. For each scan line that crosses the polygon, the edge intersections are
sorted from left to right, and then the pixel positions between, and including, each intersection pair are set to the specified fill color. In the example of Figure 46, the four pixel intersection positions with the polygon boundaries define two stretches of interior pixels. Thus, the fill color is applied to the five pixels from \(x = 10\) to \(x = 14\) and to the seven pixels from \(x = 18\) to \(x = 24\). If a pattern fill is to be applied to the polygon, then the color for each pixel along a scan line is determined from its overlap position with the fill pattern.

However, the scan-line fill algorithm for a polygon is not quite as simple as Figure 46 might suggest. Whenever a scan line passes through a vertex, it intersects two polygon edges at that point. In some cases, this can result in an odd number of boundary intersections for a scan line. Figure 47 shows two scan lines that cross a polygon fill area and intersect a vertex. Scan line \(y\) intersects an even number of edges, and the two pairs of intersection points along this scan line correctly identify the interior pixel spans. But scan line \(y'\) intersects five polygon edges. To identify the interior pixels for scan line \(y\), we must count the vertex intersection as only one point. Thus, as we process scan lines, we need to distinguish between these cases.

We can detect the topological difference between scan line \(y\) and scan line \(y'\) in Figure 47 by noting the position of the intersecting edges relative to the scan line. For scan line \(y\), the two edges sharing an intersection vertex are on opposite sides of the scan line. But for scan line \(y'\), the two intersecting edges are both above the scan line. Thus, a vertex that has adjoining edges on opposite sides of an intersecting scan line should be counted as just one boundary intersection point. We can identify these vertices by tracing around the polygon boundary in either clockwise or counterclockwise order and observing the relative changes in vertex \(y\) coordinates as we move from one edge to the next. If the three endpoint \(y\) values of two consecutive edges monotonically increase or decrease, we need to count the shared (middle) vertex as a single intersection point for the scan line passing through that vertex. Otherwise, the shared vertex represents a local extremum (minimum or maximum) on the polygon boundary, and the two edge intersections with the scan line passing through that vertex can be added to the intersection list.

One method for implementing the adjustment to the vertex-intersection count is to shorten some polygon edges to split those vertices that should be counted as one intersection. We can process nonhorizontal edges around the polygon boundary in the order specified, either clockwise or counterclockwise. As we...
process each edge, we can check to determine whether that edge and the next nonhorizontal edge have either monotonically increasing or decreasing endpoint $y$ values. If so, the lower edge can be shortened to ensure that only one intersection point is generated for the scan line going through the common vertex joining the two edges. Figure 48 illustrates the shortening of an edge. When the endpoint $y$ coordinates of the two edges are increasing, the $y$ value of the upper endpoint for the current edge is decreased by 1, as in Figure 48(a). When the endpoint $y$ values are monotonically decreasing, as in Figure 48(b), we decrease the $y$ coordinate of the upper endpoint of the edge following the current edge.

Typically, certain properties of one part of a scene are related in some way to the properties in other parts of the scene, and these coherence properties can be used in computer-graphics algorithms to reduce processing. Coherence methods often involve incremental calculations applied along a single scan line or between successive scan lines. For example, in determining fill-area edge intersections, we can set up incremental coordinate calculations along any edge by exploiting the fact that the slope of the edge is constant from one scan line to the next. Figure 49 shows two successive scan lines crossing the left edge of a triangle. The slope of this edge can be expressed in terms of the scan-line intersection coordinates:

$$m = \frac{y_{k+1} - y_k}{x_{k+1} - x_k}$$

Because the change in $y$ coordinates between the two scan lines is simply

$$y_{k+1} - y_k = 1$$

the $x$-intersection value $x_{k+1}$ on the upper scan line can be determined from the $x$-intersection value $x_k$ on the preceding scan line as

$$x_{k+1} = x_k + \frac{1}{m}$$

Each successive $x$ intercept can thus be calculated by adding the inverse of the slope and rounding to the nearest integer.
An obvious parallel implementation of the fill algorithm is to assign each scan line that crosses the polygon to a separate processor. Edge intersection calculations are then performed independently. Along an edge with slope $m$, the intersection $x_k$ value for scan line $k$ above the initial scan line can be calculated as

$$x_k = x_0 + \frac{k}{m} \quad (62)$$

In a sequential fill algorithm, the increment of $x$ values by the amount $\frac{1}{m}$ along an edge can be accomplished with integer operations by recalling that the slope $m$ is the ratio of two integers:

$$m = \frac{\Delta y}{\Delta x}$$

where $\Delta x$ and $\Delta y$ are the differences between the edge endpoint $x$ and $y$ coordinate values. Thus, incremental calculations of $x$ intercepts along an edge for successive scan lines can be expressed as

$$x_{k+1} = x_k + \frac{\Delta x}{\Delta y} \quad (63)$$

Using this equation, we can perform integer evaluation of the $x$ intercepts by initializing a counter to 0, then incrementing the counter by the value of $\Delta x$ each time we move up to a new scan line. Whenever the counter value becomes equal to or greater than $\Delta y$, we increment the current $x$ intersection value by 1 and decrease the counter by the value $\Delta y$. This procedure is equivalent to maintaining integer and fractional parts for $x$ intercepts and incrementing the fractional part until we reach the next integer value.

As an example of this integer-incrementing scheme, suppose that we have an edge with slope $m = \frac{7}{3}$. At the initial scan line, we set the counter to 0 and the counter increment to 3. As we move up to the next three scan lines along this edge, the counter is successively assigned the values 3, 6, and 9. On the third scan line above the initial scan line, the counter now has a value greater than 7. So we increment the $x$ intersection coordinate by 1 and reset the counter to the value $9 - 7 = 2$. We continue determining the scan-line intersections in this way until we reach the upper endpoint of the edge. Similar calculations are carried out to obtain intersections for edges with negative slopes.

We can round to the nearest pixel $x$ intersection value, instead of truncating to obtain integer positions, by modifying the edge-intersection algorithm so that the increment is compared to $\Delta y/2$. This can be done with integer arithmetic by incrementing the counter with the value $2\Delta x$ at each step and comparing the increment to $\Delta y$. When the increment is greater than or equal to $\Delta y$, we increase the $x$ value by 1 and decrement the counter by the value of $2\Delta y$. In our previous example with $m = \frac{7}{3}$, the counter values for the first few scan lines above the initial scan line on this edge would now be 6, 12 (reduced to $-2$), 4, 10 (reduced to $-4$), 2, 8 (reduced to $-6$), 0, 6, and 12 (reduced to $-2$). Now $x$ would be incremented on scan lines 2, 4, 6, 9, and so forth, above the initial scan line for this edge. The extra calculations required for each edge are $2\Delta x = \Delta x + \Delta x$ and $2\Delta y = \Delta y + \Delta y$, which are carried out as preprocessing steps.

To perform a polygon fill efficiently, we can first store the polygon boundary in a sorted edge table that contains all the information necessary to process the scan lines efficiently. Proceeding around the edges in either a clockwise or a counterclockwise order, we can use a bucket sort to store the edges, sorted on the smallest $y$ value of each edge, in the correct scan-line positions. Only nonhorizontal edges are entered into the sorted edge table. As the edges are processed, we can also
shorten certain edges to resolve the vertex-intersection question. Each entry in the table for a particular scan line contains the maximum y value for that edge, the x-intercept value (at the lower vertex) for the edge, and the inverse slope of the edge. For each scan line, the edges are in sorted order from left to right. Figure 50 shows a polygon and the associated sorted edge table.

Next, we process the scan lines from the bottom of the polygon to its top, producing an active edge list for each scan line crossing the polygon boundaries. The active edge list for a scan line contains all edges crossed by that scan line, with iterative coherence calculations used to obtain the edge intersections.

Implementation of edge-intersection calculations can be facilitated by storing $\Delta x$ and $\Delta y$ values in the sorted edge list. Also, to ensure that we correctly fill the interior of specified polygons, we can apply the considerations discussed in Section 8. For each scan line, we fill in the pixel spans for each pair of x intercepts starting from the leftmost x intercept value and ending at one position before the rightmost x intercept. Each polygon edge can be shortened by one unit in the y direction at the top endpoint. These measures also guarantee that pixels in adjacent polygons will not overlap.

11 Scan-Line Fill of Convex Polygons

When we apply a scan-line fill procedure to a convex polygon, there can be no more than a single interior span for each screen scan line. So we need to process the polygon edges only until we have found two boundary intersections for each scan line crossing the polygon interior.

The general polygon scan-line algorithm discussed in the preceding section can be simplified considerably for convex-polygon fill. We again use coordinate extents to determine which edges cross a scan line. Intersection calculations with these edges then determine the interior pixel span for that scan line, where any
vertex crossing is counted as a single boundary intersection point. When a scan line intersects a single vertex (at an apex, for example), we plot only that point. Some graphics packages further restrict fill areas to be triangles. This makes filling even easier because each triangle has just three edges to process.

12 Scan-Line Fill for Regions with Curved Boundaries

Because an area with curved boundaries is described with nonlinear equations, a scan-line fill generally takes more time than a polygon scan-line fill. We can use the same general approach detailed in Section 10, but the boundary intersection calculations are performed with curve equations. In addition, the slope of the boundary is continuously changing, so we cannot use the straightforward incremental calculations that are possible with straight-line edges.

For simple curves such as circles or ellipses, we can apply fill methods similar to those for convex polygons. Each scan line crossing a circle or ellipse interior has just two boundary intersections; and we can determine these two intersection points along the boundary of a circle or an ellipse using the incremental calculations in the midpoint method. Then we simply fill in the horizontal pixel spans from one intersection point to the other. Symmetries between quadrants (and between octants for circles) are used to reduce the boundary calculations.

Similar methods can be used to generate a fill area for a curve section. For example, an area bounded by an elliptical arc and a straight line section (Figure 51) can be filled using a combination of curve and line procedures. Symmetries and incremental calculations are exploited whenever possible to reduce computations.

Filling other curve areas can involve considerably more processing. We could use similar incremental methods in combination with numerical techniques to determine the scan-line intersections, but usually such curve boundaries are approximated with straight-line segments.

13 Fill Methods for Areas with Irregular Boundaries

Another approach for filling a specified area is to start at an inside position and “paint” the interior, point by point, out to the boundary. This is a particularly useful technique for filling areas with irregular borders, such as a design created with a paint program. Generally, these methods require an input starting position inside the area to be filled and some color information about either the boundary or the interior.

We can fill irregular regions with a single color or with a color pattern. For a pattern fill, we overlay a color mask. As each pixel within the region is processed, its color is determined by the corresponding values in the overlaid pattern.

Boundary-Fill Algorithm

If the boundary of some region is specified in a single color, we can fill the interior of this region, pixel by pixel, until the boundary color is encountered. This
method, called the **boundary-fill algorithm**, is employed in interactive painting packages, where interior points are easily selected. Using a graphics tablet or other interactive device, an artist or designer can sketch a figure outline, select a fill color from a color menu, specify the area boundary color, and pick an interior point. The figure interior is then painted in the fill color. Both inner and outer boundaries can be set up to define an area for boundary fill, and Figure 52 illustrates examples for specifying color regions.

Basically, a boundary-fill algorithm starts from an interior point \((x, y)\) and tests the color of neighboring positions. If a tested position is not displayed in the boundary color, its color is changed to the fill color and its neighbors are tested. This procedure continues until all pixels are processed up to the designated boundary color for the area.

Figure 53 shows two methods for processing neighboring pixels from a current test position. In Figure 53(a), four neighboring points are tested. These are the pixel positions that are right, left, above, and below the current pixel. Areas filled by this method are called **4-connected**. The second method, shown in Figure 53(b), is used to fill more complex figures. Here the set of neighboring positions to be tested includes the four diagonal pixels, as well as those in the cardinal directions. Fill methods using this approach are called **8-connected**. An 8-connected boundary-fill algorithm would correctly fill the interior of the area defined in Figure 54, but a 4-connected boundary-fill algorithm would fill only part of that region.

The following procedure illustrates a recursive method for painting a 4-connected area with a solid color, specified in parameter `fillColor`, up to a boundary color specified with parameter `borderColor`. We can extend this procedure to fill an 8-connected region by including four additional statements to test the diagonal positions \((x \pm 1, y \pm 1)\).

![Figure 52](image1.png)  
*Example of color boundaries for a boundary-fill procedure.*

![Figure 53](image2.png)  
*Fill methods applied to a 4-connected area (a) and to an 8-connected area (b). Hollow circles represent pixels to be tested from the current test position, shown as a solid color.*

![Figure 54](image3.png)  
The area defined within the color boundary (a) is only partially filled in (b) using a 4-connected boundary-fill algorithm.
Recursive boundary-fill algorithms may not fill regions correctly if some interior pixels are already displayed in the fill color. This occurs because the algorithm checks the next pixels both for boundary color and for fill color. Encountering a pixel with the fill color can cause a recursive branch to terminate, leaving other interior pixels unfilled. To avoid this, we can first change the color of any interior pixels that are initially set to the fill color before applying the boundary-fill procedure.

Also, because this procedure requires considerable stacking of neighboring points, more efficient methods are generally employed. These methods fill horizontal pixel spans across scan lines, instead of proceeding to 4-connected or 8-connected neighboring points. Then we need only stack a beginning position for each horizontal pixel span, instead of stacking all unprocessed neighboring positions around the current position. Starting from the initial interior point with this method, we first fill in the contiguous span of pixels on this starting scan line. Then we locate and stack starting positions for spans on the adjacent scan lines, where spans are defined as the contiguous horizontal string of positions bounded by pixels displayed in the border color. At each subsequent step, we retrieve the next start position from the top of the stack and repeat the process.

An example of how pixel spans could be filled using this approach is illustrated for the 4-connected fill region in Figure 55. In this example, we first process scan lines successively from the start line to the top boundary. After all upper scan lines are processed, we fill in the pixel spans on the remaining scan lines in order down to the bottom boundary. The leftmost pixel position for each horizontal span is located and stacked, in left-to-right order across successive scan lines, as shown in Figure 55. In (a) of this figure, the initial span has been filled, and starting positions 1 and 2 for spans on the next scan lines (below and above) are stacked. In Figure 55(b), position 2 has been unstacked and processed to produce the filled span shown, and the starting pixel (position 3) for the single span on the next scan line has been stacked. After position 3 is processed, the filled spans and stacked positions are as shown in Figure 55(c). Figure 55(d) shows the filled pixels after processing all spans in the upper-right portion of the specified area. Position 5 is next processed, and spans are filled in the upper-left portion of the region; then position 4 is picked up to continue the processing for the lower scan lines.

```c
void boundaryFill4 (int x, int y, int fillColor, int borderColor)
{
    int interiorColor;
    /* Set current color to fillColor, then perform the following operations. */
    getPixel (x, y, interiorColor);
    if ((interiorColor != borderColor) && (interiorColor != fillColor)) {
        setPixel (x, y); // Set color of pixel to fillColor.
        boundaryFill4 (x + 1, y, fillColor, borderColor);
        boundaryFill4 (x - 1, y, fillColor, borderColor);
        boundaryFill4 (x, y + 1, fillColor, borderColor);
        boundaryFill4 (x, y - 1, fillColor, borderColor)
    }
}
```
Boundary fill across pixel spans for a 4-connected area: (a) Initial scan line with a filled pixel span, showing the position of the initial point (hollow) and the stacked positions for pixel spans on adjacent scan lines. (b) Filled pixel span on the first scan line above the initial scan line and the current contents of the stack. (c) Filled pixel spans on the first two scan lines above the initial scan line and the current contents of the stack. (d) Completed pixel spans for the upper-right portion of the defined region and the remaining stacked positions to be processed.
Flood-Fill Algorithm

Sometimes we want to fill in (or recolor) an area that is not defined within a single color boundary. Figure 56 shows an area bordered by several different color regions. We can paint such areas by replacing a specified interior color instead of searching for a particular boundary color. This fill procedure is called a flood-fill algorithm. We start from a specified interior point \((x, y)\) and reassign all pixel values that are currently set to a given interior color with the desired fill color. If the area that we want to paint has more than one interior color, we can first reassign pixel values so that all interior points have the same color. Using either a 4-connected or 8-connected approach, we then step through pixel positions until all interior points have been repainted. The following procedure flood fills a 4-connected region recursively, starting from the input position.

```c
void floodFill4 (int x, int y, int fillColor, int interiorColor)
{
    int color;

    /* Set current color to fillColor, then perform the following operations. */
    getPixel (x, y, color);
    if (color = interiorColor) {
        setPixel (x, y); // Set color of pixel to fillColor.
        floodFill4 (x + 1, y, fillColor, interiorColor);
        floodFill4 (x - 1, y, fillColor, interiorColor);
        floodFill4 (x, y + 1, fillColor, interiorColor);
        floodFill4 (x, y - 1, fillColor, interiorColor)
    }
}
```

We can modify the above procedure to reduce the storage requirements of the stack by filling horizontal pixel spans, as discussed for the boundary-fill algorithm. In this approach, we stack only the beginning positions for those pixel spans having the value `interiorColor`. The steps in this modified flood-fill algorithm are similar to those illustrated in Figure 55 for a boundary fill. Starting at the first position of each span, the pixel values are replaced until a value other than `interiorColor` is encountered.

14 Implementation Methods for Fill Styles

There are two basic procedures for filling an area on raster systems, once the definition of the fill region has been mapped to pixel coordinates. One procedure first determines the overlap intervals for scan lines that cross the area. Then, pixel positions along these overlap intervals are set to the fill color. Another method for area filling is to start from a given interior position and “paint” outward, pixel-by-pixel, from this point until we encounter specified boundary conditions. The scan-line approach is usually applied to simple shapes such as circles or regions with polyline boundaries, and general graphics packages use this fill method. Fill algorithms that use a starting interior point are useful for filling areas with more complex boundaries and in interactive painting systems.
Fill Styles

We can implement a pattern fill by determining where the pattern overlaps those scan lines that cross a fill area. Beginning from a specified start position for a pattern fill, we map the rectangular patterns vertically across scan lines and horizontally across pixel positions on the scan lines. Each replication of the pattern array is performed at intervals determined by the width and height of the mask. Where the pattern overlaps the fill area, pixel colors are set according to the values stored in the mask.

Hatch fill could be applied to regions by drawing sets of line segments to display either single hatching or cross-hatching. Spacing and slope for the hatch lines could be set as parameters in a hatch table. Alternatively, hatch fill can be specified as a pattern array that produces sets of diagonal lines.

A reference point \((xp, yp)\) for the starting position of a fill pattern can be set at any convenient position, inside or outside the fill region. For instance, the reference point could be set at a polygon vertex; or the reference point could be chosen as the lower-left corner of the bounding rectangle (or bounding box) determined by the coordinate extents of the region. To simplify selection of the reference coordinates, some packages always use the coordinate origin of the display window as the pattern start position. Always setting \((xp, yp)\) at the coordinate origin also simplifies the tiling operations when each element of a pattern is to be mapped to a single pixel. For example, if the row positions in the pattern array are referenced from bottom to top, starting with the value 1, a color value is then assigned to pixel position \((x, y)\) in screen coordinates from pattern position \((y \mod ny + 1, x \mod nx + 1)\). Here, \(ny\) and \(nx\) specify the number of rows and number of columns in the pattern array. Setting the pattern start position at the coordinate origin, however, effectively attaches the pattern fill to the screen background rather than to the fill regions. Adjacent or overlapping areas filled with the same pattern would show no apparent boundary between the areas. Also, repositioning and refilling an object with the same pattern can result in a shift in the assigned pixel values over the object interior. A moving object would appear to be transparent against a stationary pattern background instead of moving with a fixed interior pattern.

Color-Blended Fill Regions

Color-blended regions can be implemented using either transparency factors to control the blending of background and object colors, or using simple logical or replace operations as shown in Figure 57, which demonstrates how these operations would combine a \(2 \times 2\) fill pattern with a background pattern for a binary (black-and-white) system.

The linear soft-fill algorithm repaints an area that was originally painted by merging a foreground color \(F\) with a single background color \(B\), where \(F \neq B\). Assuming we know the values for \(F\) and \(B\), we can check the current contents of the frame buffer to determine how these colors were combined. The current color \(P\) of each pixel within the area to be refilled is some linear combination of \(F\) and \(B\):

\[
P = tF + (1 - t)B
\]

(64)

where the transparency factor \(t\) has a value between 0 and 1 for each pixel. For values of \(t\) less than 0.5, the background color contributes more to the interior color of the region than does the fill color. If our color values are represented using separate red, green, and blue (RGB) components, Equation 64 holds for
FIGURE 57
Combining a fill pattern with a background pattern using logical operations and, or, and xor (exclusive or), and using simple replacement.

We can thus calculate the value of parameter \( t \) using one of the RGB color components as follows:

\[
\begin{align*}
    t &= \frac{P_k - B_k}{F_k - B_k} \\
    \end{align*}
\]  

(66)

where \( k = R, G, \text{or } B; \text{ and } F_k \neq B_k \). Theoretically, parameter \( t \) has the same value for each RGB component, but the round-off calculations to obtain integer codes can result in different values of \( t \) for different components. We can minimize this round-off error by selecting the component with the largest difference between \( F \) and \( B \). This value of \( t \) is then used to mix the new fill color \( NF \) with the background color. We can accomplish this mixing using either a modified flood-fill or boundary-fill procedure, as described in Section 13.

Similar color-blending procedures can be applied to an area whose foreground color is to be merged with multiple background color areas, such as a checkerboard pattern. When two background colors \( B_1 \) and \( B_2 \) are mixed with foreground color \( F \), the resulting pixel color \( P \) is

\[
\begin{align*}
    P &= t_0F + t_1B_1 + (1 - t_0 - t_1)B_2 \\
    \end{align*}
\]  

(67)

where the sum of the color-term coefficients \( t_0, t_1, \) and \( (1 - t_0 - t_1) \) must equal 1. We can set up two simultaneous equations using two of the three RGB color components to solve for the two proportionality parameters, \( t_0 \) and \( t_1 \). These parameters are then used to mix the new fill color with the two background colors to obtain the new pixel color. With three background colors and one foreground color, or with two background and two foreground colors, we need all three RGB equations to obtain the relative amounts of the four colors. For some foreground and background color combinations, however, the system of two or three RGB
equations cannot be solved. This occurs when the color values are all very similar or when they are all proportional to each other.

15 Implementation Methods for Antialiasing

Line segments and other graphics primitives generated by the raster algorithms discussed earlier in this chapter have a jagged, or stair-step, appearance because the sampling process digitizes coordinate points on an object to discrete integer pixel positions. This distortion of information due to low-frequency sampling (undersampling) is called aliasing. We can improve the appearance of displayed raster lines by applying antialiasing methods that compensate for the undersampling process.

An example of the effects of undersampling is shown in Figure 58. To avoid losing information from such periodic objects, we need to set the sampling frequency to at least twice that of the highest frequency occurring in the object, referred to as the Nyquist sampling frequency (or Nyquist sampling rate) $f_s$:

\[ f_s = 2f_{\text{max}} \]  

Another way to state this is that the sampling interval should be no larger than one-half the cycle interval (called the Nyquist sampling interval). For $x$-interval sampling, the Nyquist sampling interval $\Delta x_s$ is

\[ \Delta x_s = \frac{\Delta x_{\text{cycle}}}{2} \]  

where $\Delta x_{\text{cycle}} = 1/f_{\text{max}}$. In Figure 58, our sampling interval is one and one-half times the cycle interval, so the sampling interval is at least three times too large. If we want to recover all the object information for this example, we need to cut the sampling interval down to one-third the size shown in the figure.

One way to increase sampling rate with raster systems is simply to display objects at higher resolution. However, even at the highest resolution possible with current technology, the jaggies will be apparent to some extent. There is a limit to how big we can make the frame buffer and still maintain the refresh rate at 60 frames or more per second. To represent objects accurately with continuous parameters, we need arbitrarily small sampling intervals. Therefore, unless hardware technology is developed to handle arbitrarily large frame buffers, increased screen resolution is not a complete solution to the aliasing problem.

With raster systems that are capable of displaying more than two intensity levels per color, we can apply antialiasing methods to modify pixel intensities. By
appropriately varying the intensities of pixels along the boundaries of primitives, we can smooth the edges to lessen their jagged appearance.

A straightforward antialiasing method is to increase sampling rate by treating the screen as if it were covered with a finer grid than is actually available. We can then use multiple sample points across this finer grid to determine an appropriate intensity level for each screen pixel. This technique of sampling object characteristics at a high resolution and displaying the results at a lower resolution is called \textit{supersampling} (or \textit{postfiltering}, because the general method involves computing intensities at subpixel grid positions and then combining the results to obtain the pixel intensities). Displayed pixel positions are spots of light covering a finite area of the screen, and not infinitesimal mathematical points. Yet in the line and fill-area algorithms we have discussed, the intensity of each pixel is determined by the location of a single point on the object boundary. By supersampling, we obtain intensity information from multiple points that contribute to the overall intensity of a pixel.

An alternative to supersampling is to determine pixel intensity by calculating the areas of overlap of each pixel with the objects to be displayed. Antialiasing by computing overlap areas is referred to as \textit{area sampling} (or \textit{prefiltering}, because the intensity of the pixel as a whole is determined without calculating subpixel intensities). Pixel overlap areas are obtained by determining where object boundaries intersect individual pixel boundaries.

Raster objects can also be antialiased by shifting the display location of pixel areas. This technique, called \textit{pixel phasing}, is applied by “micropositioning” the electron beam in relation to object geometry. For example, pixel positions along a straight-line segment can be moved closer to the defined line path to smooth out the raster stair-step effect.

\textbf{Supersampling Straight-Line Segments}

We can perform supersampling in several ways. For a straight-line segment, we can divide each pixel into a number of subpixels and count the number of subpixels that overlap the line path. The intensity level for each pixel is then set to a value that is proportional to this subpixel count. An example of this method is given in Figure 59. Each square pixel area is divided into nine equal-sized square subpixels, and the shaded regions show the subpixels that would be selected by Bresenham’s algorithm. This scheme provides for three intensity settings above zero, because the maximum number of subpixels that can be selected within any pixel is three. For this example, the pixel at position (10, 20) is set to the maximum intensity (level 3); pixels at (11, 21) and (12, 21) are each set to the next highest intensity (level 2); and pixels at (11, 20) and (12, 22) are each set to the lowest intensity above zero (level 1). Thus, the line intensity is spread out over a greater number of pixels to smooth the original jagged effect. This procedure displays a somewhat blurred line in the vicinity of the stair steps (between horizontal runs). If we want to use more intensity levels to antialias the line with this method, we increase the number of sampling positions across each pixel. Sixteen subpixels gives us four intensity levels above zero; twenty-five subpixels gives us five levels; and so on.

In the supersampling example of Figure 59, we considered pixel areas of finite size, but we treated the line as a mathematical entity with zero width. Actually, displayed lines have a width approximately equal to that of a pixel. If we take the finite width of the line into account, we can perform supersampling by setting pixel intensity proportional to the number of subpixels inside the polygon representing the line area. A subpixel can be considered to be inside the...
line if its lower-left corner is inside the polygon boundaries. An advantage of this supersampling procedure is that the number of possible intensity levels for each pixel is equal to the total number of subpixels within the pixel area. For the example in Figure 59, we can represent this line with finite width by positioning the polygon boundaries parallel to the line path as in Figure 60. In addition, each pixel can now be set to one of nine possible brightness levels above zero.

Another advantage of supersampling with a finite-width line is that the total line intensity is distributed over more pixels. In Figure 60, we now have the pixel at grid position (10, 21) turned on (at intensity level 1), and we also pick up contributions from pixels immediately below and immediately to the left of position (10, 21). Also, if we have a color display, we can extend the method to take background colors into account. A particular line might cross several different color areas, and we can average subpixel intensities to obtain pixel color settings. For instance, if five subpixels within a particular pixel area are determined to be inside the boundaries for a red line and the remaining four subpixels fall within a blue background area, we can calculate the color for this pixel as

\[
\text{pixel color} = \frac{(5 \cdot \text{red} + 4 \cdot \text{blue})}{9}
\]

The trade-off for these gains from supersampling a finite-width line is that identifying interior subpixels requires more calculations than simply determining which subpixels are along the line path. Also, we need to take into account the positioning of the line boundaries in relation to the line path. This positioning depends on the slope of the line. For a 45° line, the line path is centered on the polygon area; but for either a horizontal or a vertical line, we want the line path to be one of the polygon boundaries. For example, a horizontal line passing through grid coordinates (10, 20) could be represented as the polygon bounded by horizontal grid lines \( y = 20 \) and \( y = 21 \). Similarly, the polygon representing a vertical line through (10, 20) can have vertical boundaries along grid lines \( x = 10 \) and \( x = 11 \). For lines with slope \( |m| < 1 \), the mathematical line path will be positioned proportionately closer to either the lower or the upper polygon boundary depending upon where the line intersects the polygon; in Figure 59, for instance, the line intersects the pixel at (10, 20) closer to the lower boundary, but intersects the pixel at (11, 20) closer to the upper boundary. Similarly, for lines with slope \( |m| > 1 \), the line path is placed closer to the left or right polygon boundary depending on where it intersects the polygon.
Subpixel Weighting Masks

Supersampling algorithms are often implemented by giving more weight to subpixels near the center of a pixel area because we would expect these subpixels to be more important in determining the overall intensity of a pixel. For the $3 \times 3$ pixel subdivisions we have considered so far, a weighting scheme as in Figure 61 could be used. The center subpixel here is weighted four times that of the corner subpixels and twice that of the remaining subpixels. Intensities calculated for each of the nine subpixels would then be averaged so that the center subpixel is weighted by a factor of $\frac{1}{4}$; the top, bottom, and side subpixels are each weighted by a factor of $\frac{1}{2}$; and the corner subpixels are each weighted by a factor of $\frac{1}{16}$. An array of values specifying the relative importance of subpixels is usually referred to as a weighting mask. Similar masks can be set up for larger subpixel grids. Also, these masks are often extended to include contributions from subpixels belonging to neighboring pixels, so that intensities can be averaged with adjacent pixels to provide a smoother intensity variation between pixels.

Area Sampling Straight-Line Segments

We perform area sampling for a straight line by setting pixel intensity proportional to the area of overlap of the pixel with the finite-width line. The line can be treated as a rectangle, and the section of the line area between two adjacent vertical (or two adjacent horizontal) screen grid lines is then a polygon. Overlap areas for pixels are calculated by determining how much of the polygon overlaps each pixel in that column (or row). In Figure 60, the pixel with screen grid coordinates $(10, 20)$ is about 90 percent covered by the line area, so its intensity would be set to 90 percent of the maximum intensity. Similarly, the pixel at $(10, 21)$ would be set to an intensity of about 15 percent of maximum. A method for estimating pixel overlap areas is illustrated by the supersampling example in Figure 60. The total number of subpixels within the line boundaries is approximately equal to the overlap area, and this estimation is improved by using finer subpixel grids.

Filtering Techniques

A more accurate method for antialiasing lines is to use filtering techniques. The method is similar to applying a weighted pixel mask, but now we imagine a continuous weighting surface (or filter function) covering the pixel. Figure 62 shows examples of rectangular, conical, and Gaussian filter functions. Methods for applying the filter function are similar to those for applying a weighting mask, but now we integrate over the pixel surface to obtain the weighted average intensity. To reduce computation, table lookups are commonly used to evaluate the integrals.

Pixel Phasing

On raster systems that can address subpixel positions within the screen grid, pixel phasing can be used to antialias objects. A line display is smoothed with this technique by moving (micropositioning) pixel positions closer to the line path. Systems incorporating pixel phasing are designed so that the electron beam can be shifted by a fraction of a pixel diameter. The electron beam is typically shifted by $\frac{1}{4}$, $\frac{1}{2}$, or $\frac{3}{4}$ of a pixel diameter to plot points closer to the true path of a line or object edge. Some systems also allow the size of individual pixels to be adjusted as an additional means for distributing intensities. Figure 63 illustrates the antialiasing effects of pixel phasing on a variety of line paths.
Compensating for Line-Intensity Differences

Antialiasing a line to soften the stair-step effect also compensates for another raster effect, illustrated in Figure 64. Both lines are plotted with the same number of pixels, yet the diagonal line is longer than the horizontal line by a factor of $\sqrt{2}$. For example, if the horizontal line had a length of 10 centimeters, the diagonal line would have a length of more than 14 centimeters. The visual effect of this is that the diagonal line appears less bright than the horizontal line, because the diagonal line is displayed with a lower intensity per unit length. A line-drawing algorithm could be adapted to compensate for this effect by adjusting the intensity of each
Unequal length lines displayed with the same number of pixels in each line.

According to its slope. Horizontal and vertical lines would be displayed with the lowest intensity, while 45° lines would be given the highest intensity. But if antialiasing techniques are applied to a display, intensities are compensated automatically. When the finite width of a line is taken into account, pixel intensities are adjusted so that the line displays a total intensity proportional to its length.

Antialiasing Area Boundaries

The antialiasing concepts that we have discussed for lines can also be applied to the boundaries of areas to remove their jagged appearance. We can incorporate these procedures into a scan-line algorithm to smooth out the boundaries as the area is generated.

If system capabilities permit the repositioning of pixels, we could smooth area boundaries by shifting pixel positions closer to the boundary. Other methods adjust pixel intensity at a boundary position according to the percent of the pixel area that is interior to the object. In Figure 65, the pixel at position \((x, y)\) has about half its area inside the polygon boundary. Therefore, the intensity at that position would be adjusted to one-half its assigned value. At the next position \((x + 1, y + 1)\) along the boundary, the intensity is adjusted to about one-third the assigned value for that point. Similar adjustments, based on the percent of pixel area coverage, are applied to the other intensity values around the boundary.

Supersampling methods can be applied by determining the number of subpixels that are in the interior of an object. A partitioning scheme with four subareas per pixel is shown in Figure 66. The original \(4 \times 4\) grid of pixels is turned into an \(8 \times 8\) grid, and we now process eight scan lines across this grid instead of four. Figure 67 shows one of the pixel areas in this grid that overlaps an object.

\(I_{\text{pixel}}\)
boundary. Along the two scan lines, we determine that three of the subpixel areas are inside the boundary. So we set the pixel intensity at 75 percent of its maximum value.

Another method for determining the percentage of pixel area within a fill region, developed by Pitteway and Watkinson, is based on the midpoint line algorithm. This algorithm selects the next pixel along a line by testing the location of the midpoint position between two pixels. As in the Bresenham algorithm, we set up a decision parameter $p$ whose sign tells us which of the next two candidate pixels is closer to the line. By slightly modifying the form of $p$, we obtain a quantity that also gives the percentage of the current pixel area that is covered by an object.

We first consider the method for a line with slope $m$ in the range from 0 to 1. In Figure 68, a straight-line path is shown on a pixel grid. Assuming that the pixel at position $(x_k, y_k)$ has been plotted, the next pixel nearest the line at $x = x_k + 1$ is either the pixel at $y_k$ or the one at $y_k + 1$. We can determine which pixel is nearer with the calculation

$$y - y_{\text{mid}} = [m(x_k + 1) + b] - (y_k + 0.5)$$

This gives the vertical distance from the actual $y$ coordinate on the line to the halfway point between pixels at position $y_k$ and $y_k + 1$. If this difference calculation is negative, the pixel at $y_k$ is closer to the line. If the difference is positive, the pixel at $y_k + 1$ is closer. We can adjust this calculation so that it produces a positive number in the range from 0 to 1 by adding the quantity $1 - m$:

$$p = [m(x_k + 1) + b] - (y_k + 0.5) + (1 - m)$$

Now the pixel at $y_k$ is nearer if $p < 1 - m$, and the pixel at $y_k + 1$ is nearer if $p > 1 - m$.

Parameter $p$ also measures the amount of the current pixel that is overlapped by the area. For the pixel at $(x_k, y_k)$ in Figure 69, the interior part of the pixel is trapezoidal and has an area that can be calculated as

$$\text{area} = m \cdot x_k + b - y_k + 0.5$$

This expression for the overlap area of the pixel at $(x_k, y_k)$ is the same as that for parameter $p$ in Equation 71. Therefore, by evaluating $p$ to determine the next pixel position along the polygon boundary, we also determine the percentage of area coverage for the current pixel.

We can generalize this algorithm to accommodate lines with negative slopes and lines with slopes greater than 1. This calculation for parameter $p$ could then

FIGURE 68
Boundary edge of a fill area passing through a pixel grid section.

FIGURE 69
Overlap area of a pixel rectangle, centered at position $(x_k, y_k)$, with the interior of a polygon fill area.
be incorporated into a midpoint line algorithm to locate pixel positions along a polygon edge and, concurrently, adjust pixel intensities along the boundary lines. Also, we can adjust the calculations to reference pixel coordinates at their lower-left coordinates and maintain area proportions, as discussed in Section 8.

At polygon vertices and for very skinny polygons, as shown in Figure 70, we have more than one boundary edge passing through a pixel area. For these cases, we need to modify the Pitteway-Watkinson algorithm by processing all edges passing through a pixel and determining the correct interior area.

Filtering techniques discussed for line antialiasing can also be applied to area edges. In addition, the various antialiasing methods can be applied to polygon areas or to regions with curved boundaries. Equations describing the boundaries are used to estimate the amount of pixel overlap with the area to be displayed, and coherence techniques are used along and between scan lines to simplify the calculations.

16 Summary

Three methods that can be used to locate pixel positions along a straight-line path are the DDA algorithm, Bresenham’s algorithm, and the midpoint method. Bresenham’s line algorithm and the midpoint line method are equivalent, and they are the most efficient. Color values for the pixel positions along the line path are efficiently stored in the frame buffer by incrementally calculating the memory addresses. Any of the line-generating algorithms can be adapted to a parallel implementation by partitioning the line segments and distributing the partitions among the available processors.

Circles and ellipses can be efficiently and accurately scan-converted using midpoint methods and taking curve symmetry into account. Other conic sections (parabolas and hyperbolas) can be plotted with similar methods. Spline curves, which are piecewise continuous polynomials, are widely used in animation and in CAD. Parallel implementations for generating curve displays can be accomplished with methods similar to those for parallel line processing.

To account for the fact that displayed lines and curves have finite widths, we can adjust the pixel dimensions of objects to coincide to the specified geometric dimensions. This can be done with an addressing scheme that references pixel positions at their lower-left corner, or by adjusting line lengths.

Scan-line methods are commonly used to fill polygons, circles, and ellipses. Across each scan line, the interior fill is applied to pixel positions between each pair of boundary intersections, left to right. For polygons, scan-line intersections with vertices can result in an odd number of intersections. This can be resolved by shortening some polygon edges. Scan-line fill algorithms can be simplified if fill areas are restricted to convex polygons. A further simplification is achieved if all fill areas in a scene are triangles. The interior pixels along each scan line are assigned appropriate color values, depending on the fill-attribute specifications. Painting programs generally display fill regions using a boundary-fill method or a flood-fill method. Each of these two fill methods requires an initial interior point. The interior is then painted pixel by pixel from the initial point out to the region boundaries.

Soft-fill procedures provide a new fill color for a region that has the same variations as the previous fill color. One example of this approach is the linear soft-fill algorithm that assumes that the previous fill was a linear combination of foreground and background colors. This same linear relationship is then
determined from the frame buffer settings and used to repaint the area in a new color.

We can improve the appearance of raster primitives by applying antialiasing procedures that adjust pixel intensities. One method for doing this is to supersample. That is, we consider each pixel to be composed of subpixels and we calculate the intensity of the subpixels and average the values of all subpixels. We can also weight the subpixel contributions according to position, giving higher weights to the central subpixels. Alternatively, we can perform area sampling and determine the percentage of area coverage for a screen pixel, then set the pixel intensity proportional to this percentage. Another method for antialiasing is to build special hardware configurations that can shift pixel positions.

REFERENCES


EXERCISES
1. Implement a polyline function using the DDA algorithm, given any number (n) of input points. A single point is to be plotted when n = 1.
2. Extend Bresenham’s line algorithm to generate lines with any slope, taking symmetry between quadrants into account.
3. Implement a polyline function, using the algorithm from the previous exercise, to display the set of straight lines connecting a list of n input points. For n = 1, the routine displays a single point.
4. Use the midpoint method to derive decision parameters for generating points along a straight-line path with slope in the range 0 < m < 1. Show that the midpoint decision parameters are the same as those in the Bresenham line algorithm.
5. Use the midpoint method to derive decision parameters that can be used to generate straight-line segments with any slope.
6. Set up a parallel version of Bresenham’s line algorithm for slopes in the range 0 < m < 1.
7. Set up a parallel version of Bresenham’s algorithm for straight lines with any slope.
8. Suppose you have a system with an 8 inch by 10 inch video monitor that can display 100 pixels per inch. If memory is organized in one byte words, the starting frame buffer address is 0, and each pixel is assigned one byte of storage, what is the frame buffer address of the pixel with screen coordinates (x, y)?
9. Suppose you have a system with a 12 inch by 14 inch video monitor that can display 120 pixels per inch. If memory is organized in one byte words, the starting frame buffer address is 0, and each pixel is assigned one byte of storage, what is the frame buffer address of the pixel with screen coordinates (x, y)?
10. Suppose you have a system with a 12 inch by 14 inch video monitor that can display 120 pixels per inch. If memory is organized in one byte words, the starting frame buffer address is 0, and each pixel is assigned 4 bits of storage, what is the frame buffer address of the pixel with screen coordinates (x, y)?
11. Incorporate the iterative techniques for calculating frame-buffer addresses (Section 3) into the Bresenham line algorithm.
12. Revise the midpoint circle algorithm to display circles with input geometric magnitudes preserved (Section 8).
13. Set up a procedure for a parallel implementation of the midpoint circle algorithm.
14. Derive decision parameters for the midpoint ellipse algorithm assuming the start position is (r_x, 0) and points are to be generated along the curve path in counterclockwise order.
15 Set up a procedure for a parallel implementation of the midpoint ellipse algorithm.

16 Devise an efficient algorithm that takes advantage of symmetry properties to display a sine function over one cycle.

17 Modify the algorithm in the preceding exercise to display a sine curve over any specified angular interval.

18 Devise an efficient algorithm, taking function symmetry into account, to display a plot of damped harmonic motion:

\[ y = A e^{-kx} \sin(\omega x + \theta) \]

where \( \omega \) is the angular frequency and \( \theta \) is the phase of the sine function. Plot \( y \) as a function of \( x \) for several cycles of the sine function or until the maximum amplitude is reduced to \( \frac{A}{10} \).

19 Use the algorithm developed in the previous exercise to write a program that displays one cycle of a sine curve. The curve should begin at the left edge of the display window and complete at the right edge, and the amplitude should be scaled so that the maximum and minimum values of the curve are equal to the maximum and minimum \( y \) values of the display window.

20 Using the midpoint method, and taking symmetry into account, develop an efficient algorithm for scan conversion of the following curve over the interval \(-10 \leq x \leq 10\):

\[ y = \frac{1}{12}x^3 \]

21 Use the algorithm developed in the previous exercise to write a program that displays a portion of a sine curve determined by an input angular interval. The curve should begin at the left edge of the display window and complete at the right edge, and the amplitude should be scaled so that the maximum and minimum values of the curve are equal to the maximum and minimum \( y \) values of the display window.

22 Use the midpoint method and symmetry considerations to scan convert the parabola

\[ x = y^2 - 5 \]

over the interval \(-10 \leq x \leq 10\).

23 Use the midpoint method and symmetry considerations to scan convert the parabola

\[ y = 50 - x^2 \]

over the interval \(-5 \leq x \leq 5\).

24 Set up a midpoint algorithm, taking symmetry considerations into account to scan convert any parabola of the form

\[ y = ax^2 + b \]

with input values for parameters \( a \), \( b \), and the range for \( x \).

25 Define an efficient polygon-mesh representation for a cylinder and justify your choice of representation.

26 Implement a general line-style function by modifying Bresenham’s line-drawing algorithm to display solid, dashed, or dotted lines.

27 Implement a line-style function using a midpoint line algorithm to display solid, dashed, or dotted lines.

28 Devise a parallel method for implementing a line-style function.

29 Devise a parallel method for implementing a line-width function.

30 A line specified by two endpoints and a width can be converted to a rectangular polygon with four vertices and then displayed using a scan-line method. Develop an efficient algorithm for computing the four vertices needed to define such a rectangle, with the line endpoints and line width as input parameters.

31 Implement a line-width function in a line-drawing program so that any one of three line widths can be displayed.

32 Write a program to output a line graph of three data sets defined over the same \( x \)-coordinate range. Input to the program is to include the three sets of data values and the labels for the graph. The data sets are to be scaled to fit within a defined coordinate range for a display window. Each data set is to be plotted with a different line style.

33 Modify the program in the previous exercise to plot the three data sets in different colors, as well as different line styles.

34 Set up an algorithm for displaying thick lines with butt caps, round caps, or projecting square caps. These options can be provided in an option menu.

35 Devise an algorithm for displaying thick polylines with a miter join, a round join, or a bevel join. These options can be provided in an option menu.

36 Implement pen and brush menu options for a line-drawing procedure, including at least two options: round and square shapes.

37 Modify a line-drawing algorithm so that the intensity of the output line is set according to its slope. That is, by adjusting pixel intensities according to the value of the slope, all lines are displayed with the same intensity per unit length.
38 Define and implement a function for controlling the line style (solid, dashed, dotted) of displayed ellipses.
39 Define and implement a function for setting the width of displayed ellipses.
40 Modify the scan-line algorithm to apply any specified rectangular fill pattern to a polygon interior, starting from a designated pattern position.
41 Write a program to scan convert the interior of a specified ellipse into a solid color.
42 Write a procedure to fill the interior of a given ellipse with a specified pattern.
43 Write a procedure for filling the interior of any specified set of fill-area vertices, including one with crossing edges, using the nonzero winding number rule to identify interior regions.
44 Modify the boundary-fill algorithm for a 4-connected region to avoid excessive stacking by incorporating scan-line methods.
45 Write a boundary-fill procedure to fill an 8-connected region.
46 Explain how an ellipse displayed with the midpoint method could be properly filled with a boundary-fill algorithm.
47 Develop and implement a flood-fill algorithm to fill the interior of any specified area.
48 Define and implement a procedure for changing the size of an existing rectangular fill pattern.
49 Write a procedure to implement a soft-fill algorithm. Carefully define what the soft-fill algorithm is to accomplish and how colors are to be combined.
50 Devise an algorithm for adjusting the height and width of characters defined as rectangular grid patterns.
51 Implement routines for setting the character up vector and the text path for controlling the display of character strings.
52 Write a program to align text as specified by input values for the alignment parameters.
53 Develop procedures for implementing marker attributes (size and color).
54 Implement an antialiasing procedure by extending Bresenham’s line algorithm to adjust pixel intensities in the vicinity of a line path.
55 Implement an antialiasing procedure for the midpoint line algorithm.
56 Develop an algorithm for antialiasing elliptical boundaries.
57 Modify the scan-line algorithm for area fill to incorporate antialiasing. Use coherence techniques to reduce calculations on successive scan lines.
58 Write a program to implement the Pitteway-Watkinson antialiasing algorithm as a scan-line procedure to fill a polygon interior, using the OpenGL point-plotting function.

IN MORE DEPTH

1 Write routines to implement Bresenham’s line-drawing algorithm and the DDA line-drawing algorithm and use them to draw the outlines of the shapes in the current snapshot of your application. Record the runtimes of the algorithms and compare the performance of the two. Next examine the polygons that represent the objects in your scene and either choose a few that would be better represented using ellipses or other curves or add a few objects with this property. Implement a midpoint algorithm to draw the ellipses or curves that represent these objects and use it to draw the outlines of those objects. Discuss ways in which you could improve the performance of the algorithms you developed if you had direct access to parallel hardware.

2 Implement the general scan-line polygon-fill algorithm to fill in the polygons that make up the objects in your scene with solid colors. Next, implement a scan-line curve-filling algorithm to fill the curved objects you added in the previous exercise. Finally, implement a boundary fill algorithm to fill all of the objects in your scene. Compare the run times of the two approaches to filling in the shapes in your scene.
So far, we have seen how we can describe a scene in terms of graphics primitives, such as line segments and fill areas, and the attributes associated with these primitives. Also, we have explored the scan-line algorithms for displaying output primitives on a raster device. Now, we take a look at transformation operations that we can apply to objects to reposition or resize them. These operations are also used in the viewing routines that convert a world-coordinate scene description to a display for an output device. In addition, they are used in a variety of other applications, such as computer-aided design (CAD) and computer animation. An architect, for example, creates a layout by arranging the orientation and size of the component parts of a design, and a computer animator develops a video sequence by moving the “camera” position or the objects in a scene along specified paths. Operations that are applied to the geometric description of an object to change its position, orientation, or size are called **geometric transformations**.

Sometimes geometric transformations are also referred to as **modeling transformations**, but some graphics packages make a
Two-Dimensional Geometric Transformations

Distinction between the two. In general, modeling transformations are used to construct a scene or to give the hierarchical description of a complex object that is composed of several parts, which in turn could be composed of simpler parts, and so forth. For example, an aircraft consists of wings, tail, fuselage, engine, and other components, each of which can be specified in terms of second-level components, and so on, down the hierarchy of component parts. Thus, the aircraft can be described in terms of these components and an associated “modeling” transformation for each one that describes how that component is to be fitted into the overall aircraft design. Geometric transformations, on the other hand, can be used to describe how objects might move around in a scene during an animation sequence or simply to view them from another angle. Therefore, some graphics packages provide two sets of transformation routines, while other packages have a single set of functions that can be used for both geometric transformations and modeling transformations.

1 Basic Two-Dimensional Geometric Transformations

The geometric-transformation functions that are available in all graphics packages are those for translation, rotation, and scaling. Other useful transformation routines that are sometimes included in a package are reflection and shearing operations. To introduce the general concepts associated with geometric transformations, we first consider operations in two dimensions.

Two-Dimensional Translation

We perform a **translation** on a single coordinate point by adding offsets to its coordinates so as to generate a new coordinate position. In effect, we are moving the original point position along a straight-line path to its new location. Similarly, a translation is applied to an object that is defined with multiple coordinate positions, such as a quadrilateral, by relocating all the coordinate positions by the same displacement along parallel paths. Then the complete object is displayed at the new location.

To translate a two-dimensional position, we add translation distances \( t_x \) and \( t_y \) to the original coordinates \((x, y)\) to obtain the new coordinate position \((x', y')\) as shown in Figure 1.

\[
  x' = x + t_x, \quad y' = y + t_y
\]  

The translation distance pair \((t_x, t_y)\) is called a **translation vector** or **shift vector**.

We can express Equations 1 as a single matrix equation by using the following column vectors to represent coordinate positions and the translation vector:

\[
  P = \begin{bmatrix} x \\ y \end{bmatrix}, \quad P' = \begin{bmatrix} x' \\ y' \end{bmatrix}, \quad T = \begin{bmatrix} t_x \\ t_y \end{bmatrix}
\]  

This allows us to write the two-dimensional translation equations in the matrix form

\[
P' = P + T
\]
Translation is a rigid-body transformation that moves objects without deformation. That is, every point on the object is translated by the same amount. A straight-line segment is translated by applying Equation 3 to each of the two line endpoints and redrawing the line between the new endpoint positions. A polygon is translated similarly. We add a translation vector to the coordinate position of each vertex and then regenerate the polygon using the new set of vertex coordinates. Figure 2 illustrates the application of a specified translation vector to move an object from one position to another.

The following routine illustrates the translation operations. An input translation vector is used to move the \( n \) vertices of a polygon from one world-coordinate position to another, and OpenGL routines are used to regenerate the translated polygon.

```cpp
class wcPt2D {
    public:
        GLfloat x, y;
};

void translatePolygon (wcPt2D * verts, GLint nVerts, GLfloat tx, GLfloat ty) {
    GLint k;

    for (k = 0; k < nVerts; k++) {
        verts[k].x = verts[k].x + tx;
        verts[k].y = verts[k].y + ty;
    }

    glBegin (GL_POLYGON);
    for (k = 0; k < nVerts; k++)
        glVertex2f (verts[k].x, verts[k].y);
    glEnd ( );
}
```

If we want to delete the original polygon, we could display it in the background color before translating it. Other methods for deleting picture components...
are available in some graphics packages. Also, if we want to save the original polygon position, we can store the translated positions in a different array.

Similar methods are used to translate other objects. To change the position of a circle or ellipse, we translate the center coordinates and redraw the figure in the new location. For a spline curve, we translate the points that define the curve path and then reconstruct the curve sections between the new coordinate positions.

**Two-Dimensional Rotation**

We generate a rotation transformation of an object by specifying a rotation axis and a rotation angle. All points of the object are then transformed to new positions by rotating the points through the specified angle about the rotation axis.

A two-dimensional rotation of an object is obtained by repositioning the object along a circular path in the $xy$ plane. In this case, we are rotating the object about a rotation axis that is perpendicular to the $xy$ plane (parallel to the coordinate $z$ axis). Parameters for the two-dimensional rotation are the rotation angle $\theta$ and a position $(x_r, y_r)$, called the rotation point (or pivot point), about which the object is to be rotated (Figure 3). The pivot point is the intersection position of the rotation axis with the $xy$ plane. A positive value for the angle $\theta$ defines a counterclockwise rotation about the pivot point, as in Figure 3, and a negative value rotates objects in the clockwise direction.

To simplify the explanation of the basic method, we first determine the transformation equations for rotation of a point position $\mathbf{P}$ when the pivot point is at the coordinate origin. The angular and coordinate relationships of the original and transformed point positions are shown in Figure 4. In this figure, $r$ is the constant distance of the point from the origin, angle $\phi$ is the original angular position of the point from the horizontal, and $\theta$ is the rotation angle. Using standard trigonometric identities, we can express the transformed coordinates in terms of angles $\theta$ and $\phi$ as

$$x' = r \cos(\phi + \theta) = r \cos \phi \cos \theta - r \sin \phi \sin \theta$$
$$y' = r \sin(\phi + \theta) = r \cos \phi \sin \theta + r \sin \phi \cos \theta$$

(4)

The original coordinates of the point in polar coordinates are

$$x = r \cos \phi, \quad y = r \sin \phi$$

(5)

Substituting expressions 5 into 4, we obtain the transformation equations for rotating a point at position $(x, y)$ through an angle $\theta$ about the origin:

$$x' = x \cos \theta - y \sin \theta$$
$$y' = x \sin \theta + y \cos \theta$$

(6)

With the column-vector representations 2 for coordinate positions, we can write the rotation equations in the matrix form

$$\mathbf{P}' = \mathbf{R} \cdot \mathbf{P}$$

(7)

where the rotation matrix is

$$\mathbf{R} = \begin{bmatrix} \cos \theta & -\sin \theta \\ \sin \theta & \cos \theta \end{bmatrix}$$

(8)

A column-vector representation for a coordinate position $\mathbf{P}$, as in Equations 2, is standard mathematical notation. However, early graphics systems sometimes used a row-vector representation for point positions. This changes the order in which the matrix multiplication for a rotation would be performed. But now, graphics packages such as OpenGL, Java, PHIGS, and GKS all follow the standard column-vector convention.
Rotation of a point about an arbitrary pivot position is illustrated in Figure 5. Using the trigonometric relationships indicated by the two right triangles in this figure, we can generalize Equations 6 to obtain the transformation equations for rotation of a point about any specified rotation position \((x_r, y_r)\):

\[
\begin{align*}
    x' &= x_r + (x - x_r) \cos \theta - (y - y_r) \sin \theta \\
    y' &= y_r + (x - x_r) \sin \theta + (y - y_r) \cos \theta
\end{align*}
\]  

(9)

These general rotation equations differ from Equations 6 by the inclusion of additive terms, as well as the multiplicative factors on the coordinate values. The matrix expression 7 could be modified to include pivot coordinates by including the matrix addition of a column vector whose elements contain the additive (translational) terms in Equations 9. There are better ways, however, to formulate such matrix equations, and in Section 2, we discuss a more consistent scheme for representing the transformation equations.

As with translations, rotations are rigid-body transformations that move objects without deformation. Every point on an object is rotated through the same angle. A straight-line segment is rotated by applying Equations 9 to each of the two line endpoints and redrawing the line between the new endpoint positions. A polygon is rotated by displacing each vertex using the specified rotation angle and then regenerating the polygon using the new vertices. We rotate a curve by repositioning the defining points for the curve and then redrawing it. A circle or an ellipse, for instance, can be rotated about a noncentral pivot point by moving the center position through the arc that subtends the specified rotation angle. In addition, we could rotate an ellipse about its center coordinates simply by rotating the major and minor axes.

In the following code example, a polygon is rotated about a specified world-coordinate pivot point. Parameters input to the rotation procedure are the original vertices of the polygon, the pivot-point coordinates, and the rotation angle \(\theta\) specified in radians. Following the transformation of the vertex positions, the polygon is regenerated using OpenGL routines.

```cpp
class wcPt2D {
public:
    GLfloat x, y;
};

void rotatePolygon (wcPt2D * verts, GLint nVerts, wcPt2D pivPt, GLdouble theta)
{
    wcPt2D * vertsRot;
    GLint k;

    for (k = 0; k < nVerts; k++) {
        vertsRot[k].x = pivPt.x + (verts[k].x - pivPt.x) * cos (theta)
            - (verts[k].y - pivPt.y) * sin (theta);
        vertsRot[k].y = pivPt.y + (verts[k].x - pivPt.x) * sin (theta)
            + (verts[k].y - pivPt.y) * cos (theta);
    }

    glBegin (GL_POLYGON);
    for (k = 0; k < nVerts; k++)
        glVertex2f (vertsRot[k].x, vertsRot[k].y);
    glEnd ( );
}
```
Two-Dimensional Geometric Transformations

Two-Dimensional Scaling

To alter the size of an object, we apply a scaling transformation. A simple two-dimensional scaling operation is performed by multiplying object positions \((x, y)\) by scaling factors \(s_x\) and \(s_y\) to produce the transformed coordinates \((x', y')\):

\[
x' = x \cdot s_x, \quad y' = y \cdot s_y
\]  

(10)

Scaling factor \(s_x\) scales an object in the \(x\) direction, while \(s_y\) scales in the \(y\) direction. The basic two-dimensional scaling equations 10 can also be written in the following matrix form:

\[
\begin{bmatrix}
x' \\
y'
\end{bmatrix} =
\begin{bmatrix}
s_x & 0 \\
0 & s_y
\end{bmatrix}
\begin{bmatrix}
x \\
y
\end{bmatrix}
\]

(11)

or

\[
P' = S \cdot P
\]

(12)

where \(S\) is the \(2 \times 2\) scaling matrix in Equation 11.

Any positive values can be assigned to the scaling factors \(s_x\) and \(s_y\). Values less than 1 reduce the size of objects; values greater than 1 produce enlargements. Specifying a value of 1 for both \(s_x\) and \(s_y\) leaves the size of objects unchanged. When \(s_x\) and \(s_y\) are assigned the same value, a uniform scaling is produced, which maintains relative object proportions. Unequal values for \(s_x\) and \(s_y\) result in a differential scaling that is often used in design applications, where pictures are constructed from a few basic shapes that can be adjusted by scaling and positioning transformations (Figure 6). In some systems, negative values can also be specified for the scaling parameters. This not only resizes an object, it reflects it about one or more of the coordinate axes.

Objects transformed with Equation 11 are both scaled and repositioned. Scaling factors with absolute values less than 1 move objects closer to the coordinate origin, while absolute values greater than 1 move coordinate positions farther from the origin. Figure 7 illustrates scaling of a line by assigning the value 0.5 to both \(s_x\) and \(s_y\) in Equation 11. Both the line length and the distance from the origin are reduced by a factor of \(\frac{1}{2}\).

We can control the location of a scaled object by choosing a position, called the fixed point, that is to remain unchanged after the scaling transformation. Coordinates for the fixed point, \((x_f, y_f)\), are often chosen at some object position, such as its centroid (see Appendix A), but any other spatial position can be selected. Objects are now resized by scaling the distances between object points and the fixed point (Figure 8). For a coordinate position \((x, y)\), the scaled coordinates \((x', y')\) are then calculated from the following relationships:

\[
x' - x_f = (x - x_f)s_x, \quad y' - y_f = (y - y_f)s_y
\]  

(13)

We can rewrite Equations 13 to separate the multiplicative and additive terms as

\[
x' = x \cdot s_x + x_f(1 - s_x)
\]

\[
y' = y \cdot s_y + y_f(1 - s_y)
\]  

(14)

where the additive terms \(x_f(1 - s_x)\) and \(y_f(1 - s_y)\) are constants for all points in the object.

Including coordinates for a fixed point in the scaling equations is similar to including coordinates for a pivot point in the rotation equations. We can set up...
a column vector whose elements are the constant terms in Equations 14, then
add this column vector to the product \( S \cdot P \) in Equation 12. In the next section,
we discuss a matrix formulation for the transformation equations that involves
only matrix multiplication.

Polygons are scaled by applying transformations 14 to each vertex, then
regenerating the polygon using the transformed vertices. For other objects,
we apply the scaling transformation equations to the parameters defining the
objects. To change the size of a circle, we can scale its radius and calculate the
new coordinate positions around the circumference. And to change the size of an ellipse, we apply scaling parameters to its two axes and then plot the new ellipse
positions about its center coordinates.

The following procedure illustrates an application of the scaling calculations
for a polygon. Coordinates for the polygon vertices and for the fixed point are
input parameters, along with the scaling factors. After the coordinate transforma-
tions, OpenGL routines are used to generate the scaled polygon.

```cpp
class wcPt2D {
    public:
        GLfloat x, y;
};

void scalePolygon (wcPt2D * verts, GLint nVerts, wcPt2D fixedPt,
                    GLfloat sx, GLfloat sy)
{
    wcPt2D vertsNew;
    GLint k;

    for (k = 0; k < nVerts; k++) {
        vertsNew [k].x = verts [k].x * sx + fixedPt.x * (1 - sx);
        vertsNew [k].y = verts [k].y * sy + fixedPt.y * (1 - sy);
    }
    glBegin (GL_POLYGON);
    for (k = 0; k < nVerts; k++)
        glVertex2f (vertsNew [k].x, vertsNew [k].y);
    glEnd ( );
}
```
Two-Dimensional Geometric Transformations

We have seen in Section 1 that each of the three basic two-dimensional transformations (translation, rotation, and scaling) can be expressed in the general matrix form

\[ \mathbf{P}' = \mathbf{M}_1 \cdot \mathbf{P} + \mathbf{M}_2 \]  \hspace{1cm} (15)

with coordinate positions \( \mathbf{P} \) and \( \mathbf{P}' \) represented as column vectors. Matrix \( \mathbf{M}_1 \) is a 2 \( \times \) 2 array containing multiplicative factors, and \( \mathbf{M}_2 \) is a two-element column matrix containing translational terms. For translation, \( \mathbf{M}_1 \) is the identity matrix. For rotation or scaling, \( \mathbf{M}_2 \) contains the translational terms associated with the pivot point or scaling fixed point. To produce a sequence of transformations with these equations, such as scaling followed by rotation and then translation, we could calculate the transformed coordinates one step at a time. First, coordinate positions are scaled, then these scaled coordinates are rotated, and finally, the rotated coordinates are translated. A more efficient approach, however, is to combine the transformations so that the final coordinate positions are obtained directly from the initial coordinates, without calculating intermediate coordinate values. We can do this by reformulating Equation 15 to eliminate the matrix addition operation.

Homogeneous Coordinates

Multiplicative and translational terms for a two-dimensional geometric transformation can be combined into a single matrix if we expand the representations to 3 \( \times \) 3 matrices. Then we can use the third column of a transformation matrix for the translation terms, and all transformation equations can be expressed as matrix multiplications. But to do so, we also need to expand the matrix representation for a two-dimensional coordinate position to a three-element column matrix. A standard technique for accomplishing this is to expand each two-dimensional coordinate-position representation (\( x, y \)) to a three-element representation (\( x_h, y_h, h \)), called homogeneous coordinates, where the homogeneous parameter \( h \) is a nonzero value such that

\[ x = \frac{x_h}{h}, \quad y = \frac{y_h}{h} \]  \hspace{1cm} (16)

Therefore, a general two-dimensional homogeneous coordinate representation could also be written as \((h \cdot x, h \cdot y, h)\). For geometric transformations, we can choose the homogeneous parameter \( h \) to be any nonzero value. Thus, each coordinate point \((x, y)\) has an infinite number of equivalent homogeneous representations. A convenient choice is simply to set \( h = 1 \). Each two-dimensional position is then represented with homogeneous coordinates \((x, y, 1)\). Other values for parameter \( h \) are needed, for example, in matrix formulations of three-dimensional viewing transformations.

The term homogeneous coordinates is used in mathematics to refer to the effect of this representation on Cartesian equations. When a Cartesian point \((x, y)\) is converted to a homogeneous representation \((x_h, y_h, h)\), equations containing \( x \) and \( y \), such as \( f(x, y) = 0 \), become homogeneous equations in the three parameters \( x_h, y_h, \) and \( h \). This just means that if each of the three parameters is replaced by any value \( v \) times that parameter, the value \( v \) can be factored out of the equations.

Expressing positions in homogeneous coordinates allows us to represent all geometric transformation equations as matrix multiplications, which is the standard method used in graphics systems. Two-dimensional coordinate positions are represented with three-element column vectors, and two-dimensional transformation operations are expressed as 3 \( \times \) 3 matrices.
Two-Dimensional Translation Matrix

Using a homogeneous-coordinate approach, we can represent the equations for a two-dimensional translation of a coordinate position using the following matrix multiplication:

\[
\begin{bmatrix}
    x' \\
    y' \\
    1
\end{bmatrix} =
\begin{bmatrix}
    1 & 0 & t_x \\
    0 & 1 & t_y \\
    0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
    x \\
    y \\
    1
\end{bmatrix}
\]  

(17)

This translation operation can be written in the abbreviated form

\[ P' = T(t_x, t_y) \cdot P \]  

(18)

with \( T(t_x, t_y) \) as the \( 3 \times 3 \) translation matrix in Equation 17. In situations where there is no ambiguity about the translation parameters, we can simply represent the translation matrix as \( T \).

Two-Dimensional Rotation Matrix

Similarly, two-dimensional rotation transformation equations about the coordinate origin can be expressed in the matrix form

\[
\begin{bmatrix}
    x' \\
    y' \\
    1
\end{bmatrix} =
\begin{bmatrix}
    \cos \theta & -\sin \theta & 0 \\
    \sin \theta & \cos \theta & 0 \\
    0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
    x \\
    y \\
    1
\end{bmatrix}
\]  

(19)

or as

\[ P' = R(\theta) \cdot P \]  

(20)

The rotation transformation operator \( R(\theta) \) is the \( 3 \times 3 \) matrix in Equation 19 with rotation parameter \( \theta \). We can also write this rotation matrix simply as \( R \).

In some graphics libraries, a two-dimensional rotation function generates only rotations about the coordinate origin, as in Equation 19. A rotation about any other pivot point must then be performed as a sequence of transformation operations. An alternative approach in a graphics package is to provide additional parameters in the rotation routine for the pivot-point coordinates. A rotation routine that includes a pivot-point parameter then sets up a general rotation matrix without the need to invoke a succession of transformation functions.

Two-Dimensional Scaling Matrix

Finally, a scaling transformation relative to the coordinate origin can now be expressed as the matrix multiplication

\[
\begin{bmatrix}
    x' \\
    y' \\
    1
\end{bmatrix} =
\begin{bmatrix}
    s_x & 0 & 0 \\
    0 & s_y & 0 \\
    0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
    x \\
    y \\
    1
\end{bmatrix}
\]  

(21)

or

\[ P' = S(s_x, s_y) \cdot P \]  

(22)

The scaling operator \( S(s_x, s_y) \) is the \( 3 \times 3 \) matrix in Equation 21 with parameters \( s_x \) and \( s_y \). And, in most cases, we can represent the scaling matrix simply as \( S \).

Some libraries provide a scaling function that can generate only scaling with respect to the coordinate origin, as in Equation 21. In this case, a scaling transformation relative to another reference position is handled as a succession of transformation operations. However, other systems do include a general scaling routine that can construct the homogeneous matrix for scaling with respect to a designated fixed point.
3 Inverse Transformations

For translation, we obtain the inverse matrix by negating the translation distances. Thus, if we have two-dimensional translation distances \( t_x \) and \( t_y \), the inverse translation matrix is

\[
T^{-1} = \begin{bmatrix}
1 & 0 & -t_x \\
0 & 1 & -t_y \\
0 & 0 & 1
\end{bmatrix}
\]  

(23)

This produces a translation in the opposite direction, and the product of a translation matrix and its inverse produces the identity matrix.

An inverse rotation is accomplished by replacing the rotation angle by its negative. For example, a two-dimensional rotation through an angle \( \theta \) about the coordinate origin has the inverse transformation matrix

\[
R^{-1} = \begin{bmatrix}
\cos \theta & \sin \theta & 0 \\
-\sin \theta & \cos \theta & 0 \\
0 & 0 & 1
\end{bmatrix}
\]

(24)

Negative values for rotation angles generate rotations in a clockwise direction, so the identity matrix is produced when any rotation matrix is multiplied by its inverse. Because only the sine function is affected by the change in sign of the rotation angle, the inverse matrix can also be obtained by interchanging rows and columns. That is, we can calculate the inverse of any rotation matrix \( R \) by evaluating its transpose \( (R^{-1} = R^T) \).

We form the inverse matrix for any scaling transformation by replacing the scaling parameters with their reciprocals. For two-dimensional scaling with parameters \( s_x \) and \( s_y \) applied relative to the coordinate origin, the inverse transformation matrix is

\[
S^{-1} = \begin{bmatrix}
\frac{1}{s_x} & 0 & 0 \\
0 & \frac{1}{s_y} & 0 \\
0 & 0 & 1
\end{bmatrix}
\]

(25)

The inverse matrix generates an opposite scaling transformation, so the multiplication of any scaling matrix with its inverse produces the identity matrix.

4 Two-Dimensional Composite Transformations

Using matrix representations, we can set up a sequence of transformations as a composite transformation matrix by calculating the product of the individual transformations. Forming products of transformation matrices is often referred to as a concatenation, or composition, of matrices. Because a coordinate position is represented with a homogeneous column matrix, we must premultiply the column matrix by the matrices representing any transformation sequence. Also, because many positions in a scene are typically transformed by the same sequence, it is more efficient to first multiply the transformation matrices to form a single composite matrix. Thus, if we want to apply two transformations to point position \( P \), the transformed location would be calculated as

\[
P' = M_2 \cdot M_1 \cdot P
\]

(26)
The coordinate position is transformed using the composite matrix $M$, rather than applying the individual transformations $M_1$ and then $M_2$.

**Composite Two-DimensionalTranslations**

If two successive translation vectors $(t_{1x}, t_{1y})$ and $(t_{2x}, t_{2y})$ are applied to a two-dimensional coordinate position $P$, the final transformed location $P'$ is calculated as

$$ P' = T(t_{2x}, t_{2y}) \cdot (T(t_{1x}, t_{1y}) \cdot P) $$

$$ = (T(t_{2x}, t_{2y}) \cdot T(t_{1x}, t_{1y})) \cdot P $$

(27)

where $P$ and $P'$ are represented as three-element, homogeneous-coordinate column vectors. We can verify this result by calculating the matrix product for the two associative groupings. Also, the composite transformation matrix for this sequence of translations is

$$ \begin{bmatrix} 1 & 0 & t_{2x} \\ 0 & 1 & t_{2y} \\ 0 & 0 & 1 \end{bmatrix} \cdot \begin{bmatrix} 1 & 0 & t_{1x} \\ 0 & 1 & t_{1y} \\ 0 & 0 & 1 \end{bmatrix} = \begin{bmatrix} 1 & 0 & t_{1x} + t_{2x} \\ 0 & 1 & t_{1y} + t_{2y} \\ 0 & 0 & 1 \end{bmatrix} $$

(28)

or

$$ T(t_{2x}, t_{2y}) \cdot T(t_{1x}, t_{1y}) = T(t_{1x} + t_{2x}, t_{1y} + t_{2y}) $$

(29)

which demonstrates that two successive translations are additive.

**Composite Two-Dimensional Rotations**

Two successive rotations applied to a point $P$ produce the transformed position

$$ P' = R(\theta_2) \cdot (R(\theta_1) \cdot P) $$

$$ = (R(\theta_2) \cdot R(\theta_1)) \cdot P $$

(30)

By multiplying the two rotation matrices, we can verify that two successive rotations are additive:

$$ R(\theta_2) \cdot R(\theta_1) = R(\theta_1 + \theta_2) $$

(31)

so that the final rotated coordinates of a point can be calculated with the composite rotation matrix as

$$ P' = R(\theta_1 + \theta_2) \cdot P $$

(32)

**Composite Two-Dimensional scalings**

Concatenating transformation matrices for two successive scaling operations in two dimensions produces the following composite scaling matrix:

$$ \begin{bmatrix} s_{2x} & 0 & 0 \\ 0 & s_{2y} & 0 \\ 0 & 0 & 1 \end{bmatrix} \cdot \begin{bmatrix} s_{1x} & 0 & 0 \\ 0 & s_{1y} & 0 \\ 0 & 0 & 1 \end{bmatrix} = \begin{bmatrix} s_{1x} \cdot s_{2x} & 0 & 0 \\ 0 & s_{1y} \cdot s_{2y} & 0 \\ 0 & 0 & 1 \end{bmatrix} $$

(33)

or

$$ S(s_{2x}, s_{2y}) \cdot S(s_{1x}, s_{1y}) = S(s_{1x} \cdot s_{2x}, s_{1y} \cdot s_{2y}) $$

(34)

The resulting matrix in this case indicates that successive scaling operations are multiplicative. That is, if we were to triple the size of an object twice in succession, the final size would be nine times that of the original.
General Two-Dimensional Pivot-Point Rotation

When a graphics package provides only a rotate function with respect to the coordinate origin, we can generate a two-dimensional rotation about any other pivot point \((x_r, y_r)\) by performing the following sequence of translate-rotate-translate operations:

1. Translate the object so that the pivot-point position is moved to the coordinate origin.
2. Rotate the object about the coordinate origin.
3. Translate the object so that the pivot point is returned to its original position.

This transformation sequence is illustrated in Figure 9. The composite transformation matrix for this sequence is obtained with the concatenation

\[
\begin{bmatrix}
1 & 0 & x_r \\
0 & 1 & y_r \\
0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
\cos \theta & -\sin \theta & 0 \\
\sin \theta & \cos \theta & 0 \\
0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
1 & 0 & -x_r \\
0 & 1 & -y_r \\
0 & 0 & 1
\end{bmatrix}
\]

\[
= \begin{bmatrix}
\cos \theta & -\sin \theta & x_r(1 - \cos \theta) + y_r \sin \theta \\
\sin \theta & \cos \theta & y_r(1 - \cos \theta) - x_r \sin \theta \\
0 & 0 & 1
\end{bmatrix}
\]

which can be expressed in the form

\[
T(x_r, y_r) \cdot R(\theta) \cdot T(-x_r, -y_r) = R(x_r, y_r, \theta)
\]  
(35)

where \(T(-x_r, -y_r) = T^{-1}(x_r, y_r)\). In general, a rotate function in a graphics library could be structured to accept parameters for pivot-point coordinates, as well as the rotation angle, and to generate automatically the rotation matrix of Equation 35.

General Two-Dimensional Fixed-Point Scaling

Figure 10 illustrates a transformation sequence to produce a two-dimensional scaling with respect to a selected fixed position \((x_f, y_f)\), when we have a function that can scale relative to the coordinate origin only. This sequence is

1. Translate the object so that the fixed point coincides with the coordinate origin.
2. Scale the object with respect to the coordinate origin.

3. Use the inverse of the translation in step (1) to return the object to its original position.

Concatenating the matrices for these three operations produces the required scaling matrix:

\[
\begin{bmatrix}
1 & 0 & x_f \\
0 & 1 & y_f \\
0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
s_x & 0 & 0 \\
0 & s_y & 0 \\
0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
1 & 0 & -x_f \\
0 & 1 & -y_f \\
0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
s_x & 0 & x_f (1 - s_x) \\
0 & s_y & y_f (1 - s_y) \\
0 & 0 & 1
\end{bmatrix}
\]

or

\[
T(x_f, y_f) \cdot S(s_x, s_y) \cdot T(-x_f, -y_f) = S(x_f, y_f, s_x, s_y)
\]

This transformation is generated automatically in systems that provide a scale function that accepts coordinates for the fixed point.

**General Two-Dimensional Scaling Directions**

Parameters \(s_x\) and \(s_y\) scale objects along the \(x\) and \(y\) directions. We can scale an object in other directions by rotating the object to align the desired scaling directions with the coordinate axes before applying the scaling transformation.

Suppose we want to apply scaling factors with values specified by parameters \(s_1\) and \(s_2\) in the directions shown in Figure 11. To accomplish the scaling without changing the orientation of the object, we first perform a rotation so that the directions for \(s_1\) and \(s_2\) coincide with the \(x\) and \(y\) axes, respectively. Then the scaling transformation \(S(s_1, s_2)\) is applied, followed by an opposite rotation to return points to their original orientations. The composite matrix resulting from the product of these three transformations is

\[
R^{-1}(\theta) \cdot S(s_1, s_2) \cdot R(\theta) = \begin{bmatrix}
s_1 \cos^2 \theta + s_2 \sin^2 \theta & (s_2 - s_1) \cos \theta \sin \theta & 0 \\
(s_2 - s_1) \cos \theta \sin \theta & s_1 \sin^2 \theta + s_2 \cos^2 \theta & 0 \\
0 & 0 & 1
\end{bmatrix}
\]

As an example of this scaling transformation, we turn a unit square into a parallelogram (Figure 12) by stretching it along the diagonal from \((0, 0)\) to \((1, 1)\). We first rotate the diagonal onto the \(y\) axis using \(\theta = 45^\circ\), then we double its length with the scaling values \(s_1 = 1\) and \(s_2 = 2\), and then we rotate again to return the diagonal to its original orientation.

In Equation 39, we assumed that scaling was to be performed relative to the origin. We could take this scaling operation one step further and concatenate the matrix with translation operators, so that the composite matrix would include parameters for the specification of a scaling fixed position.
Matrix Concatenation Properties

Multiplication of matrices is associative. For any three matrices, \( M_1, M_2, \) and \( M_3, \) the matrix product \( M_3 \cdot M_2 \cdot M_1 \) can be performed by first multiplying \( M_3 \) and \( M_2 \) or by first multiplying \( M_2 \) and \( M_1 \):

\[
M_3 \cdot M_2 \cdot M_1 = (M_3 \cdot M_2) \cdot M_1 = M_3 \cdot (M_2 \cdot M_1)
\]

Therefore, depending upon the order in which the transformations are specified, we can construct a composite matrix either by multiplying from left to right (premultiplying) or by multiplying from right to left (postmultiplying). Some graphics packages require that transformations be specified in the order in which they are to be applied. In that case, we would first invoke transformation \( M_1, \) then \( M_2, \) then \( M_3. \) As each successive transformation routine is called, its matrix is concatenated on the left of the previous matrix product. Other graphics systems, however, postmultiply matrices, so that this transformation sequence would have to be invoked in the reverse order: the last transformation invoked (which is \( M_1 \) for this example) is the first to be applied, and the first transformation that is called (\( M_3 \) in this case) is the last to be applied.

Transformation products, on the other hand, may not be commutative. The matrix product \( M_2 \cdot M_1 \) is not equal to \( M_1 \cdot M_2, \) in general. This means that if we want to translate and rotate an object, we must be careful about the order in which the composite matrix is evaluated (Figure 13). For some special cases—such as a sequence of transformations that are all of the same kind—the multiplication of transformation matrices is commutative. As an example, two successive rotations could be performed in either order and the final position would be the same. This commutative property holds also for two successive translations or two successive scalings. Another commutative pair of operations is rotation and uniform scaling (\( s_x = s_y \)).

Reversing the order in which a sequence of transformations is performed may affect the transformed position of an object. In (a), an object is first translated in the \( x \) direction, then rotated counterclockwise through an angle of \( 45^\circ. \) In (b), the object is first rotated \( 45^\circ \) counterclockwise, then translated in the \( x \) direction.
General Two-Dimensional Composite Transformations
and Computational Efficiency

A two-dimensional transformation, representing any combination of translations, rotations, and scalings, can be expressed as

\[
\begin{bmatrix}
  x' \\
  y' \\
  1
\end{bmatrix} =
\begin{bmatrix}
  rs_{xx} & rs_{xy} & trs_x \\
  rs_{yx} & rs_{yy} & trs_y \\
  0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
  x \\
  y \\
  1
\end{bmatrix}
\tag{41}
\]

The four elements \(rs_{ji}\) are the multiplicative rotation-scaling terms in the transformation, which involve only rotation angles and scaling factors. Elements \(trs_x\) and \(trs_y\) are the translational terms, containing combinations of translation distances, pivot-point and fixed-point coordinates, rotation angles, and scaling parameters. For example, if an object is to be scaled and rotated about its centroid coordinates \((x_c, y_c)\) and then translated, the values for the elements of the composite transformation matrix are

\[
T(t_x, t_y) \cdot R(x_c, y_c, \theta) \cdot S(x_c, y_c, s_x, s_y)
\]

\[
= \begin{bmatrix}
  s_x \cos \theta & -s_x \sin \theta & x_c(1 - s_x \cos \theta) + y_cs_y \sin \theta + t_x \\
  s_y \cos \theta & s_y \sin \theta & y_c(1 - s_y \cos \theta) - x_cs_x \sin \theta + t_y \\
  0 & 0 & 1
\end{bmatrix}
\tag{42}
\]

Although Equation 41 requires nine multiplications and six additions, the explicit calculations for the transformed coordinates are

\[
x' = x \cdot rs_{xx} + y \cdot rs_{xy} + trs_x, \quad y' = x \cdot rs_{yx} + y \cdot rs_{yy} + trs_y
\tag{43}
\]

Thus, we need actually perform only four multiplications and four additions to transform coordinate positions. This is the maximum number of computations required for any transformation sequence, once the individual matrices have been concatenated and the elements of the composite matrix evaluated. Without concatenation, the individual transformations would be applied one at a time, and the number of calculations could be increased significantly. An efficient implementation for the transformation operations, therefore, is to formulate transformation matrices, concatenate any transformation sequence, and calculate transformed coordinates using Equations 43. On parallel systems, direct matrix multiplications with the composite transformation matrix of Equation 41 can be equally efficient.

Because rotation calculations require trigonometric evaluations and several multiplications for each transformed point, computational efficiency can become an important consideration in rotation transformations. In animations and other applications that involve many repeated transformations and small rotation angles, we can use approximations and iterative calculations to reduce computations in the composite transformation equations. When the rotation angle is small, the trigonometric functions can be replaced with approximation values based on the first few terms of their power series expansions. For small-enough angles (less than \(10^\circ\)), \(\cos \theta\) is approximately 1.0 and \(\sin \theta\) has a value very close to the value of \(\theta\) in radians. If we are rotating in small angular steps about the origin, for instance, we can set \(\cos \theta\) to 1.0 and reduce transformation calculations at each step to two multiplications and two additions for each set of coordinates to be rotated. These rotation calculations are

\[
x' = x - y \sin \theta, \quad y' = x \sin \theta + y
\tag{44}
\]

where \(\sin \theta\) is evaluated once for all steps, assuming the rotation angle does not change. The error introduced by this approximation at each step decreases as the rotation angle decreases; but even with small rotation angles, the accumulated
error over many steps can become quite large. We can control the accumulated error by estimating the error in \( x' \) and \( y' \) at each step and resetting object positions when the error accumulation becomes too great. Some animation applications automatically reset object positions at fixed intervals, such as every 360° or every 180°.

Composite transformations often involve inverse matrices. For example, transformation sequences for general scaling directions and for some reflections and shears (Section 5) require inverse rotations. As we have noted, the inverse matrix representations for the basic geometric transformations can be generated with simple procedures. An inverse translation matrix is obtained by changing the signs of the translation distances, and an inverse rotation matrix is obtained by performing a matrix transpose (or changing the sign of the sine terms). These operations are much simpler than direct inverse matrix calculations.

**Two-Dimensional Rigid-Body Transformation**

If a transformation matrix includes only translation and rotation parameters, it is a **rigid-body transformation matrix**. The general form for a two-dimensional rigid-body transformation matrix is

\[
\begin{bmatrix}
  r_{xx} & r_{xy} & t_x \\
  r_{yx} & r_{yy} & t_y \\
  0 & 0 & 1
\end{bmatrix}
\] (45)

where the four elements \( r_{jk} \) are the multiplicative rotation terms, and the elements \( t_x \) and \( t_y \) are the translational terms. A rigid-body change in coordinate position is also sometimes referred to as a **rigid-motion** transformation. All angles and distances between coordinate positions are unchanged by the transformation. In addition, matrix 45 has the property that its upper-left \( 2 \times 2 \) submatrix is an **orthogonal matrix**. This means that if we consider each row (or each column) of the submatrix as a vector, then the two row vectors \((r_{xx}, r_{xy})\) and \((r_{yx}, r_{yy})\) (or the two column vectors) form an orthogonal set of unit vectors. Such a set of vectors is also referred to as an **orthonormal** vector set. Each vector has unit length as follows:

\[
r_{xx}^2 + r_{xy}^2 = r_{yx}^2 + r_{yy}^2 = 1
\] (46)

and the vectors are perpendicular (their dot product is 0):

\[
r_{xx}r_{yx} + r_{xy}r_{yy} = 0
\] (47)

Therefore, if these unit vectors are transformed by the rotation submatrix, then the vector \((r_{xx}, r_{xy})\) is converted to a unit vector along the \( x \) axis and the vector \((r_{yx}, r_{yy})\) is transformed into a unit vector along the \( y \) axis of the coordinate system:

\[
\begin{bmatrix}
  r_{xx} & r_{xy} \\
  r_{yx} & r_{yy}
\end{bmatrix}
\begin{bmatrix}
  0 \\
  0
\end{bmatrix}
= \begin{bmatrix}
  1 \\
  0
\end{bmatrix}
\]

(48)

\[
\begin{bmatrix}
  r_{xx} & r_{xy} \\
  r_{yx} & r_{yy}
\end{bmatrix}
\begin{bmatrix}
  0 \\
  1
\end{bmatrix}
= \begin{bmatrix}
  0 \\
  1
\end{bmatrix}
\]

(49)

For example, the following rigid-body transformation first rotates an object through an angle \( \theta \) about a pivot point \((x_s, y_s)\) and then translates the object:

\[
T(t_x, t_y) \cdot R(x_r, y_r, \theta) = \begin{bmatrix}
  \cos \theta & -\sin \theta & x_s(1 - \cos \theta) + y_s \sin \theta + t_x \\
  \sin \theta & \cos \theta & y_s(1 - \cos \theta) - x_s \sin \theta + t_y \\
  0 & 0 & 1
\end{bmatrix}
\] (50)
Here, orthogonal unit vectors in the upper-left $2 \times 2$ submatrix are $(\cos \theta, -\sin \theta)$ and $(\sin \theta, \cos \theta)$, and
\[
\begin{bmatrix}
\cos \theta & -\sin \theta & 0 \\
\sin \theta & \cos \theta & 0 \\
0 & 0 & 1
\end{bmatrix} \cdot 
\begin{bmatrix}
\cos \theta \\
-\sin \theta \\
1
\end{bmatrix} = 
\begin{bmatrix}
1 \\
0 \\
1
\end{bmatrix}
\] (51)

Similarly, unit vector $(\sin \theta, \cos \theta)$ is converted by the preceding transformation matrix to the unit vector $(0, 1)$ in the $y$ direction.

**Constructing Two-Dimensional Rotation Matrices**

The orthogonal property of rotation matrices is useful for constructing the matrix when we know the final orientation of an object, rather than the amount of angular rotation necessary to put the object into that position. This orientation information could be determined by the alignment of certain objects in a scene or by reference positions within the coordinate system. For example, we might want to rotate an object to align its axis of symmetry with the viewing (camera) direction, or we might want to rotate one object so that it is above another object. Figure 14 shows an object that is to be aligned with the unit direction vectors $\mathbf{u}'$ and $\mathbf{v}'$. Assuming that the original object orientation, as shown in Figure 14(a), is aligned with the coordinate axes, we construct the desired transformation by assigning the elements of $\mathbf{u}'$ to the first row of the rotation matrix and the elements of $\mathbf{v}'$ to the second row. In a modeling application, for instance, we can use this method to obtain the transformation matrix within an object’s local coordinate system when we know what its orientation is to be within the overall world-coordinate scene. A similar transformation is the conversion of object descriptions from one coordinate system to another, and we take up these methods in more detail in Section 8.

**Two-Dimensional Composite-Matrix Programming Example**

An implementation example for a sequence of geometric transformations is given in the following program. Initially, the composite matrix, `compMatrix`, is constructed as the identity matrix. For this example, a left-to-right concatenation order is used to construct the composite transformation matrix, and we invoke the transformation routines in the order that they are to be executed. As each of the basic transformation routines (scale, rotate, and translate) is invoked, a matrix is set up for that transformation and left-concatenated with the composite matrix. When all transformations have been specified, the composite transformation is applied to transform a triangle. The triangle is first scaled with respect to its centroid position, then the triangle is rotated about its centroid, and, lastly, it is translated. Figure 15 shows the original and final positions of the triangle that is transformed by this sequence. Routines in OpenGL are used to display the initial and final position of the triangle.
A triangle (a) is transformed into position (b) using the composite-matrix calculations in procedure transformVerts2D.

```c
#include <GL/glut.h>
#include <stdlib.h>
#include <math.h>

/* Set initial display-window size. */
GLsizei winWidth = 600, winHeight = 600;

/* Set range for world coordinates. */
GLfloat xwcMin = 0.0, xwcMax = 225.0;
GLfloat ywcMin = 0.0, ywcMax = 225.0;

class wcPt2D {
    public:
        GLfloat x, y;
};

typedef GLfloat Matrix3x3 [3][3];
Matrix3x3 matComposite;

const GLdouble pi = 3.14159;

void init (void)
{
    /* Set color of display window to white. */
    glClearColor (1.0, 1.0, 1.0, 0.0);
}

/* Construct the 3 x 3 identity matrix. */
void matrix3x3SetIdentity (Matrix3x3 matIdent3x3)
{
    GLint row, col;

    for (row = 0; row < 3; row++)
        for (col = 0; col < 3; col++)
            matIdent3x3 [row][col] = (row == col);
}
```
/* Premultiply matrix m1 times matrix m2, store result in m2. */
void matrix3x3PreMultiply (Matrix3x3 m1, Matrix3x3 m2)
{
    GLint row, col;
    Matrix3x3 matTemp;
    for (row = 0; row < 3; row++)
        for (col = 0; col < 3; col++)
            matTemp [row][col] = m1 [row][0] * m2 [0][col] + m1 [row][1] * m2 [1][col] + m1 [row][2] * m2 [2][col];
    for (row = 0; row < 3; row++)
        for (col = 0; col < 3; col++)
            m2 [row][col] = matTemp [row][col];
}

void translate2D (GLfloat tx, GLfloat ty)
{
    Matrix3x3 matTransl;
    /* Initialize translation matrix to identity. */
    matrix3x3SetIdentity (matTransl);
    matTransl [0][2] = tx;
    matTransl [1][2] = ty;
    /* Concatenate matTransl with the composite matrix. */
    matrix3x3PreMultiply (matTransl, matComposite);
}

void rotate2D (wcPt2D pivotPt, GLfloat theta)
{
    Matrix3x3 matRot;
    /* Initialize rotation matrix to identity. */
    matrix3x3SetIdentity (matRot);
    matRot [0][0] = cos (theta);
    matRot [0][1] = -sin (theta);
    matRot [0][2] = pivotPt.x * (1 - cos (theta)) + pivotPt.y * sin (theta);
    matRot [1][0] = sin (theta);
    matRot [1][1] = cos (theta);
    matRot [1][2] = pivotPt.y * (1 - cos (theta)) - pivotPt.x * sin (theta);
    /* Concatenate matRot with the composite matrix. */
    matrix3x3PreMultiply (matRot, matComposite);
}

void scale2D (GLfloat sx, GLfloat sy, wcPt2D fixedPt)
{
    Matrix3x3 matScale;
/* Set geometric transformation parameters. */
wcPt2D pivPt, fixedPt;
pivPt = centroidPt;
fixedPt = centroidPt;

GLfloat tx = 0.0, ty = 100.0;
GLfloat sx = 0.5, sy = 0.5;
GLdouble theta = pi/2.0;

g1Clear (GL_COLOR_BUFFER_BIT); // Clear display window.
g1Color3f (0.0, 0.0, 1.0); // Set initial fill color to blue.
triangle (verts); // Display blue triangle.

/* Initialize composite matrix to identity. */
matrix3x3SetIdentity (matComposite);

/* Construct composite matrix for transformation sequence. */
scale2D (sx, sy, fixedPt); // First transformation: Scale.
rotate2D (pivPt, theta); // Second transformation: Rotate
translate2D (tx, ty); // Final transformation: Translate.

/* Apply composite matrix to triangle vertices. */
transformVerts2D (nVerts, verts);
g1Color3f (1.0, 0.0, 0.0); // Set color for transformed triangle.
triangle (verts); // Display red transformed triangle.

g1Flush ( );
}

void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ( );
    gluOrtho2D (xwcMin, xwcMax, ywcMin, ywcMax);
    g1Clear (GL_COLOR_BUFFER_BIT);
}

void main (int argc, char ** argv)
{
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowSize (50, 50);
    glutCreateWindow ("Geometric Transformation Sequence");
    init ( );
    glutDisplayFunc (displayFcn);
    glutReshapeFunc (winReshapeFcn);
    glutMainLoop ( );
}
/ Initialize scaling matrix to identity. */
matrix3x3SetIdentity (matScale);

matScale [0][0] = sx;
matScale [0][2] = (1 - sx) * fixedPt.x;
matScale [1][1] = sy;
matScale [1][2] = (1 - sy) * fixedPt.y;

/* Concatenate matScale with the composite matrix. */
matrix3x3PreMultiply (matScale, matComposite);

}/* Using the composite matrix, calculate transformed coordinates. */
void transformVerts2D (GLint nVerts, wcPt2D * verts)
{
GLint k;
GLfloat temp;
for (k = 0; k < nVerts; k++) {
    temp = matComposite [0][0] * verts[k].x + matComposite [0][1] * verts[k].y + matComposite [0][2];
    verts[k].y = matComposite [1][0] * verts[k].x + matComposite [1][1] * verts[k].y + matComposite [1][2];
    verts[k].x = temp;
}
}

void triangle (wcPt2D * verts)
{
GLint k;

glBegin (GL_TRIANGLES);
    for (k = 0; k < 3; k++)
        glVertex2f (verts[k].x, verts[k].y);
    glEnd ( );
}

void displayFcn (void)
{
/* Define initial position for triangle. */
GLint nVerts = 3;
wcPt2D verts[3] = { {50.0, 25.0}, {150.0, 25.0}, {100.0, 100.0} };

/* Calculate position of triangle centroid. */
wcPt2D centroidPt:

GLint k, xSum = 0, ySum = 0;
for (k = 0; k < nVerts; k++) {
    xSum += verts[k].x;
    ySum += verts[k].y;
}
centroidPt.x = GLfloat (xSum) / GLfloat (nVerts);
centroidPt.y = GLfloat (ySum) / GLfloat (nVerts);
5 Other Two-Dimensional Transformations

Basic transformations such as translation, rotation, and scaling are standard components of graphics libraries. Some packages provide a few additional transformations that are useful in certain applications. Two such transformations are reflection and shear.

Reflection

A transformation that produces a mirror image of an object is called a reflection. For a two-dimensional reflection, this image is generated relative to an axis of reflection by rotating the object 180° about the reflection axis. We can choose an axis of reflection in the $xy$ plane or perpendicular to the $xy$ plane. When the reflection axis is a line in the $xy$ plane, the rotation path about this axis is in a plane perpendicular to the $xy$ plane. For reflection axes that are perpendicular to the $xy$ plane, the rotation path is in the $xy$ plane. Some examples of common reflections follow.

Reflection about the line $y = 0$ (the $x$ axis) is accomplished with the transformation matrix

$$\begin{bmatrix} 1 & 0 & 0 \\ 0 & -1 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

This transformation retains $x$ values, but “flips” the $y$ values of coordinate positions. The resulting orientation of an object after it has been reflected about the $x$ axis is shown in Figure 16. To envision the rotation transformation path for this reflection, we can think of the flat object moving out of the $xy$ plane and rotating 180° through three-dimensional space about the $x$ axis and back into the $xy$ plane on the other side of the $x$ axis.

A reflection about the line $x = 0$ (the $y$ axis) flips $x$ coordinates while keeping $y$ coordinates the same. The matrix for this transformation is

$$\begin{bmatrix} -1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

Figure 17 illustrates the change in position of an object that has been reflected about the line $x = 0$. The equivalent rotation in this case is 180° through three-dimensional space about the $y$ axis.

We flip both the $x$ and $y$ coordinates of a point by reflecting relative to an axis that is perpendicular to the $xy$ plane and that passes through the coordinate origin. This reflection is sometimes referred to as a reflection relative to the coordinate origin, and it is equivalent to reflecting with respect to both coordinate axes. The matrix representation for this reflection is

$$\begin{bmatrix} -1 & 0 & 0 \\ 0 & -1 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

An example of reflection about the origin is shown in Figure 18. The reflection matrix 54 is the same as the rotation matrix $R(\theta)$ with $\theta = 180^\circ$. We are simply rotating the object in the $xy$ plane half a revolution about the origin.
Reflection 54 can be generalized to any reflection point in the $xy$ plane (Figure 19). This reflection is the same as a $180^\circ$ rotation in the $xy$ plane about the reflection point.

If we choose the reflection axis as the diagonal line $y = x$ (Figure 20), the reflection matrix is

$$
\begin{bmatrix}
0 & 1 & 0 \\
1 & 0 & 0 \\
0 & 0 & 1
\end{bmatrix}
$$

We can derive this matrix by concatenating a sequence of rotation and coordinate-axis reflection matrices. One possible sequence is shown in Figure 21. Here, we first perform a clockwise rotation with respect to the origin through a $45^\circ$ angle, which rotates the line $y = x$ onto the $x$ axis. Next, we perform a reflection with respect to the $x$ axis. The final step is to rotate the line $y = x$ back to its original position with a counterclockwise rotation through $45^\circ$. Another equivalent sequence of transformations is to first reflect the object about the $x$ axis, then rotate it counterclockwise $90^\circ$.

To obtain a transformation matrix for reflection about the diagonal $y = -x$, we could concatenate matrices for the transformation sequence: (1) clockwise rotation by $45^\circ$, (2) reflection about the $y$ axis, and (3) counterclockwise rotation by $45^\circ$. The resulting transformation matrix is

$$
\begin{bmatrix}
0 & -1 & 0 \\
-1 & 0 & 0 \\
0 & 0 & 1
\end{bmatrix}
$$

Figure 22 shows the original and final positions for an object transformed with this reflection matrix.

Reflections about any line $y = mx + b$ in the $xy$ plane can be accomplished with a combination of translate-rotate-reflect transformations. In general, we
first translate the line so that it passes through the origin. Then we can rotate the line onto one of the coordinate axes and reflect about that axis. Finally, we restore the line to its original position with the inverse rotation and translation transformations.

We can implement reflections with respect to the coordinate axes or coordinate origin as scaling transformations with negative scaling factors. Also, elements of the reflection matrix can be set to values other than ±1. A reflection parameter with a magnitude greater than 1 shifts the mirror image of a point farther from the reflection axis, and a parameter with magnitude less than 1 brings the mirror image of a point closer to the reflection axis. Thus, a reflected object can also be enlarged, reduced, or distorted.

Shear

A transformation that distorts the shape of an object such that the transformed shape appears as if the object were composed of internal layers that had been caused to slide over each other is called a shear. Two common shearing transformations are those that shift coordinate x values and those that shift y values.

An x-direction shear relative to the x axis is produced with the transformation matrix

\[
\begin{bmatrix}
1 & s_h \cdot x & 0 \\
0 & 1 & 0 \\
0 & 0 & 1
\end{bmatrix}
\]

which transforms coordinate positions as

\[
x' = x + s_h \cdot y, \quad y' = y
\]  

Any real number can be assigned to the shear parameter \( s_h \). A coordinate position \((x, y)\) is then shifted horizontally by an amount proportional to its perpendicular distance \((y\) value\) from the \(x\) axis. Setting parameter \( s_h \) to the value 2, for example, changes the square in Figure 23 into a parallelogram. Negative values for \( s_h \) shift coordinate positions to the left.
We can generate $x$-direction shears relative to other reference lines with

$$
\begin{bmatrix}
1 & sh_x & -sh_x \cdot y_{ref} \\
0 & 1 & 0 \\
0 & 0 & 1
\end{bmatrix}
$$

(59)

Now, coordinate positions are transformed as

$$
x' = x + sh_x(y - y_{ref}), \quad y' = y
$$

(60)

An example of this shearing transformation is given in Figure 24 for a shear parameter value of $\frac{1}{2}$ relative to the line $y_{ref} = -1$.

A $y$-direction shear relative to the line $x = x_{ref}$ is generated with the transformation matrix

$$
\begin{bmatrix}
1 & 0 & 0 \\
sh_y & 1 & -sh_y \cdot x_{ref} \\
0 & 0 & 1
\end{bmatrix}
$$

(61)

which generates the transformed coordinate values

$$
x' = x, \quad y' = y + sh_y(x - x_{ref})
$$

(62)

This transformation shifts a coordinate position vertically by an amount proportional to its distance from the reference line $x = x_{ref}$. Figure 25 illustrates the conversion of a square into a parallelogram with $sh_y = 0.5$ and $x_{ref} = -1$.

Shearing operations can be expressed as sequences of basic transformations. The $x$-direction shear matrix 57, for example, can be represented as a composite transformation involving a series of rotation and scaling matrices. This composite transformation scales the unit square of Figure 23 along its diagonal, while maintaining the original lengths and orientations of edges parallel to the $x$ axis. Shifts in the positions of objects relative to shearing reference lines are equivalent to translations.
6 Raster Methods for Geometric Transformations

The characteristics of raster systems suggest an alternate method for performing certain two-dimensional transformations. Raster systems store picture information as color patterns in the frame buffer. Therefore, some simple object transformations can be carried out rapidly by manipulating an array of pixel values. Few arithmetic operations are needed, so the pixel transformations are particularly efficient.

Functions that manipulate rectangular pixel arrays are called raster operations and moving a block of pixel values from one position to another is termed a block transfer, a bitblt, or a pixblt. Routines for performing some raster operations are usually available in a graphics package.

Figure 26 illustrates a two-dimensional translation implemented as a block transfer of a refresh-buffer area. All bit settings in the rectangular area shown are copied as a block into another part of the frame buffer. We can erase the pattern at the original location by assigning the background color to all pixels within that block (assuming that the pattern to be erased does not overlap other objects in the scene).

Rotations in 90-degree increments are accomplished easily by rearranging the elements of a pixel array. We can rotate a two-dimensional object or pattern 90° counterclockwise by reversing the pixel values in each row of the array, then interchanging rows and columns. A 180° rotation is obtained by reversing the order of the elements in each row of the array, then reversing the order of the rows.

Figure 27 demonstrates the array manipulations that can be used to rotate a pixel block by 90° and by 180°.

For array rotations that are not multiples of 90°, we need to do some extra processing. The general procedure is illustrated in Figure 28. Each destination pixel area is mapped onto the rotated array and the amount of overlap with the rotated pixel areas is calculated. A color for a destination pixel can then be computed by averaging the colors of the overlapped source pixels, weighted by their percentage of area overlap. Alternatively, we could use an approximation method, as in antialiasing, to determine the color of the destination pixels.

We can use similar methods to scale a block of pixels. Pixel areas in the original block are scaled, using specified values for $s_x$ and $s_y$, and then mapped onto a set of destination pixels. The color of each destination pixel is then assigned according to its area of overlap with the scaled pixel areas (Figure 29).

An object can be reflected using raster transformations that reverse row or column values in a pixel block, combined with translations. Shears are produced with shifts in the positions of array values along rows or columns.
7 OpenGL Raster Transformations

You should already be familiar with most of the OpenGL functions for performing raster operations. A translation of a rectangular array of pixel-color values from one buffer area to another can be accomplished in OpenGL as the following copy operation:

\[
\text{glCopyPixels (xmin, ymin, width, height, GL\_COLOR);} \]

The first four parameters in this function give the location and dimensions of the pixel block; and the OpenGL symbolic constant GL\_COLOR specifies that it is color values are to be copied. This array of pixels is to be copied to a rectangular area of a refresh buffer whose lower-left corner is at the location specified by the current raster position. Pixel-color values are copied as either RGBA values or color-table indices, depending on the current setting for the color mode. Both the region to be copied (the source) and the destination area should lie completely within the bounds of the screen coordinates. This translation can be carried out on any of the OpenGL buffers used for refreshing, or even between different buffers. A source buffer for the glCopyPixels function is chosen with the glReadBuffer routine, and a destination buffer is selected with the glDrawBuffer routine.

We can rotate a block of pixel-color values in 90-degree increments by first saving the block in an array, then rearranging the elements of the array and placing it back in the refresh buffer. A block of RGB color values in a buffer can be saved in an array with the function

\[
\text{glReadPixels (xmin, ymin, width, height, GL\_RGB,}
\text{ GL\_UNSIGNED\_BYTE, colorArray);} \]

If color-table indices are stored at the pixel positions, we replace the constant GL\_RGB with GL\_COLOR\_INDEX. To rotate the color values, we rearrange the rows and columns of the color array, as described in the previous section. Then we put the rotated array back in the buffer with

\[
\text{glDrawPixels (width, height, GL\_RGB, GL\_UNSIGNED\_BYTE,}
\text{ colorArray);} \]

The lower-left corner of this array is placed at the current raster position. We select the source buffer containing the original block of pixel values with glReadBuffer, and we designate a destination buffer with glDrawBuffer.
A two-dimensional scaling transformation can be performed as a raster operation in OpenGL by specifying scaling factors and then invoking either glCopyPixels or glDrawPixels. For the raster operations, we set the scaling factors with

\[ \text{glPixelZoom} \left( \text{sx}, \text{sy} \right); \]

where parameters \( \text{sx} \) and \( \text{sy} \) can be assigned any nonzero floating-point values. Positive values greater than 1.0 increase the size of an element in the source array, and positive values less than 1.0 decrease element size. A negative value for \( \text{sx} \) or \( \text{sy} \), or both, produces a reflection and scales the array elements. Thus, if \( \text{sx} = \text{sy} = -3.0 \), the source array is reflected with respect to the current raster position and each color element of the array is mapped to a 3 \( \times \) 3 block of destination pixels. If the center of a destination pixel lies within the rectangular area of a scaled color element of an array, it is assigned the color of that array element. Destination pixels whose centers are on the left or top boundary of the scaled array element are also assigned the color of that element. The default value for both \( \text{sx} \) and \( \text{sy} \) is 1.0.

We can also combine raster transformations with logical operations to produce various effects. With the \textit{exclusive or} operator, for example, two successive copies of a pixel array to the same buffer area restores the values that were originally present in that area. This technique can be used in an animation application to translate an object across a scene without altering the background pixels.

\section{Transformations between Two-Dimensional Coordinate Systems}

Computer-graphics applications involve coordinate transformations from one reference frame to another during various stages of scene processing. The viewing routines transform object descriptions from world coordinates to device coordinates. For modeling and design applications, individual objects are typically defined in their own local Cartesian references. These local-coordinate descriptions must then be transformed into positions and orientations within the overall scene coordinate system. A facility-management program for office layouts, for instance, has individual coordinate descriptions for chairs and tables and other furniture that can be placed into a floor plan, with multiple copies of the chairs and other items in different positions.

Also, scenes are sometimes described in non-Cartesian reference frames that take advantage of object symmetries. Coordinate descriptions in these systems must be converted to Cartesian world coordinates for processing. Some examples of non-Cartesian systems are polar coordinates, spherical coordinates, elliptical coordinates, and parabolic coordinates. Here, we consider only the transformations involved in converting from one two-dimensional Cartesian frame to another.

Figure 30 shows a Cartesian \( x'y' \) system specified with coordinate origin \((x_0, y_0)\) and orientation angle \( \theta \) in a Cartesian \( xy \) reference frame. To transform object descriptions from \( xy \) coordinates to \( x'y' \) coordinates, we set up a transformation that superimposes the \( x'y' \) axes onto the \( xy \) axes. This is done in two steps:

1. Translate so that the origin \((x_0, y_0)\) of the \( x'y' \) system is moved to the origin \((0, 0)\) of the \( xy \) system.
2. Rotate the \( x' \) axis onto the \( x \) axis.
Translation of the coordinate origin is accomplished with the matrix transformation
\[
T(-x_0, -y_0) = \begin{bmatrix} 1 & 0 & -x_0 \\ 0 & 1 & -y_0 \\ 0 & 0 & 1 \end{bmatrix}
\] (63)

The orientation of the two systems after the translation operation would then appear as in Figure 31. To get the axes of the two systems into coincidence, we then perform the clockwise rotation
\[
R(-\theta) = \begin{bmatrix} \cos \theta & \sin \theta & 0 \\ -\sin \theta & \cos \theta & 0 \\ 0 & 0 & 1 \end{bmatrix}
\] (64)

Concatenating these two transformation matrices gives us the complete composite matrix for transforming object descriptions from the \(xy\) system to the \(x'y'\) system:
\[
M_{xy,x'y'} = R(-\theta) \cdot T(-x_0, -y_0)
\] (65)

An alternate method for describing the orientation of the \(x'y'\) coordinate system is to specify a vector \(V\) that indicates the direction for the positive \(y'\) axis, as shown in Figure 32. We can specify vector \(V\) as a point in the \(xy\) reference frame relative to the origin of the \(xy\) system, which we can convert to the unit vector \(v = \frac{V}{|V|} = (v_x, v_y)\) (66).

We obtain the unit vector \(u\) along the \(x'\) axis by applying a 90° clockwise rotation to vector \(v\):
\[
\begin{align*}
u &= (v_y, -v_x) \\
&= (u_x, u_y)
\end{align*}
\] (67)

In Section 4, we noted that the elements of any rotation matrix could be expressed as elements of a set of orthonormal vectors. Therefore, the matrix to rotate the \(x'y'\) system into coincidence with the \(xy\) system can be written as
\[
R = \begin{bmatrix} u_x & u_y & 0 \\ v_x & v_y & 0 \\ 0 & 0 & 1 \end{bmatrix}
\] (68)

For example, suppose that we choose the orientation for the \(y'\) axis as \(V = (-1, 0)\). Then the \(x'\) axis is in the positive \(y\) direction, and the rotation transformation matrix is
\[
\begin{bmatrix} 0 & 1 & 0 \\ -1 & 0 & 0 \\ 0 & 0 & 1 \end{bmatrix}
\]
Equivalently, we can obtain this rotation matrix from Equation 64 by setting the orientation angle as $\theta = 90^\circ$.

In an interactive application, it may be more convenient to choose the direction of $\mathbf{V}$ relative to position $\mathbf{P}_0$ than to specify it relative to the $xy$-coordinate origin. Unit vectors $\mathbf{u}$ and $\mathbf{v}$ would then be oriented as shown in Figure 33. The components of $\mathbf{v}$ are now calculated as

$$\mathbf{v} = \frac{\mathbf{P}_1 - \mathbf{P}_0}{|\mathbf{P}_1 - \mathbf{P}_0|} \tag{69}$$

and $\mathbf{u}$ is obtained as the perpendicular to $\mathbf{v}$ that forms a right-handed Cartesian system.

## 9 OpenGL Functions for Two-Dimensional Geometric Transformations

In the core library of OpenGL, a separate function is available for each of the basic geometric transformations. Because OpenGL is designed as a three-dimensional graphics application programming interface (API), all transformations are specified in three dimensions. Internally, all coordinate positions are represented as four-element column vectors, and all transformations are represented using $4 \times 4$ matrices. Fortunately, performing two-dimensional transformations within OpenGL is generally just a matter of using a value for the transformation in the third ($z$) dimension that causes no change in that dimension.

To perform a translation, we invoke the translation routine and set the components for the three-dimensional translation vector. In the rotation function, we specify the angle and the orientation for a rotation axis that intersects the coordinate origin. In addition, a scaling function is used to set the three coordinate scaling factors relative to the coordinate origin. In each case, the transformation routine sets up a $4 \times 4$ matrix that is applied to the coordinates of objects that are referenced after the transformation call.

### Basic OpenGL Geometric Transformations

A $4 \times 4$ translation matrix is constructed with the following routine:

```c
glTranslate* (tx, ty, tz);
```

Translation parameters $tx$, $ty$, and $tz$ can be assigned any real-number values, and the single suffix code to be affixed to this function is either f (float) or d (double). For two-dimensional applications, we set $tz = 0.0$; and a two-dimensional position is represented as a four-element column matrix with the $z$ component equal to 0.0. The translation matrix generated by this function is used to transform
positions of objects defined after this function is invoked. For example, we translate subsequently defined coordinate positions 25 units in the $x$ direction and $-10$ units in the $y$ direction with the statement
\[
glTranslatef (25.0, -10.0, 0.0);
\]
Similarly, a $4 \times 4$ rotation matrix is generated with
\[
glRotate* (\theta, vx, vy, vz);
\]
where the vector $v = (vx, vy, vz)$ can have any floating-point values for its components. This vector defines the orientation for a rotation axis that passes through the coordinate origin. If $v$ is not specified as a unit vector, then it is normalized automatically before the elements of the rotation matrix are computed. The suffix code can be either $f$ or $d$, and parameter $\theta$ is to be assigned a rotation angle in degrees, which the routine converts to radians for the trigonometric calculations. The rotation specified here will be applied to positions defined after this function call. Rotation in two-dimensional systems is rotation about the $z$ axis, specified as a unit vector with $x$ and $y$ components of zero, and a $z$ component of 1.0. For example, the statement
\[
glRotatef (90.0, 0.0, 0.0, 1.0);
\]
sets up the matrix for a $90^\circ$ rotation about the $z$ axis. We should note here that internally, this function generates a rotation matrix using quaternions. This method is more efficient when rotation is about an arbitrarily-specific axis.

We obtain a $4 \times 4$ scaling matrix with respect to the coordinate origin with the following routine:
\[
glScale* (sx, sy, sz);
\]
The suffix code is again either $f$ or $d$, and the scaling parameters can be assigned any real-number values. Scaling in a two-dimensional system involves changes in the $x$ and $y$ dimensions, so a typical two-dimensional scaling operation has a $z$ scaling factor of 1.0 (which causes no change in the $z$ coordinate of positions). Because the scaling parameters can be any real-number value, this function will also generate reflections when negative values are assigned to the scaling parameters. For example, the following statement produces a matrix that scales by a factor of 2 in the $x$ direction, scales by a factor of 3 in the $y$ direction, and reflects with respect to the $x$ axis:
\[
glScalef (2.0, -3.0, 1.0);
\]
A zero value for any scaling parameter can cause a processing error because an inverse matrix cannot be calculated. The scale-reflect matrix is applied to subsequently defined objects.

It is important to note that internally OpenGL uses composite matrices to hold transformations. As a result, transformations are cumulative—that is, if we apply a translation and then apply a rotation, objects whose positions are specified after that will have both transformations applied to them. If that is not the behavior we desired, we must be able to remove the effects of previous transformations. This requires additional functions for manipulating the composite matrices.
OpenGL Matrix Operations

The glMatrixMode routine is used to set the projection mode, which designates the matrix that is to be used for the projection transformation. This transformation determines how a scene is to be projected onto the screen. We use the same routine to set up a matrix for the geometric transformations. In this case, however, the matrix is referred to as the modelview matrix, and it is used to store and combine the geometric transformations. It is also used to combine the geometric transformations with the transformation to a viewing-coordinate system. We specify the modelview mode with the statement

\[
\text{glMatrixMode (GL_MODELVIEW);}
\]

which designates the 4 x 4 modelview matrix as the current matrix. The OpenGL transformation routines discussed in the previous section are all applied to whatever composite matrix is the current matrix, so it is important to use glMatrixMode to change to the modelview matrix before applying geometric transformations. Following this call, OpenGL transformation routines are used to modify the modelview matrix, which is then applied to transform coordinate positions in a scene. Two other modes that we can set with the glMatrixMode function are the texture mode and the color mode. The texture matrix is used for mapping texture patterns to surfaces, and the color matrix is used to convert from one color model to another. We discuss viewing, projection, texture, and color transformations in later chapters. For now, we limit our discussion to the details of the geometric transformations. The default argument for the glMatrixMode function is GL_MODELVIEW.

Once we are in the modelview mode (or any other mode), a call to a transformation routine generates a matrix that is multiplied by the current matrix for that mode. In addition, we can assign values to the elements of the current matrix, and there are two functions in the OpenGL library for this purpose. With the following function, we assign the identity matrix to the current matrix:

\[
\text{glLoadIdentity ( );}
\]

Alternatively, we can assign other values to the elements of the current matrix using

\[
\text{glLoadMatrix* (elements16);}
\]

A single-subscripted, 16-element array of floating-point values is specified with parameter elements16, and a suffix code of either f or d is used to designate the data type. The elements in this array must be specified in column-major order. That is, we first list the four elements in the first column, and then we list the four elements in the second column, the third column, and finally the fourth column. To illustrate this ordering, we initialize the modelview matrix with the following code:

\[
\text{glMatrixMode (GL_MODELVIEW);}
\]

\[
\text{GLfloat elems [16];}
\]

\[
\text{GLint k;}
\]

\[
\text{for (k = 0; k < 16; k++)}
\]

\[
\quad \text{elems [k] = float (k);}
\]

\[
\text{glLoadMatrixf (elems);}
\]
which produces the matrix

\[
M = \begin{bmatrix}
0.0 & 4.0 & 8.0 & 12.0 \\
1.0 & 5.0 & 9.0 & 13.0 \\
2.0 & 6.0 & 10.0 & 14.0 \\
3.0 & 7.0 & 11.0 & 15.0 \\
\end{bmatrix}
\]

We can also concatenate a specified matrix with the current matrix as follows:

\[
glMultMatrix\* \text{ (otherElements16);} \]

Again, the suffix code is either f or d, and parameter otherElements16 is a
16-element, single-subscripted array that lists the elements of some other matrix
in column-major order. The current matrix is postmultiplied by the matrix specified
in glMultMatrix, and this product replaces the current matrix. Thus, assuming
that the current matrix is the modelview matrix, which we designate as \( M \), then
the updated modelview matrix is computed as

\[
M = M \cdot M'
\]

where \( M' \) represents the matrix whose elements are specified by parameter
otherElements16 in the preceding glMultMatrix statement.

The glMultMatrix function can also be used to set up any transformation
sequence with individually defined matrices. For example,

\[
glMatrixMode \text{ (GL_MODELVIEW);} \\
glLoadIdentity \text{ ()}; \quad // \text{Set current matrix to the identity.} \\
glMultMatrixf \text{ (elemsM2);} \quad // \text{Postmultiply identity with matrix M2.} \\
glMultMatrixf \text{ (elemsM1);} \quad // \text{Postmultiply M2 with matrix M1.}
\]

produces the following current modelview matrix:

\[
M = M_2 \cdot M_1
\]

The first transformation to be applied in this sequence is the last one specified in
the code. Thus, if we set up a transformation sequence in an OpenGL program,
we can think of the individual transformations as being loaded onto a stack, so the
last operation specified is the first one applied. This is not what actually happens,
but the stack analogy may help you remember that in an OpenGL program, a
transformation sequence is applied in the opposite order from which it is specified.

It is also important to keep in mind that OpenGL stores matrices in column-
major order. In addition, a reference to a matrix element such as \( m_{jk} \) in OpenGL is
a reference to the element in column \( j \) and row \( k \). This is the reverse of the standard
mathematical convention, where the row number is referenced first. However, we
can usually avoid errors in row-column references by always specifying matrices
in OpenGL as 16-element, single-subscript arrays and listing the elements in a
column-major order.
OpenGL actually maintains a stack of composite matrices for each of the four modes that we can select with the glMatrixMode routine.

10 OpenGL Geometric-Transformation Programming Examples

In the following code segment, we apply each of the basic geometric transformations, one at a time, to a rectangle. Initially, the modelview matrix is the identity matrix and we display a blue rectangle. Next, we reset the current color to red, specify two-dimensional translation parameters, and display the red translated rectangle (Figure 34). Because we do not want to combine transformations, we next reset the current matrix to the identity. Then a rotation matrix is constructed and concatenated with the current matrix (the identity matrix). When the original rectangle is again referenced, it is rotated about the z axis and displayed in red (Figure 35). We repeat this process once more to generate the scaled and reflected rectangle shown in Figure 36.

```c
glMatrixMode (GL_MODELVIEW);

glColor3f (0.0, 0.0, 1.0);
glRecti (50, 100, 200, 150); // Display blue rectangle.

glColor3f (1.0, 0.0, 0.0);
glTranslatef (-200.0, -50.0, 0.0); // Set translation parameters.
glRecti (50, 100, 200, 150); // Display red, translated rectangle.

glLoadIdentity ( ); // Reset current matrix to identity.
glRotatef (90.0, 0.0, 0.0, 1.0); // Set 90-deg. rotation about z axis.
glRecti (50, 100, 200, 150); // Display red, rotated rectangle.

glLoadIdentity ( ); // Reset current matrix to identity.
glScalef (-0.5, 1.0, 1.0); // Set scale-reflection parameters.
glRecti (50, 100, 200, 150); // Display red, transformed rectangle.
```

**FIGURE 34**
Translating a rectangle using the OpenGL function glTranslatef
(−200.0, −50.0, 0.0).
The basic geometric transformations are translation, rotation, and scaling. Translation moves an object in a straight-line path from one position to another. Rotation moves an object from one position to another along a circular path around a specified rotation axis. For two-dimensional applications, the rotation path is in the $xy$ plane about an axis that is parallel to the $z$ axis. Scaling transformations change the dimensions of an object relative to a fixed position.

We can express two-dimensional transformations as $3 \times 3$ matrix operators, so that sequences of transformations can be concatenated into a single composite matrix. Performing geometric transformations with matrices is an efficient formulation because it allows us to reduce computations by applying a composite matrix to an object description to obtain its transformed position. To do this, we express coordinate positions as column matrices. We choose a column-matrix representation for coordinate points because this is the standard mathematical convention, and most graphics packages now follow this convention. A three-element column matrix (vector) is referred to as a homogeneous-coordinate representation. For geometric transformations, the homogeneous coefficient is assigned the value 1.

As with two-dimensional systems, transformations between three-dimensional Cartesian coordinate systems are accomplished with a sequence of translate-rotate transformations that brings the two systems into coincidence.
TABLE 1
Summary of OpenGL Geometric Transformation Functions

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glTranslate*</td>
<td>Specifies translation parameters.</td>
</tr>
<tr>
<td>glRotate*</td>
<td>Specifies parameters for rotation about any axis through the origin.</td>
</tr>
<tr>
<td>glScale*</td>
<td>Specifies scaling parameters with respect to coordinate origin.</td>
</tr>
<tr>
<td>glMatrixMode</td>
<td>Specifies current matrix for geometric-viewing transformations, projection transformations, texture transformations, or color transformations.</td>
</tr>
<tr>
<td>glLoadIdentity</td>
<td>Sets current matrix to identity.</td>
</tr>
<tr>
<td>glLoadMatrix* (elems);</td>
<td>Sets elements of current matrix.</td>
</tr>
<tr>
<td>glMultMatrix* (elems);</td>
<td>Postmultiplies the current matrix by the specified matrix.</td>
</tr>
<tr>
<td>glPixelZoom</td>
<td>Specifies two-dimensional scaling parameters for raster operations.</td>
</tr>
</tbody>
</table>

However, in a three-dimensional system, we must specify two of the three axis directions, not just one (as in a two-dimensional system).

The OpenGL basic library contains three functions for applying individual translate, rotate, and scale transformations to coordinate positions. Each function generates a matrix that is premultiplied by the modelview matrix. Thus, a sequence of geometric-transformation functions must be specified in reverse order: the last transformation invoked is the first to be applied to coordinate positions. Transformation matrices are applied to subsequently defined objects. In addition to accumulating transformation sequences in the modelview matrix, we can set this matrix to the identity or some other matrix. We can also form products with the modelview matrix and any specified matrices. Several operations are available in OpenGL for performing raster transformations. A block of pixels can be translated, rotated, scaled, or reflected with these OpenGL raster operations.

Table 1 summarizes the OpenGL geometric-transformation functions and matrix routines discussed in this chapter.

REFERENCES


Additional programming examples using OpenGL geometric-transformation functions are given in Woo, et al. (1999). Programming examples for the OpenGL geometric-transformation functions are also available at Nate Robins’s tutorial website: http://www.xmission.com/~nate/opengl.html. Finally, a complete listing of OpenGL geometric-transformation functions is provided in Shreiner (2000).
Two-Dimensional Geometric Transformations

EXERCISES
1. Write an animation program that implements the example two-dimensional rotation procedure of Section 1. An input polygon is to be rotated repeatedly in small steps around a pivot point in the $xy$ plane. Small angles are to be used for each successive step in the rotation, and approximations to the sine and cosine functions are to be used to speed up the calculations. To avoid excessive accumulation of round-off errors, reset the original coordinate values for the object at the start of each new revolution.

2. Show that the composition of two rotations is additive by concatenating the matrix representations for $R(\theta_1)$ and $R(\theta_2)$ to obtain
   \[ R(\theta_1) \cdot R(\theta_2) = R(\theta_1 + \theta_2) \]

3. Modify the two-dimensional transformation matrix (39), for scaling in an arbitrary direction, to include coordinates for any specified scaling fixed point $(x_f, y_f)$.

4. Prove that the multiplication of transformation matrices for each of the following sequences is commutative:
   (a) Two successive rotations.
   (b) Two successive translations.
   (c) Two successive scalings.

5. Prove that a uniform scaling and a rotation form a commutative pair of operations but that, in general, scaling and rotation are not commutative operations.

6. Multiple the individual scale, rotate, and translate matrices in Equation 42 to verify the elements in the composite transformation matrix.

7. Modify the example program in Section 4 so that transformation parameters can be specified as user input.

8. Modify the program from the previous exercise so that the transformation sequence can be applied to any polygon, with vertices specified as user input.

9. Modify the example program in Section 4 so that the order of the geometric transformation sequence can be specified as user input.

10. Show that transformation matrix (55), for a reflection about the line $y = x$, is equivalent to a reflection relative to the $x$ axis followed by a counterclockwise rotation of $90^\circ$.

11. Show that transformation matrix (56), for a reflection about the line $y = -x$, is equivalent to a reflection relative to the $y$ axis followed by a counterclockwise rotation of $90^\circ$.

12. Show that two successive reflections about either the $x$ axis or the $y$ axis is equivalent to a single rotation in the $xy$ plane about the coordinate origin.

13. Determine the form of the two-dimensional transformation matrix for a reflection about any line: $y = mx + b$.

14. Show that two successive reflections about any line in the $xy$ plane that intersects the coordinate origin is equivalent to a rotation in the $xy$ plane about the origin.

15. Determine a sequence of basic transformations that is equivalent to the $x$-direction shearing matrix (57).

16. Determine a sequence of basic transformations that is equivalent to the $y$-direction shearing matrix (61).

17. Set up a shearing procedure to display two-dimensional italic characters, given a vector font definition. That is, all character shapes in this font are defined with straight-line segments, and italic characters are formed with shearing transformations. Determine an appropriate value for the shear parameter by comparing italics and plain text in some available font. Define a simple vector font for input to your routine.

18. Derive the following equations for transforming a coordinate point $P = (x, y)$ in one two-dimensional Cartesian system to the coordinate values $(x', y')$ in another Cartesian system that is rotated counterclockwise by an angle $\theta$ relative to the first system. The transformation equations can be obtained by projecting point $P$ onto each of the four axes and analyzing the resulting right triangles.

   \[ x' = x \cos \theta + y \sin \theta \quad y' = -x \sin \theta + y \cos \theta \]

19. Write a procedure to compute the elements of the matrix for transforming object descriptions from one two-dimensional Cartesian coordinate system to another. The second coordinate system is to be defined with an origin point $P_0$ and a vector $V$ that gives the direction for the positive $y'$ axis of this system.

20. Set up procedures for implementing a block transfer of a rectangular area of a frame buffer, using one function to read the area into an array and another function to copy the array into the designated transfer area.

21. Determine the results of performing two successive block transfers into the same area of a frame buffer using the various Boolean operations.

22. What are the results of performing two successive block transfers into the same area of a frame buffer using the binary arithmetic operations?
23 Implement a routine to perform block transfers in a frame buffer using any specified Boolean operation or a replacement (copy) operation.

24 Write a routine to implement rotations in increments of 90° in frame-buffer block transfers.

25 Write a routine to implement rotations by any specified angle in a frame-buffer block transfer.

26 Write a routine to implement scaling as a raster transformation of a pixel block.

27 Write a program to display an animation of a black square on a white background tracing a circular, clockwise path around the display window with the path’s center at the display window’s center (like the tip of the minute hand on a clock). The orientation of the square should not change. Use only basic OpenGL geometric transformations to do this.

28 Repeat the previous exercise using OpenGL matrix operations.

29 Modify the program in Exercise 27 to have the square rotate clockwise about its own center as it moves along its path. The square should complete one revolution about its center for each quarter of its path around the window that it completes. Use only basic OpenGL geometric transformations to do this.

30 Repeat the previous exercise using OpenGL matrix operations.

31 Modify the program in Exercise 29 to have the square additionally “pulse” as it moves along its path. That is, for every revolution about its own center that it makes, it should go through one pulse cycle that begins with the square at full size, reduces smoothly in size down to 50% of normal size by the end of the cycle. Do this using only basic OpenGL geometric transformations.

32 Repeat the previous exercise using only OpenGL matrix operations.

IN MORE DEPTH

1 In this exercise, you’ll set up the routines necessary to make a crude animation of the objects in your application using two-dimensional geometric transformations. Decide on some simple motion behaviors for the objects in your application that can be achieved with the types of transformations discussed in this chapter (translations, rotations, scaling, shearing, and reflections). These behaviors may be motion patterns that certain objects will always be exhibiting, or they may be trajectories that are triggered or guided by user input (you can generate fixed example trajectories since we have not yet included user input). Set up the transformation matrices needed to produce these behaviors via composition of matrices. The matrices should be defined in homogeneous coordinates. If two or more objects act as a single “unit” in certain behaviors that are easier to model in terms of relative positions, you can use the techniques in Section 8 to convert the local transformations of the objects relative to each other (in their own coordinate frame) into transformations in the world coordinate frame.

2 Use the matrices you designed in the previous exercise to produce an animation of the behaviors of the objects in your scene. You should employ the OpenGL matrix operations and have the matrices produce small changes in position for each of the objects in the scene. The scene should then be redrawn several times per second to produce the animation, with the transformations being applied each time. Set the animation up so that it “loops”; that is, the behaviors should either be cyclical, or once the trajectories you designed for the objects have completed, the positions of all of the objects in the scene should be reset to their starting positions and the animation begun again.
We now examine in more detail the procedures for displaying views of a two-dimensional picture on an output device. Typically, a graphics package allows a user to specify which part of a defined picture is to be displayed and where that part is to be placed on the display device. Any convenient Cartesian coordinate system, referred to as the world-coordinate reference frame, can be used to define the picture. For a two-dimensional picture, a view is selected by specifying a region of the xy plane that contains the total picture or any part of it. A user can select a single area for display, or several areas could be selected for simultaneous display or for an animated panning sequence across a scene. The picture parts within the selected areas are then mapped onto specified areas of the device coordinates. When multiple view areas are selected, these areas can be placed in separate display locations, or some areas could be inserted into other, larger display areas.
Two-dimensional viewing transformations from world to device coordinates involve translation, rotation, and scaling operations, as well as procedures for deleting those parts of the picture that are outside the limits of a selected scene area.

1 The Two-Dimensional Viewing Pipeline

A section of a two-dimensional scene that is selected for display is called a **clipping window** because all parts of the scene outside the selected section are “clipped” off. The only part of the scene that shows up on the screen is what is inside the clipping window. Sometimes the clipping window is alluded to as the **world window** or the **viewing window**. And, at one time, graphics systems referred to the clipping window simply as “the window,” but there are now so many windows in use on computers that we need to distinguish between them. For example, a window-management system can create and manipulate several areas on a video screen, each of which is called “a window,” for the display of graphics and text. So we will always use the term **clipping window** to refer to a selected section of a scene that is eventually converted to pixel patterns within a display window on the video monitor. Graphics packages allow us also to control the placement within the display window using another “window” called the **viewport**. Objects inside the clipping window are mapped to the viewport, and it is the viewport that is then positioned within the display window. The clipping window selects what we want to see; the viewport indicates where it is to be viewed on the output device.

By changing the position of a viewport, we can view objects at different positions on the display area of an output device. Multiple viewports can be used to display different sections of a scene at different screen positions. Also, by varying the size of viewports, we can change the size and proportions of displayed objects. We achieve zooming effects by successively mapping different-sized clipping windows onto a fixed-size viewport. As the clipping windows are made smaller, we zoom in on some part of a scene to view details that are not shown with the larger clipping windows. Similarly, more overview is obtained by zooming out from a section of a scene with successively larger clipping windows. And panning effects are achieved by moving a fixed-size clipping window across the various objects in a scene.

Usually, clipping windows and viewports are rectangles in standard position, with the rectangle edges parallel to the coordinate axes. Other window or viewport geometries, such as general polygon shapes and circles, are used in some applications, but these shapes take longer to process. We first consider only rectangular viewports and clipping windows, as illustrated in Figure 1.
Two-Dimensional Viewing

The mapping of a two-dimensional, world-coordinate scene description to device coordinates is called a two-dimensional viewing transformation. Sometimes this transformation is simply referred to as the window-to-viewport transformation or the windowing transformation. But, in general, viewing involves more than just the transformation from clipping-window coordinates to viewport coordinates. In analogy with three-dimensional viewing, we can describe the steps for two-dimensional viewing as indicated in Figure 2. Once a world-coordinate scene has been constructed, we could set up a separate two-dimensional, viewing-coordinate reference frame for specifying the clipping window. But the clipping window is often just defined in world coordinates, so viewing coordinates for two-dimensional applications are the same as world coordinates. (For a three-dimensional scene, however, we need a separate viewing frame to specify the parameters for the viewing position, direction, and orientation.)

To make the viewing process independent of the requirements of any output device, graphics systems convert object descriptions to normalized coordinates and apply the clipping routines. Some systems use normalized coordinates in the range from 0 to 1, and others use a normalized range from \(-1\) to 1. Depending upon the graphics library in use, the viewport is defined either in normalized coordinates or in screen coordinates after the normalization process. At the final step of the viewing transformation, the contents of the viewport are transferred to positions within the display window.

Clipping is usually performed in normalized coordinates. This allows us to reduce computations by first concatenating the various transformation matrices. Clipping procedures are of fundamental importance in computer graphics. They are used not only in viewing transformations, but also in window-manager systems, in painting and drawing packages to erase picture sections, and in many other applications.

2 The Clipping Window

To achieve a particular viewing effect in an application program, we could design our own clipping window with any shape, size, and orientation we choose. For example, we might like to use a star pattern, an ellipse, or a figure with spline boundaries as a clipping window. But clipping a scene using a concave polygon or a clipping window with nonlinear boundaries requires more processing than clipping against a rectangle. We need to perform more computations to determine where an object intersects a circle than to find out where it intersects a straight line. The simplest window edges to clip against are straight lines that are parallel to the coordinate axes. Therefore, graphics packages commonly allow only rectangular clipping windows aligned with the \(x\) and \(y\) axes.

If we want some other shape for a clipping window, then we must implement our own clipping and coordinate-transformation algorithms, or we could just edit
the picture to produce a certain shape for the display frame around the scene. For example, we could trim the edges of a picture with any desired pattern by overlaying polygons that are filled with the background color. In this way, we could generate any desired border effects or even put interior holes in the picture.

Rectangular clipping windows in standard position are easily defined by giving the coordinates of two opposite corners of each rectangle. If we would like to get a rotated view of a scene, we could either define a rectangular clipping window in a rotated viewing-coordinate frame or, equivalently, we could rotate the world-coordinate scene. Some systems provide options for selecting a rotated, two-dimensional viewing frame, but usually the clipping window must be specified in world coordinates.

**Viewing-Corodinate Clipping Window**

A general approach to the two-dimensional viewing transformation is to set up a viewing-coordinate system within the world-coordinate frame. This viewing frame provides a reference for specifying a rectangular clipping window with any selected orientation and position, as in Figure 3. To obtain a view of the world-coordinate scene as determined by the clipping window of Figure 3, we just need to transfer the scene description to viewing coordinates. Although many graphics packages do not provide functions for specifying a clipping window in a two-dimensional viewing-coordinate system, this is the standard approach for defining a clipping region for a three-dimensional scene.

We choose an origin for a two-dimensional viewing-coordinate frame at some world position \( P_0 = (x_0, y_0) \), and we can establish the orientation using a world vector \( V \) that defines the \( y_{\text{view}} \) direction. Vector \( V \) is called the two-dimensional view up vector. An alternative method for specifying the orientation of the viewing frame is to give a rotation angle relative to either the \( x \) or \( y \) axis in the world frame. From this rotation angle, we can then obtain the view up vector. Once we have established the parameters that define the viewing-coordinate frame, we transform the scene description to the viewing system. This involves a sequence of transformations equivalent to superimposing the viewing frame on the world frame.

The first step in the transformation sequence is to translate the viewing origin to the world origin. Next, we rotate the viewing system to align it with the world frame. Given the orientation vector \( V \), we can calculate the components of unit vectors \( v = (v_x, v_y) \) and \( u = (u_x, u_y) \) for the \( y_{\text{view}} \) and \( x_{\text{view}} \) axes, respectively. These unit vectors are used to form the first and second rows of the rotation matrix \( R \) that aligns the viewing \( x_{\text{view}}, y_{\text{view}} \) axes with the world \( x_w, y_w \) axes.

Object positions in world coordinates are then converted to viewing coordinates with the composite two-dimensional transformation matrix

\[
M_{WC, VC} = R \cdot T
\]  

where \( T \) is the translation matrix that takes the viewing origin \( P_0 \) to the world origin, and \( R \) is the rotation matrix that rotates the viewing frame of reference into coincidence with the world-coordinate system. Figure 4 illustrates the steps in this coordinate transformation.

**World-Coordinate Clipping Window**

A routine for defining a standard, rectangular clipping window in world coordinates is typically provided in a graphics-programming library. We simply specify two world-coordinate positions, which are then assigned to the two opposite
corners of a standard rectangle. Once the clipping window has been established, the scene description is processed through the viewing routines to the output device.

If we want to obtain a rotated view of a two-dimensional scene, as discussed in the previous section, we perform exactly the same steps as described there, but without considering a viewing frame of reference. Thus, we simply rotate (and possibly translate) objects to a desired position and set up the clipping window—all in world coordinates. For example, we could display a rotated view of the triangle in Figure 5(a) by rotating it into the position we want and setting up a standard clipping rectangle. In analogy with the coordinate transformation described in the previous section, we could also translate the triangle to the world origin and define a clipping window around the triangle. In that case, we define an orientation vector and choose a reference point such as the triangle’s centroid. Then we translate the reference point to the world origin and rotate the orientation vector onto the y_world axis using transformation matrix 1. With the triangle in the desired orientation, we can use a standard clipping window in world coordinates to capture the view of the rotated triangle. The transformed position of the triangle and the selected clipping window are shown in Figure 5(b).

### 3 Normalization and Viewport Transformations

With some graphics packages, the normalization and window-to-viewport transformations are combined into one operation. In this case, the viewport coordinates are often given in the range from 0 to 1 so that the viewport is positioned within a unit square. After clipping, the unit square containing the viewport is mapped to the output display device. In other systems, the normalization and clipping routines are applied before the viewport transformation. For these systems, the viewport boundaries are specified in screen coordinates relative to the display-window position.

**Mapping the Clipping Window into a Normalized Viewport**

To illustrate the general procedures for the normalization and viewport transformations, we first consider a viewport defined with normalized coordinate values between 0 and 1. Object descriptions are transferred to this normalized space using a transformation that maintains the same relative placement of a point in...
the viewport as it had in the clipping window. If a coordinate position is at the center of the clipping window, for instance, it would be mapped to the center of the viewport. Figure 6 illustrates this window-to-viewport mapping. Position \((x_w, y_w)\) in the clipping window is mapped to position \((x_v, y_v)\) in the associated viewport.

To transform the world-coordinate point into the same relative position within the viewport, we require that

\[
\begin{align*}
x_v - x_{v\min} &= \frac{x_w - x_{w\min}}{x_{w\max} - x_{w\min}} \\
y_v - y_{v\min} &= \frac{y_w - y_{w\min}}{y_{w\max} - y_{w\min}}
\end{align*}
\]

Solving these expressions for the viewport position \((x_v, y_v)\), we have

\[
\begin{align*}
x_v &= s_x x_w + t_x \\
y_v &= s_y y_w + t_y
\end{align*}
\]

where the scaling factors are

\[
\begin{align*}
s_x &= \frac{x_{v\max} - x_{v\min}}{x_{w\max} - x_{w\min}} \\
s_y &= \frac{y_{v\max} - y_{v\min}}{y_{w\max} - y_{w\min}}
\end{align*}
\]

and the translation factors are

\[
\begin{align*}
t_x &= \frac{x_{w\max} x_{v\min} - x_{w\min} x_{v\max}}{x_{w\max} - x_{w\min}} \\
t_y &= \frac{y_{w\max} y_{v\min} - y_{w\min} y_{v\max}}{y_{w\max} - y_{w\min}}
\end{align*}
\]

Because we are simply mapping world-coordinate positions into a viewport that is positioned near the world origin, we can also derive Equations 3 using any transformation sequence that converts the rectangle for the clipping window into the viewport rectangle. For example, we could obtain the transformation from world coordinates to viewport coordinates with the following sequence:

1. Scale the clipping window to the size of the viewport using a fixed-point position of \((x_{w\min}, y_{w\min})\).
2. Translate \((x_{w\min}, y_{w\min})\) to \((x_{v\min}, y_{v\min})\).
The scaling transformation in step (1) can be represented with the two-dimensional matrix

\[
S = \begin{bmatrix}
s_x & 0 & x_{w\min}(1 - s_x) \\
0 & s_y & y_{w\min}(1 - s_y) \\
0 & 0 & 1
\end{bmatrix}
\]  \( (6) \)

where \( s_x \) and \( s_y \) are the same as in Equations 4. The two-dimensional matrix representation for the translation of the lower-left corner of the clipping window to the lower-left viewport corner is

\[
T = \begin{bmatrix}
1 & 0 & x_{v\min} - x_{w\min} \\
0 & 1 & y_{v\min} - y_{w\min} \\
0 & 0 & 1
\end{bmatrix}
\]  \( (7) \)

And the composite matrix representation for the transformation to the normalized viewport is

\[
M_{\text{window, normviewport}} = T \cdot S = \begin{bmatrix}
s_x & 0 & t_x \\
0 & s_y & t_y \\
0 & 0 & 1
\end{bmatrix}
\]  \( (8) \)

which gives us the same result as in Equations 3. Any other clipping-window reference point, such as the top-right corner or the window center, could be used for the scale–translate operations. Alternatively, we could first translate any clipping-window position to the corresponding location in the viewport, and then scale relative to that viewport location.

The window-to-viewport transformation maintains the relative placement of object descriptions. An object inside the clipping window is mapped to a corresponding position inside the viewport. Similarly, an object outside the clipping window is outside the viewport.

Relative proportions of objects, on the other hand, are maintained only if the aspect ratio of the viewport is the same as the aspect ratio of the clipping window. In other words, we keep the same object proportions if the scaling factors \( s_x \) and \( s_y \) are the same. Otherwise, world objects will be stretched or contracted in either the \( x \) or \( y \) directions (or both) when displayed on the output device.

The clipping routines can be applied using either the clipping-window boundaries or the viewport boundaries. After clipping, the normalized coordinates are transformed into device coordinates. And the unit square can be mapped onto the output device using the same procedures as in the window-to-viewport transformation, with the area inside the unit square transferred to the total display area of the output device.

### Mapping the Clipping Window into a Normalized Square

Another approach to two-dimensional viewing is to transform the clipping window into a normalized square, clip in normalized coordinates, and then transfer the scene description to a viewport specified in screen coordinates. This transformation is illustrated in Figure 7 with normalized coordinates in the range from \(-1\) to \(1\). The clipping algorithms in this transformation sequence are now standardized so that objects outside the boundaries \( x = \pm 1 \) and \( y = \pm 1 \) are detected and removed from the scene description. At the final step of the viewing transformation, the objects in the viewport are positioned within the display window.

We transfer the contents of the clipping window into the normalization square using the same procedures as in the window-to-viewport transformation. The matrix for the normalization transformation is obtained from Equation 8 by substituting \(-1\) for \( x_{v\min} \) and \( y_{v\min} \) and substituting \(+1\) for \( x_{v\max} \) and \( y_{v\max} \).
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**Figure 7**
A point \((x_w, y_w)\) in a clipping window is mapped to a normalized coordinate position \((x_{\text{norm}}, y_{\text{norm}})\), then to a screen-coordinate position \((x_v, y_v)\) in a viewport. Objects are clipped against the normalization square before the transformation to viewport coordinates occurs.

Making these substitutions in the expressions for \(t_x, t_y, s_x,\) and \(s_y\), we have

\[
M_{\text{window, normsquare}} = \begin{bmatrix}
\frac{2}{x_{\text{w}}^{\max} - x_{\text{w}}^{\min}} & 0 & -\frac{x_{\text{w}}^{\max} + x_{\text{w}}^{\min}}{x_{\text{w}}^{\max} - x_{\text{w}}^{\min}} \\
0 & \frac{2}{y_{\text{w}}^{\max} - y_{\text{w}}^{\min}} & -\frac{y_{\text{w}}^{\max} + y_{\text{w}}^{\min}}{y_{\text{w}}^{\max} - y_{\text{w}}^{\min}} \\
0 & 0 & 1
\end{bmatrix}
\]

(9)

Similarly, after the clipping algorithms have been applied, the normalized square with edge length equal to 2 is transformed into a specified viewport. This time, we get the transformation matrix from Equation 8 by substituting \(-1\) for \(x_{\text{w}}^{\min}\) and \(y_{\text{w}}^{\min}\) and substituting \(+1\) for \(x_{\text{w}}^{\max}\) and \(y_{\text{w}}^{\max}\):

\[
M_{\text{normsquare, viewport}} = \begin{bmatrix}
\frac{2}{x_{\text{v}}^{\max} - x_{\text{v}}^{\min}} & 0 & \frac{x_{\text{v}}^{\max} + x_{\text{v}}^{\min}}{x_{\text{v}}^{\max} - x_{\text{v}}^{\min}} \\
0 & \frac{2}{y_{\text{v}}^{\max} - y_{\text{v}}^{\min}} & \frac{y_{\text{v}}^{\max} + y_{\text{v}}^{\min}}{y_{\text{v}}^{\max} - y_{\text{v}}^{\min}} \\
0 & 0 & 1
\end{bmatrix}
\]

(10)

The last step in the viewing process is to position the viewport area in the display window. Typically, the lower-left corner of the viewport is placed at a coordinate position specified relative to the lower-left corner of the display window. Figure 8 demonstrates the positioning of a viewport within a display window.

As before, we maintain the initial proportions of objects by choosing the aspect ratio of the viewport to be the same as the clipping window. Otherwise, objects

**Figure 8**
A viewport at coordinate position \((x_s, y_s)\) within a display window.
will be stretched or contracted in the $x$ or $y$ directions. Also, the aspect ratio of the display window can affect the proportions of objects. If the viewport is mapped to the entire area of the display window and the size of the display window is changed, objects may be distorted unless the aspect ratio of the viewport is also adjusted.

**Display of Character Strings**

Character strings can be handled in one of two ways when they are mapped through the viewing pipeline to a viewport. The simplest mapping maintains a constant character size. This method could be employed with bitmap character patterns. But outline fonts could be transformed the same as other primitives; we just need to transform the defining positions for the line segments in the outline character shapes. Algorithms for determining the pixel patterns for the transformed characters are then applied when the other primitives in the scene are processed.

**Split-Screen Effects and Multiple Output Devices**

By selecting different clipping windows and associated viewports for a scene, we can provide simultaneous display of two or more objects, multiple picture parts, or different views of a single scene. And we can position these views in different parts of a single display window or in multiple display windows on the screen. In a design application, for example, we can display a wire-frame view of an object in one viewport while also displaying a fully rendered view of the object in another viewport. In addition, we could list other information or menus in a third viewport.

It is also possible that two or more output devices could be operating concurrently on a particular system, and we can set up a clipping-window/viewport pair for each output device. A mapping to a selected output device is sometimes referred to as a **workstation transformation**. In this case, viewports could be specified in the coordinates of a particular display device, or each viewport could be specified within a unit square, which is then mapped to a chosen output device. Some graphics systems provide a pair of workstation functions for this purpose. One function is used to designate a clipping window for a selected output device, identified by a *workstation number*, and the other function is used to set the associated viewport for that device.

## 4 OpenGL Two-Dimensional Viewing Functions

Actually, the basic OpenGL library has no functions specifically for two-dimensional viewing because it is designed primarily for three-dimensional applications. But we can adapt the three-dimensional viewing routines to a two-dimensional scene, and the core library contains a viewport function. In addition, the GLU library provides a function for specifying a two-dimensional clipping window, and we have GLUT library functions for handling display windows. Therefore, we can use these two-dimensional routines, along with the OpenGL viewport function, for all the viewing operations we need.

**OpenGL Projection Mode**

Before we select a clipping window and a viewport in OpenGL, we need to establish the appropriate mode for constructing the matrix to transform from world coordinates to screen coordinates. With OpenGL, we cannot set up a
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separate two-dimensional viewing-coordinate system as in Figure 3, and we must set the parameters for the clipping window as part of the projection transformation. Therefore, we must first select the projection mode. We do this with the same function we used to set the modelview mode for the geometric transformations. Subsequent commands for defining a clipping window and viewport will then be applied to the projection matrix.

\[ \text{glMatrixMode(GL_PROJECTION);} \]

This designates the projection matrix as the current matrix, which is originally set to the identity matrix. However, if we are going to loop back through this statement for other views of a scene, we can also set the initialization as

\[ \text{glLoadIdentity();} \]

This ensures that each time we enter the projection mode, the matrix will be reset to the identity matrix so that the new viewing parameters are not combined with the previous ones.

**GLU Clipping-Window Function**

To define a two-dimensional clipping window, we can use the GLU function:

\[ \text{gluOrtho2D(xwmin, xwmax, ywmin, ywmax);} \]

Coordinate positions for the clipping-window boundaries are given as double-precision numbers. This function specifies an orthogonal projection for mapping the scene to the screen. For a three-dimensional scene, this means that objects would be projected along parallel lines that are perpendicular to the two-dimensional xy display screen. But for a two-dimensional application, objects are already defined in the xy plane. Therefore, the orthogonal projection has no effect on our two-dimensional scene other than to convert object positions to normalized coordinates. Nevertheless, we must specify the orthogonal projection because our two-dimensional scene is processed through the full three-dimensional OpenGL viewing pipeline. In fact, we could specify the clipping window using the three-dimensional OpenGL core-library version of the \text{gluOrtho2D} function.

Normalized coordinates in the range from −1 to 1 are used in the OpenGL clipping routines. And the \text{gluOrtho2D} function sets up a three-dimensional version of transformation matrix 9 for mapping objects within the clipping window to normalized coordinates. Objects outside the normalized square (and outside the clipping window) are eliminated from the scene to be displayed.

If we do not specify a clipping window in an application program, the default coordinates are \((x_{\text{wmin}}, y_{\text{wmin}}) = (-1.0, -1.0)\) and \((x_{\text{wmax}}, y_{\text{wmax}}) = (1.0, 1.0)\). Thus the default clipping window is the normalized square centered on the coordinate origin with a side length of 2.0.

**OpenGL Viewport Function**

We specify the viewport parameters with the OpenGL function

\[ \text{glViewport(xvmin, yvmin, vpWidth, vpHeight);} \]

where all parameter values are given in integer screen coordinates relative to the display window. Parameters \(x_{\text{vmin}}\) and \(y_{\text{vmin}}\) specify the position of the lower-left corner of the viewport relative to the lower-left corner of the display window, and the pixel width and height of the viewport are set with parameters \(vp\text{Width}\) and \(vp\text{Height}\).
and vpHeight. If we do not invoke the glViewport function in a program, the default viewport size and position are the same as the size and position of the display window.

After the clipping routines have been applied, positions within the normalized square are transformed into the viewport rectangle using Matrix 10. Coordinates for the upper-right corner of the viewport are calculated for this transformation matrix in terms of the viewport width and height:

\[
x_{\text{vmax}} = x_{\text{vmin}} + \text{vpWidth}, \quad y_{\text{vmax}} = y_{\text{vmin}} + \text{vpHeight}
\]  

(11)

For the final transformation, pixel colors for the primitives within the viewport are loaded into the refresh buffer at the specified screen locations.

Multiple viewports can be created in OpenGL for a variety of applications (see Section 3). We can obtain the parameters for the currently active viewport using the query function

\[
\text{glGetInteger}v \left( \text{GL\_VIEWPORT}, \text{vpArray} \right);
\]

where vpArray is a single-subscript, four-element array. This Get function returns the parameters for the current viewport to vpArray in the order xvmin, yvmin, vpWidth, and vpHeight. In an interactive application, for example, we can use this function to obtain parameters for the viewport that contains the screen cursor.

Creating a GLUT Display Window

Because the GLUT library interfaces with any window-management system, we use the GLUT routines for creating and manipulating display windows so that our example programs will be independent of any specific machine. To access these routines, we first need to initialize GLUT with the following function:

\[
\text{glutInit} \left( \&\text{argc}, \text{argv} \right);
\]

Parameters for this initialization function are the same as those for the main procedure, and we can use glutInit to process command-line arguments.

We have three functions in GLUT for defining a display window and choosing its dimensions and position:

\[
\begin{align*}
\text{glutInitWindowPosition} \left( \text{xTopLeft}, \text{yTopLeft} \right); \\
\text{glutInitWindowSize} \left( \text{dwWidth}, \text{dwHeight} \right); \\
\text{glutCreateWindow} \left( \text{"Title of Display Window"} \right);
\end{align*}
\]

The first of these functions gives the integer, screen-coordinate position for the top-left corner of the display window, relative to the top-left corner of the screen. If either coordinate is negative, the display-window position on the screen is determined by the window-management system. With the second function, we choose a width and height for the display window in positive integer pixel dimensions. If we do not use these two functions to specify a size and position, the default size is 300 by 300 and the default position is \((-1, -1)\), which leaves the positioning of the display window to the window-management system. In any case, the display-window size and position specified with GLUT routines might be ignored, depending on the state of the window-management system or the other requirements currently in effect for it. Thus, the window system might position and size the display window differently. The third function creates the display window, with the specified size and position, and assigns a title, although the use
Setting the GLUT Display-Window Mode and Color

Various display-window parameters are selected with the GLUT function

```c
glutInitDisplayMode (mode);
```

We use this function to choose a color mode (RGB or index) and different buffer combinations, and the selected parameters are combined with the logical or operation. The default mode is single buffering and the RGB (or RGBA) color mode, which is the same as setting this mode with the statement

```c
glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
```

The color mode specification GLUT_RGB is equivalent to GLUT_RGBA. A background color for the display window is chosen in RGB mode with the OpenGL routine

```c
glClearColor (red, green, blue, alpha);
```

In color-index mode, we set the display-window color with

```c
glClearIndex (index);
```

where parameter `index` is assigned an integer value corresponding to a position within the color table.

GLUT Display-Window Identifier

Multiple display windows can be created for an application, and each is assigned a positive-integer `display-window identifier`, starting with the value 1 for the first window that is created. At the time that we initiate a display window, we can record its identifier with the statement

```c
windowID = glutCreateWindow ("A Display Window");
```

Once we have saved the integer display-window identifier in variable name `windowID`, we can use the identifier number to change display parameters or to delete the display window.

Deleting a GLUT Display Window

The GLUT library also includes a function for deleting a display window that we have created. If we know the display window’s identifier, we can eliminate it with the statement

```c
glutDestroyWindow (windowID);
```

Current GLUT Display Window

When we specify any display-window operation, it is applied to the `current display window`, which is either the last display window that we created or the one we select with the following command:

```c
glutSetWindow (windowID);
```
In addition, at any time, we can query the system to determine which window is the current display window:

```c
currentWindowID = glutGetWindow();
```

A value of 0 is returned by this function if there are no display windows or if the current display window was destroyed.

**Relocating and Resizing a GLUT Display Window**

We can reset the screen location for the current display window with

```c
glutPositionWindow (xNewTopLeft, yNewTopLeft);
```

where the coordinates specify the new position for the upper-left display-window corner, relative to the upper-left corner of the screen. Similarly, the following function resets the size of the current display window:

```c
glutReshapeWindow (dwNewWidth, dwNewHeight);
```

With the following command, we can expand the current display window to fill the screen:

```c
glutFullScreen ( );
```

The exact size of the display window after execution of this routine depends on the window-management system. A subsequent call to either `glutPositionWindow` or `glutReshapeWindow` will cancel the request for an expansion to full-screen size.

Whenever the size of a display window is changed, its aspect ratio may change and objects may be distorted from their original shapes. We can adjust for a change in display-window dimensions using the statement

```c
glutReshapeFunc (winReshapeFcn);
```

This GLUT routine is activated when the size of a display window is changed, and the new width and height are passed to its argument: the function `winReshapeFcn`, in this example. Thus, `winReshapeFcn` is the “callback function” for the “reshape event.” We can then use this callback function to change the parameters for the viewport so that the original aspect ratio of the scene is maintained. In addition, we could also reset the clipping-window boundaries, change the display-window color, adjust other viewing parameters, and perform any other tasks.

**Managing Multiple GLUT Display Windows**

The GLUT library also has a number of routines for manipulating a display window in various ways. These routines are particularly useful when we have multiple display windows on the screen and we want to rearrange them or locate a particular display window.

We use the following routine to convert the current display window to an icon in the form of a small picture or symbol representing the window:

```c
glutIconifyWindow ( );
```

The label on this icon will be the same name that we assigned to the window, but we can change this with the following command:

```c
glutSetIconTitle ("Icon Name");
```
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We also can change the name of the display window with a similar command:

```c
    glutSetWindowTitle ("New Window Name");
```

With multiple display windows open on the screen, some windows may overlap or totally obscure other display windows. We can choose any display window to be in front of all other windows by first designating it as the current window, and then issuing the “pop-window” command:

```c
    glutSetWindow (windowID);
    glutPopWindow ();
```

In a similar way, we can “push” the current display window to the back so that it is behind all other display windows. This sequence of operations is

```c
    glutSetWindow (windowID);
    glutPushWindow ();
```

We can also take the current window off the screen with

```c
    glutHideWindow ();
```

In addition, we can return a “hidden” display window, or one that has been converted to an icon, by designating it as the current display window and then invoking the function

```c
    glutShowWindow ();
```

### GLUT Subwindows

Within a selected display window, we can set up any number of second-level display windows, which are called subwindows. This provides a means for partitioning display windows into different display sections. We create a subwindow with the following function:

```c
    glutCreateSubWindow (windowID, xBottomLeft, yBottomLeft,
        width, height);
```

Parameter `windowID` identifies the display window in which we want to set up the subwindow. With the remaining parameters, we specify the subwindow’s size and the placement of its lower-left corner relative to the lower-left corner of the display window.

Subwindows are assigned a positive integer identifier in the same way that first-level display windows are numbered, and we can place a subwindow inside another subwindow. Also, each subwindow can be assigned an individual display mode and other parameters. We can even reshape, reposition, push, pop, hide, and show subwindows, just as we can with first-level display windows. But we cannot convert a GLUT subwindow to an icon.

### Selecting a Display-Window Screen-Cursor Shape

We can use the following GLUT routine to request a shape for the screen cursor that is to be used with the current window:

```c
    glutSetCursor (shape);
```
The possible cursor shapes that we can select are an arrow pointing in a chosen direction, a bidirectional arrow, a rotating arrow, a crosshair, a wristwatch, a question mark, or even a skull and crossbones. For example, we can assign the symbolic constant `GLUT_CURSOR_UP_DOWN` to parameter `shape` to obtain an up-down arrow. A rotating arrow is chosen with `GLUT_CURSOR_CYCLE`, a wristwatch shape is selected with `GLUT_CURSOR_WAIT`, and a skull and crossbones is obtained with the constant `GLUT_CURSOR_DESTROY`. A cursor shape can be assigned to a display window to indicate a particular kind of application, such as an animation. However, the exact shapes that we can use are system dependent.

**Viewing Graphics Objects in a GLUT Display Window**

After we have created a display window and selected its position, size, color, and other characteristics, we indicate what is to be shown in that window. If more than one display window has been created, we first designate the one we want as the current display window. Then we invoke the following function to assign something to that window:

```c
glutDisplayFunc (pictureDescrip);
```

The argument is a routine that describes what is to be displayed in the current window. This routine, called `pictureDescrip` for this example, is referred to as a *callback function* because it is the routine that is to be executed whenever GLUT determines that the display-window contents should be renewed. Routine `pictureDescrip` usually contains the OpenGL primitives and attributes that define a picture, although it could specify other constructs such as a menu display.

If we have set up multiple display windows, then we repeat this process for each of the display windows or subwindows. Also, we may need to call `glutDisplayFunc` after the `glutPopWindow` command if the display window has been damaged during the process of redisplaying the windows. In this case, the following function is used to indicate that the contents of the current display window should be renewed:

```c
glutPostRedisplay ( );
```

This routine is also used when an additional object, such as a pop-up menu, is to be shown in a display window.

**Executing the Application Program**

When the program setup is complete and the display windows have been created and initialized, we need to issue the final GLUT command that signals execution of the program:

```c
glutMainLoop ( );
```

At this time, display windows and their graphic contents are sent to the screen. The program also enters the GLUT *processing loop* that continually checks for new "events," such as interactive input from a mouse or a graphics tablet.
Other GLUT Functions

The GLUT library provides a wide variety of routines to handle processes that are system dependent and to add features to the basic OpenGL library. For example, this library contains functions for generating bitmap and outline characters, and it provides functions for loading values into a color table. In addition, some GLUT functions are available for displaying three-dimensional objects, either as solids or in a wireframe representation. These objects include a sphere, a torus, and the five regular polyhedra (cube, tetrahedron, octahedron, dodecahedron, and icosahedron).

Sometimes it is convenient to designate a function that is to be executed when there are no other events for the system to process. We can do that with

```c
    glutIdleFunc (function);
```

The parameter for this GLUT routine could reference a background function or a procedure to update parameters for an animation when no other processes are taking place.

We also have GLUT functions for obtaining and processing interactive input and for creating and managing menus. Individual routines are provided by GLUT for input devices such as a mouse, keyboard, graphics tablet, and spaceball.

Finally, we can use the following function to query the system about some of the current state parameters:

```c
    glutGet (stateParam);
```

This function returns an integer value corresponding to the symbolic constant we select for its argument. For example, we can obtain the x-coordinate position for the top-left corner of the current display window, relative to the top-left corner of the screen, with the constant GLUT_WINDOW_X; and we can retrieve the current display-window width or the screen width with GLUT_WINDOW_WIDTH or GLUT_SCREEN_WIDTH.

OpenGL Two-Dimensional Viewing Program Example

As a demonstration of the use of the OpenGL viewport function, we use a split-screen effect to show two views of a triangle in the xy plane with its centroid at the world-coordinate origin. First, a viewport is defined in the left half of the display window, and the original triangle is displayed there in blue. Using the same clipping window, we then define another viewport for the right half of the display window, and the fill color is changed to red. The triangle is then rotated about its centroid and displayed in the second viewport.

```c
#include <GL/glut.h>

class wcPt2D {
  public:
    GLfloat x, y;
};
```
```c
#include <GL/glut.h>

void init (void)
{
    /* Set color of display window to white. */
    glClearColor (1.0, 1.0, 1.0, 0.0);

    /* Set parameters for world-coordinate clipping window. */
    glMatrixMode (GL_PROJECTION);
    gluOrtho2D (-100.0, 100.0, -100.0, 100.0);

    /* Set mode for constructing geometric transformation matrix. */
    glMatrixMode (GL_MODELVIEW);
}

void triangle (wcPt2D *verts)
{
    GLint k;

    glBegin (GL_TRIANGLES);
    for (k = 0; k < 3; k++)
        glVertex2f (verts [k].x, verts [k].y);
    glEnd ( );
}

void displayFcn (void)
{
    /* Define initial position for triangle. */
    wcPt2D verts [3] = { {-50.0, -25.0}, {50.0, -25.0}, {0.0, 50.0} };

    glClear (GL_COLOR_BUFFER_BIT); // Clear display window.
    glClearColor (0.0, 0.0, 1.0, 0.0); // Set fill color to blue.
    glViewport (0, 0, 300, 300); // Set left viewport.
    triangle (verts); // Display triangle.

    /* Rotate triangle and display in right half of display window. */
    glClearColor (1.0, 0.0, 0.0, 0.0); // Set fill color to red.
    glViewport (300, 0, 300, 300); // Set right viewport.
    glRotatef (90.0, 0.0, 0.0, 1.0); // Rotate about z axis.
    triangle (verts); // Display red rotated triangle.
}

void main (int argc, char ** argv)
{
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowSize (600, 300);
    glutCreateWindow ("Split-Screen Example");

    init ( );
    glutDisplayFunc (displayFcn);

    glutMainLoop ( );
}
```

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5 Clipping Algorithms

Generally, any procedure that eliminates those portions of a picture that are either inside or outside a specified region of space is referred to as a clipping algorithm or simply clipping. Usually a clipping region is a rectangle in standard position, although we could use any shape for a clipping application.

The most common application of clipping is in the viewing pipeline, where clipping is applied to extract a designated portion of a scene (either two-dimensional or three-dimensional) for display on an output device. Clipping methods are also used to antialias object boundaries, to construct objects using solid-modeling methods, to manage a multiwindow environment, and to allow parts of a picture to be moved, copied, or erased in drawing and painting programs.

Clipping algorithms are applied in two-dimensional viewing procedures to identify those parts of a picture that are within the clipping window. Everything outside the clipping window is then eliminated from the scene description that is transferred to the output device for display. An efficient implementation of clipping in the viewing pipeline is to apply the algorithms to the normalized boundaries of the clipping window. This reduces calculations, because all geometric and viewing transformation matrices can be concatenated and applied to a scene description before clipping is carried out. The clipped scene can then be transferred to screen coordinates for final processing.

In the following sections, we explore two-dimensional algorithms for

- Point clipping
- Line clipping (straight-line segments)
- Fill-area clipping (polygons)
- Curve clipping
- Text clipping

Point, line, and polygon clipping are standard components of graphics packages. But similar methods can be applied to other objects, particularly conics, such as circles, ellipses, and spheres, in addition to spline curves and surfaces. Usually, however, objects with nonlinear boundaries are approximated with straight-line segments or polygon surfaces to reduce computations.

Unless otherwise stated, we assume that the clipping region is a rectangular window in standard position, with boundary edges at coordinate positions \( x_{\text{w min}}, x_{\text{w max}}, y_{\text{w min}}, \text{ and } y_{\text{w max}} \). These boundary edges typically correspond to a normalized square, in which the \( x \) and \( y \) values range either from 0 to 1 or from \(-1\) to 1.

6 Two-Dimensional Point Clipping

For a clipping rectangle in standard position, we save a two-dimensional point \( P = (x, y) \) for display if the following inequalities are satisfied:

\[
\begin{align*}
x_{\text{w min}} & \leq x \leq x_{\text{w max}} \\
y_{\text{w min}} & \leq y \leq y_{\text{w max}}
\end{align*}
\]  

(12)

If any of these four inequalities is not satisfied, the point is clipped (not saved for display).

Although point clipping is applied less often than line or polygon clipping, it is useful in various situations, particularly when pictures are modeled with particle systems. For example, point clipping can be applied to scenes involving
clouds, sea foam, smoke, or explosions that are modeled with “particles,” such as the center coordinates for small circles or spheres.

7 Two-Dimensional Line Clipping

Figure 9 illustrates possible positions for straight-line segments in relationship to a standard clipping window. A line-clipping algorithm processes each line in a scene through a series of tests and intersection calculations to determine whether the entire line or any part of it is to be saved. The expensive part of a line-clipping procedure is in calculating the intersection positions of a line with the window edges. Therefore, a major goal for any line-clipping algorithm is to minimize the intersection calculations. To do this, we can first perform tests to determine whether a line segment is completely inside the clipping window or completely outside. It is easy to determine whether a line is completely inside a clipping window, but it is more difficult to identify all lines that are entirely outside the window. If we are unable to identify a line as completely inside or completely outside a clipping rectangle, we must then perform intersection calculations to determine whether any part of the line crosses the window interior.

We test a line segment to determine if it is completely inside or outside a selected clipping-window edge by applying the point-clipping tests of the previous section. When both endpoints of a line segment are inside all four clipping boundaries, such as the line from \( P_1 \) to \( P_2 \) in Figure 9, the line is completely inside the clipping window and we save it. And when both endpoints of a line segment are outside any one of the four boundaries (as with line \( P_3P_4 \) in Figure 9), that line is completely outside the window and it is eliminated from the scene description. But if both these tests fail, the line segment intersects at least one clipping boundary and it may or may not cross into the interior of the clipping window.

One way to formulate the equation for a straight-line segment is to use the following parametric representation, where the coordinate positions \((x_0, y_0)\) and \((x_{\text{end}}, y_{\text{end}})\) designate the two line endpoints:

\[
\begin{align*}
    x &= x_0 + u(x_{\text{end}} - x_0) \\
    y &= y_0 + u(y_{\text{end}} - y_0) \\
    0 &\leq u \leq 1
\end{align*}
\]  

We can use this parametric representation to determine where a line segment crosses each clipping-window edge by assigning the coordinate value for that edge to either \( x \) or \( y \) and solving for parameter \( u \). For example, the left window boundary is at position \( x w_{\text{min}} \), so we substitute this value for \( x \), solve for \( u \), and calculate the corresponding \( y \)-intersection value. If this value of \( u \) is outside the range from 0 to 1, the line segment does not intersect that window border line.

![Figure 9](image-url)  
**Clipping straight-line segments using a standard rectangular clipping window.**
However, if the value of \( u \) is within the range from 0 to 1, part of the line is inside that border. We can then process this inside portion of the line segment against the other clipping boundaries until either we have clipped the entire line or we find a section that is inside the window.

Processing line segments in a scene using the simple clipping approach described in the preceding paragraph is straightforward, but not very efficient. It is possible to reformulate the initial testing and the intersection calculations to reduce processing time for a set of line segments, and a number of faster line clippers have been developed. Some of the algorithms are designed explicitly for two-dimensional pictures and some are easily adapted to sets of three-dimensional line segments.

### Cohen-Sutherland Line Clipping

This is one of the earliest algorithms to be developed for fast line clipping, and variations of this method are widely used. Processing time is reduced in the Cohen-Sutherland method by performing more tests before proceeding to the intersection calculations. Initially, every line endpoint in a picture is assigned a four-digit binary value, called a **region code**, and each bit position is used to indicate whether the point is inside or outside one of the clipping-window boundaries. We can reference the window edges in any order, and Figure 10 illustrates one possible ordering with the bit positions numbered 1 through 4 from right to left. Thus, for this ordering, the rightmost position (bit 1) references the left clipping-window boundary, and the leftmost position (bit 4) references the top window boundary. A value of 1 (or *true*) in any bit position indicates that the endpoint is outside that window border. Similarly, a value of 0 (or *false*) in any bit position indicates that the endpoint is not outside (it is inside or on) the corresponding window edge. Sometimes, a region code is referred to as an **"out" code** because a value of 1 in any bit position indicates that the spatial point is outside the corresponding clipping boundary.

Each clipping-window edge divides two-dimensional space into an inside half space and an outside half space. Together, the four window borders create nine regions, and Figure 11 lists the value for the binary code in each of these regions. Thus, an endpoint that is below and to the left of the clipping window is assigned the region code 0101, and the region-code value for any endpoint inside the clipping window is 0000.

Bit values in a region code are determined by comparing the coordinate values \((x, y)\) of an endpoint to the clipping boundaries. Bit 1 is set to 1 if \( x < x_{\text{min}} \), and
the other three bit values are determined similarly. Instead of using inequality testing, we can determine the values for a region-code more efficiently using bit-processing operations and the following two steps: (1) Calculate differences between endpoint coordinates and clipping boundaries. (2) Use the resultant sign bit of each difference calculation to set the corresponding value in the region code. For the ordering scheme shown in Figure 10, bit 1 is the sign bit of \( x - x_{\text{min}} \); bit 2 is the sign bit of \( x_{\text{max}} - x \); bit 3 is the sign bit of \( y - y_{\text{min}} \); and bit 4 is the sign bit of \( y_{\text{max}} - y \).

Once we have established region codes for all line endpoints, we can quickly determine which lines are completely inside the clip window and which are completely outside. Any lines that are completely contained within the window edges have a region code of 0000 for both endpoints, and we save these line segments. Any line that has a region-code value of 1 in the same bit position for each endpoint is completely outside the clipping rectangle, and we eliminate that line segment. As an example, a line that has a region code of 1001 for one endpoint and a code of 0101 for the other endpoint is completely to the left of the clipping window, as indicated by the value of 1 in the first bit position of each region code.

We can perform the inside-outside tests for line segments using logical operators. When the or operation between two endpoint region codes for a line segment is false (0000), the line is inside the clipping window. Therefore, we save the line and proceed to test the next line in the scene description. When the and operation between the two endpoint region codes for a line is true (not 0000), the line is completely outside the clipping window, and we can eliminate it from the scene description.

Lines that cannot be identified as being completely inside or completely outside a clipping window by the region-code tests are next checked for intersection with the window border lines. As shown in Figure 12, line segments can intersect clipping boundary lines without entering the interior of the window. Therefore, several intersection calculations might be necessary to clip a line segment, depending on the order in which we process the clipping boundaries. As we process each clipping-window edge, a section of the line is clipped, and the remaining part of the line is checked against the other window borders. We continue eliminating sections until either the line is totally clipped or the remaining part of the line is inside the clipping window. For the following discussion, we assume that the window edges are processed in the following order: left, right, bottom, top. To determine whether a line crosses a selected clipping boundary, we can check corresponding bit values in the two endpoint region codes. If one of these bit values is 1 and the other is 0, the line segment crosses that boundary.

**Figure 12**
Lines extending from one clipping-window region to another may cross into the clipping window, or they could intersect one or more clipping boundaries without entering the window.
Figure 12 illustrates two line segments that cannot be identified immediately as completely inside or completely outside the clipping window. The region codes for the line from \( P_1 \) to \( P_2 \) are 0100 and 1001. Thus, \( P_1 \) is inside the left clipping boundary and \( P_2 \) is outside that boundary. We then calculate the intersection position \( P'_2 \), and we clip off the line section from \( P_2 \) to \( P'_2 \). The remaining portion of the line is inside the right border line, and so we next check the bottom border. Endpoint \( P_1 \) is below the bottom clipping edge and \( P'_2 \) is above it, so we determine the intersection position at this boundary (\( P'_1 \)). We eliminate the line section from \( P_1 \) to \( P'_1 \) and proceed to the top window edge. There we determine the intersection position to be \( P'_2 \). The final step is to clip off the section above the top boundary and save the interior segment from \( P'_1 \) to \( P'_2 \). For the second line, we find that point \( P_3 \) is outside the left boundary and \( P_4 \) is inside. Thus, we calculate the intersection position \( P'_3 \) and eliminate the line section from \( P_3 \) to \( P'_3 \). By checking region codes for the endpoints \( P_3 \) and \( P_4 \), we find that the remainder of the line is below the clipping window and can be eliminated as well.

It is possible, when clipping a line segment using this approach, to calculate an intersection position at all four clipping boundaries, depending on how the line endpoints are processed and what ordering we use for the boundaries. Figure 13 shows the four intersection positions that could be calculated for a line segment that is processed against the clipping-window edges in the order left, right, bottom, top. Therefore, variations of this basic approach have been developed in an effort to reduce the intersection calculations.

To determine a boundary intersection for a line segment, we can use the slope-intercept form of the line equation. For a line with endpoint coordinates \((x_0, y_0)\) and \((x_{\text{end}}, y_{\text{end}})\), the \( y \) coordinate of the intersection point with a vertical clipping border line can be obtained with the calculation

\[
y = y_0 + m(x - x_0)
\]

where the \( x \) value is set to either \( x_{w_{\text{min}}} \) or \( x_{w_{\text{max}}} \), and the slope of the line is calculated as \( m = (y_{\text{end}} - y_0)/(x_{\text{end}} - x_0) \). Similarly, if we are looking for the intersection with a horizontal border, the \( x \) coordinate can be calculated as

\[
x = x_0 + \frac{y - y_0}{m}
\]

with \( y \) set either to \( y_{w_{\text{min}}} \) or to \( y_{w_{\text{max}}} \).
An implementation of the two-dimensional, Cohen-Sutherland line-clipping algorithm is given in the following procedures.

```cpp
class wcPt2D {
    public:
        GLfloat x, y;
};

inline GLint round (const GLfloat a) { return GLint (a + 0.5); }

/* Define a four-bit code for each of the outside regions of a rectangular clipping window. */
const GLint winLeftBitCode = 0x1;
const GLint winRightBitCode = 0x2;
const GLint winBottomBitCode = 0x4;
const GLint winTopBitCode = 0x8;

/* A bit-mask region code is also assigned to each endpoint of an input line segment, according to its position relative to the four edges of an input rectangular clip window. */
inline GLint inside (GLint code) { return GLint (!code); }
inline GLint reject (GLint code1, GLint code2) { return GLint (code1 & code2); }
inline GLint accept (GLint code1, GLint code2) { return GLint (!(code1 | code2)); }

GLubyte encode (wcPt2D pt, wcPt2D winMin, wcPt2D winMax)
{
    GLubyte code = 0x00;
    if (pt.x < winMin.x)
        code = code | winLeftBitCode;
    if (pt.x > winMax.x)
        code = code | winRightBitCode;
    if (pt.y < winMin.y)
        code = code | winBottomBitCode;
    if (pt.y > winMax.y)
        code = code | winTopBitCode;
    return (code);
}
```
void swapPts (wcPt2D * p1, wcPt2D * p2)
{
    wcPt2D tmp;
    tmp = *p1; *p1 = *p2; *p2 = tmp;
}

void swapCodes (GLubyte * c1, GLubyte * c2)
{
    GLubyte tmp;
    tmp = *c1; *c1 = *c2; *c2 = tmp;
}

void lineClipCohSuth (wcPt2D winMin, wcPt2D winMax, wcPt2D p1, wcPt2D p2)
{
    GLubyte code1, code2;
    GLint done = false, plotLine = false;
    GLfloat m;
    while (!done) {
        code1 = encode (p1, winMin, winMax);
        code2 = encode (p2, winMin, winMax);
        if (accept (code1, code2)) {
            done = true;
            plotLine = true;
        }
        else
            if (reject (code1, code2))
                done = true;
            else {
                /* Label the endpoint outside the display window as p1. */
                if (inside (code1)) {
                    swapPts (&p1, &p2);
                    swapCodes (&code1, &code2);
                }
                /* Use slope m to find line-clipEdge intersection. */
                if (p2.x != p1.x)
                    m = (p2.y - p1.y) / (p2.x - p1.x);
                if (code1 & winLeftBitCode) {
                    p1.x += (winMin.x - p1.x) * m;
                    p1.y = winMin.y;
                }
                else
                    if (code1 & winRightBitCode) {
                        p1.y += (winMax.x - p1.x) * m;
                        p1.x = winMax.x;
                    }
                    else
                        if (code1 & winBottomBitCode) {
                            /* Need to update p1.x for nonvertical lines only. */
                            if (p2.x != p1.x)
                                p1.x += (winMin.y - p1.y) / m;
                            p1.y = winMin.y;
                        }

Liang-Barsky Line Clipping

Faster line-clipping algorithms have been developed that do more line testing before proceeding to the intersection calculations. One of the earliest efforts in this direction is an algorithm developed by Cyrus and Beck, which is based on analysis of the parametric line equations. Later, Liang and Barsky independently devised an even faster form of the parametric line-clipping algorithm.

For a line segment with endpoints \((x_0, y_0)\) and \((x_{\text{end}}, y_{\text{end}})\), we can describe the line with the parametric form

\[
\begin{align*}
x &= x_0 + u \Delta x \\
y &= y_0 + u \Delta y
\end{align*}
\]

where \(\Delta x = x_{\text{end}} - x_0\) and \(\Delta y = y_{\text{end}} - y_0\). In the Liang-Barsky algorithm, the parametric line equations are combined with the point-clipping conditions \(12\) to obtain the inequalities

\[
\begin{align*}
x w_{\text{min}} &\leq x_0 + u \Delta x \leq x w_{\text{max}} \\
y w_{\text{min}} &\leq y_0 + u \Delta y \leq y w_{\text{max}}
\end{align*}
\]

which can be expressed as

\[
u p_k \leq q_k, \quad k = 1, 2, 3, 4
\]

where parameters \(p\) and \(q\) are defined as

\[
\begin{align*}
p_1 &= -\Delta x, & q_1 &= x_0 - x w_{\text{min}} \\
p_2 &= \Delta x, & q_2 &= x w_{\text{max}} - x_0 \\
p_3 &= -\Delta y, & q_3 &= y_0 - y w_{\text{min}} \\
p_4 &= \Delta y, & q_4 &= y w_{\text{max}} - y_0
\end{align*}
\]

Any line that is parallel to one of the clipping-window edges has \(p_k = 0\) for the value of \(k\) corresponding to that boundary, where \(k = 1, 2, 3,\) and \(4\) correspond to the left, right, bottom, and top boundaries, respectively. If, for that value of \(k\), we also find \(q_k < 0\), then the line is completely outside the boundary and can be eliminated from further consideration. If \(q_k \geq 0\), the line is inside the parallel clipping border.

When \(p_k < 0\), the infinite extension of the line proceeds from the outside to the inside of the infinite extension of this particular clipping-window edge. If \(p_k > 0\), the line proceeds from the inside to the outside. For a nonzero value of \(p_k\), we can calculate the value of \(u\) that corresponds to the point where the infinitely
extended line intersects the extension of window edge \( k \) as
\[ u = \frac{q_k}{p_k} \] (20)

For each line, we can calculate values for parameters \( u_1 \) and \( u_2 \) that define that part of the line that lies within the clip rectangle. The value of \( u_1 \) is determined by looking at the rectangle edges for which the line proceeds from the outside to the inside \((p < 0)\). For these edges, we calculate \( r_k = \frac{q_k}{p_k} \). The value of \( u_1 \) is taken as the largest of the set consisting of 0 and the various values of \( r \). Conversely, the value of \( u_2 \) is determined by examining the boundaries for which the line proceeds from inside to outside \((p > 0)\). A value of \( r_k \) is calculated for each of these boundaries, and the value of \( u_2 \) is the minimum of the set consisting of 1 and the calculated \( r \) values. If \( u_1 > u_2 \), the line is completely outside the clip window and it can be rejected. Otherwise, the endpoints of the clipped line are calculated from the two values of parameter \( u \).

This algorithm is implemented in the following code sections. Line intersection parameters are initialized to the values \( u_1 = 0 \) and \( u_2 = 1 \). For each clipping boundary, the appropriate values for \( p \) and \( q \) are calculated and used by the function \texttt{clipTest} to determine whether the line can be rejected or whether the intersection parameters are to be adjusted. When \( p < 0 \), parameter \( r \) is used to update \( u_1 \); when \( p > 0 \), parameter \( r \) is used to update \( u_2 \). If updating \( u_1 \) or \( u_2 \) results in \( u_1 > u_2 \), we reject the line. Otherwise, we update the appropriate \( u \) parameter only if the new value results in a shortening of the line. When \( p = 0 \) and \( q < 0 \), we can eliminate the line because it is parallel to and outside this boundary. If the line has not been rejected after all four values of \( p \) and \( q \) have been tested, the endpoints of the clipped line are determined from values of \( u_1 \) and \( u_2 \).

```cpp
class wcPt2D
{
    private:
        GLfloat x, y;

    public:
        /* Default Constructor: initialize position as (0.0, 0.0). */
        wcPt3D ( ) {
            x = y = 0.0;
        }

        setCoords (GLfloat xCoord, GLfloat yCoord) {
            x = xCoord;
            y = yCoord;
        }

        GLfloat getx ( ) const {
            return x;
        }

        GLfloat gety ( ) const {
            return y;
        }
};

inline GLint round (const GLfloat a) { return GLint (a + 0.5); }
```
GLint clipTest (GLfloat p, GLfloat q, GLfloat * u1, GLfloat * u2)
{
    GLfloat r;
    GLint returnValue = true;
    if (p < 0.0) {
        r = q / p;
        if (r > *u2)
            returnValue = false;
        else
            if (r > *u1)
                *u1 = r;
    } else
    if (p > 0.0) {
        r = q / p;
        if (r < *u1)
            returnValue = false;
        else if (r < *u2)
            *u2 = r;
    } else
    /* Thus p = 0 and line is parallel to clipping boundary. */
    if (q < 0.0)
    /* Line is outside clipping boundary. */
        returnValue = false;

    return (returnValue);
}

void lineClipLiangBarsk (wcPt2D winMin, wcPt2D winMax, wcPt2D p1, wcPt2D p2)
{
    GLfloat u1 = 0.0, u2 = 1.0, dx = p2.getx () - p1.getx (), dy;
    if (clipTest (-dx, p1.getx () - winMin.getx (), &u1, &u2))
        if (clipTest (dx, winMax.getx () - p1.getx (), &u1, &u2))
            dy = p2.gety () - p1.gety ();
        if (clipTest (-dy, p1.gety () - winMin.gety (), &u1, &u2))
            if (clipTest (dy, winMax.gety () - p1.gety (), &u1, &u2))
                if (u2 < 1.0) {
                    p2.setCoords (p1.getx () + u2 * dx, p1.gety () + u2 * dy);
                } if (u1 > 0.0) {
                    p1.setCoords (p1.getx () + u1 * dx, p1.gety () + u1 * dy);
                } lineBres (round (p1.getx ()), round (p1.gety ()),
                        round (p2.getx ()), round (p2.gety ()));
    }
}

In general, the Liang-Barsky algorithm is more efficient than the Cohen-Sutherland line-clipping algorithm. Each update of parameters u1 and u2 requires only one division; and window intersections of the line are computed only once, when the final values of u1 and u2 have been computed. In contrast, the Cohen
and Sutherland algorithm can calculate intersections repeatedly along a line path, even though the line may be completely outside the clip window. In addition, each Cohen-Sutherland intersection calculation requires both a division and a multiplication. The two-dimensional Liang-Barsky algorithm can be extended to clip three-dimensional lines.

Nicholl-Lee-Nicholl Line Clipping

By creating more regions around the clipping window, the Nicholl-Lee-Nicholl (NLN) algorithm avoids multiple line-intersection calculations. In the Cohen-Sutherland method, for example, multiple intersections could be calculated along the path of a line segment before an intersection on the clipping rectangle is located or the line is completely rejected. These extra intersection calculations are eliminated in the NLN algorithm by carrying out more region testing before intersection positions are calculated. Compared to both the Cohen-Sutherland and the Liang-Barsky algorithms, the Nicholl-Lee-Nicholl algorithm performs fewer comparisons and divisions. The trade-off is that the NLN algorithm can be applied only to two-dimensional clipping, whereas both the Liang-Barsky and the Cohen-Sutherland methods are easily extended to three-dimensional scenes.

Initial testing to determine whether a line segment is completely inside the clipping window or outside the window limits can be accomplished with region-code tests, as in the previous two algorithms. If a trivial acceptance or rejection of the line is not possible, the NLN algorithm proceeds to set up additional clipping regions.

For a line with endpoints \(P_0\) and \(P_{\text{end}}\), we first determine the position of point \(P_0\) for the nine possible regions relative to the clipping window. Only the three regions shown in Figure 14 need be considered. If \(P_0\) lies in any one of the other six regions, we can move it to one of the three regions in Figure 14 using a symmetry transformation. For example, the region directly above the clip window can be transformed to the region left of the window using a reflection about the line \(y = -x\), or we could use a 90° counterclockwise rotation.

Assuming that \(P_0\) and \(P_{\text{end}}\) are not both inside the clipping window, we next determine the position of \(P_{\text{end}}\) relative to \(P_0\). To do this, we create some new regions in the plane, depending on the location of \(P_0\). Boundaries of the new regions are semi-infinite line segments that start at the position of \(P_0\) and pass through the clipping-window corners. If \(P_0\) is inside the clipping window, we set up the four regions shown in Figure 15. Then, depending on which one of the four regions (L, T, R, or B) contains \(P_{\text{end}}\), we compute the line-intersection position with the corresponding window boundary.

If \(P_0\) is in the region to the left of the window, we set up the four regions labeled L, LT, LR, and LB in Figure 16. These four regions again determine a unique
clipping-window edge for the line segment, relative to the position of $P_{\text{end}}$. For instance, if $P_{\text{end}}$ is in any one of the three regions labeled L, we clip the line at the left window border and save the line segment from this intersection point to $P_{\text{end}}$. If $P_{\text{end}}$ is in region LT, we save the line segment from the left window boundary to the top boundary. Similar processing is carried out for regions LR and LB. However, if $P_{\text{end}}$ is not in any of the four regions L, LT, LR, or LB, the entire line is clipped.

For the third case, when $P_0$ is to the left and above the clipping window, we use the regions in Figure 17. In this case, we have the two possibilities shown, depending on the position of $P_0$ within the top-left corner of the clipping window. When $P_0$ is closer to the left clipping boundary of the window, we use the regions in (a) of this figure. Otherwise, when $P_0$ is closer to the top clipping boundary of the window, we use the regions in (b). If $P_{\text{end}}$ is in one of the regions T, L, TR, TB, LR, or LB, this determines a unique clipping-window border for the intersection calculations. Otherwise, the entire line is rejected.

To determine the region in which $P_{\text{end}}$ is located, we compare the slope of the line segment to the slopes of the boundaries of the NLN regions. For example, if $P_0$ is left of the clipping window (Figure 16), then $P_{\text{end}}$ is in region LT if

$$\text{slope} P_0 P_{TR} < \text{slope} P_0 P_{\text{end}} < \text{slope} P_0 P_{TL}$$

or

$$\frac{y_T - y_0}{x_R - x_0} < \frac{y_{\text{end}} - y_0}{x_{\text{end}} - x_0} < \frac{y_T - y_0}{x_L - x_0}$$

or

$$\frac{y_R - y_0}{x_T - x_0} < \frac{y_{\text{end}} - y_0}{x_{\text{end}} - x_0} < \frac{y_R - y_0}{x_B - x_0}$$

or

$$\frac{y_T - y_0}{x_L - x_0} < \frac{y_{\text{end}} - y_0}{x_{\text{end}} - x_0} < \frac{y_T - y_0}{x_R - x_0}$$

or

$$\frac{y_B - y_0}{x_T - x_0} < \frac{y_{\text{end}} - y_0}{x_{\text{end}} - x_0} < \frac{y_B - y_0}{x_L - x_0}$$

FIGURE 17
The two possible sets of clipping regions used in the NLN algorithm when $P_0$ is above and to the left of the clipping window.
Two-Dimensional Viewing

We clip the entire line if

\[(y_T - y_0)(x_{\text{end}} - x_0) < (x_L - x_0)(y_{\text{end}} - y_0)\]  \(23\)

The coordinate-difference calculations and product calculations used in the slope tests are saved and also used in the intersection calculations. From the parametric equations

\[x = x_0 + (x_{\text{end}} - x_0)u\]
\[y = y_0 + (y_{\text{end}} - y_0)u\]

we calculate an \(x\)-intersection position on the left window boundary as \(x = x_L\), with \(u = (x_L - x_0)/(x_{\text{end}} - x_0)\), so that the \(y\)-intersection position is

\[y = y_0 + \frac{y_{\text{end}} - y_0}{x_{\text{end}} - x_0}(x_L - x_0)\]  \(24\)

An intersection position on the top boundary has \(y = y_T\) and \(u = (y_T - y_0)/(y_{\text{end}} - y_0)\), with

\[x = x_0 + \frac{x_{\text{end}} - x_0}{y_{\text{end}} - y_0}(y_T - y_0)\]  \(25\)

Line Clipping Using Nonrectangular Polygon Clip Windows

In some applications, it may be desirable to clip lines against arbitrarily shaped polygons. Methods based on parametric line equations, such as either the Cyrus-Beck algorithm or the Liang-Barsky algorithm, can be readily extended to clip lines against convex polygon windows. We do this by modifying the algorithm to include the parametric equations for the boundaries of the clipping region. Preliminary screening of line segments can be accomplished by processing lines against the coordinate extents of the clipping polygon.

For concave-polygon clipping regions, we could still apply these parametric clipping procedures if we first split the concave polygon into a set of convex polygons. Another approach is simply to add one or more edges to the concave clipping area so that it is modified to a convex-polygon shape. Then a series of clipping operations can be applied using the modified convex polygon components, as illustrated in Figure 18. The line segment \(P_1P_2\) in (a) of this figure is to be clipped by the concave window with vertices \(V_1, V_2, V_3, V_4,\) and \(V_5\). Two convex clipping regions are obtained, in this case, by adding a line segment from \(V_4\) to \(V_1\). Then the line is clipped in two passes: (1) Line \(P_1P_2\) is clipped by the convex polygon with vertices \(V_1, V_2, V_3,\) and \(V_4\) to yield the clipped segment \(P_1P_2'\) [Figure 18(b)]. (2) The internal line segment \(P_1P_2'\) is clipped off using the convex polygon with vertices \(V_1, V_5,\) and \(V_4\) [Figure 18(c)] to yield the final clipped line segment \(P_1P_2''\).

Line Clipping Using Nonlinear Clipping-Window Boundaries

Circles or other curved-boundary clipping regions are also possible, but they require more processing because the intersection calculations involve nonlinear equations. At the first step, lines could be clipped against the bounding rectangle (coordinate extents) of the curved clipping region. Lines that are outside the coordinate extents are eliminated. To identify lines that are inside a circle, for instance, we could calculate the distance of the line endpoints from the circle center. If the square of this distance for both endpoints of a line is less than or equal to the radius squared, we can save the entire line. The remaining lines are then processed through the intersection calculations, which must solve simultaneous circle-line equations.
8 Polygon Fill-Area Clipping

Graphics packages typically support only fill areas that are polygons, and often only convex polygons. To clip a polygon fill area, we cannot apply a line-clipping method to the individual polygon edges directly because this approach would not, in general, produce a closed polyline. Instead, a line clipper would often produce a disjoint set of lines with no complete information about how we might form a closed boundary around the clipped fill area. Figure 19 illustrates a possible output from a line-clipping procedure applied to the edges of a polygon fill area. What we require is a procedure that will output one or more closed polylines for the boundaries of the clipped fill area, so that the polygons can be scan-converted to fill the interiors with the assigned color or pattern, as in Figure 20.

We can process a polygon fill area against the borders of a clipping window using the same general approach as in line clipping. A line segment is defined by its two endpoints, and these endpoints are processed through a line-clipping procedure by constructing a new set of clipped endpoints at each clipping-window boundary. Similarly, we need to maintain a fill area as an entity as it is processed through the clipping stages. Thus, we can clip a polygon fill area by determining the new shape for the polygon as each clipping-window edge is processed, as demonstrated in Figure 21. Of course, the interior fill for the polygon would not be applied until the final clipped border had been determined.
Just as we first tested a line segment to determine whether it could be completely saved or completely clipped, we can do the same with a polygon fill area by checking its coordinate extents. If the minimum and maximum coordinate values for the fill area are inside all four clipping boundaries, the fill area is saved for further processing. If these coordinate extents are all outside any of the clipping-window borders, we eliminate the polygon from the scene description (Figure 22).

When we cannot identify a fill area as being completely inside or completely outside the clipping window, we then need to locate the polygon intersection positions with the clipping boundaries. One way to implement convex-polygon
clipping is to create a new vertex list at each clipping boundary, and then pass this new vertex list to the next boundary clipper. The output of the final clipping stage is the vertex list for the clipped polygon (Figure 23). For concave-polygon clipping, we would need to modify this basic approach so that multiple vertex lists could be generated.

**Sutherland-Hodgman Polygon Clipping**

An efficient method for clipping a convex-polygon fill area, developed by Sutherland and Hodgman, is to send the polygon vertices through each clipping stage so that a single clipped vertex can be immediately passed to the next stage. This eliminates the need for an output set of vertices at each clipping stage, and it allows the boundary-clipping routines to be implemented in parallel. The final output is a list of vertices that describe the edges of the clipped polygon fill area.

Because the Sutherland-Hodgman algorithm produces only one list of output vertices, it cannot correctly generate the two output polygons in Figure 20(b) that are the result of clipping the concave polygon shown in Figure 20(a). However, more processing steps can be added to the algorithm to allow it to produce multiple output vertex lists, so that general concave-polygon clipping could be accommodated. And the basic Sutherland-Hodgman algorithm is able to process concave polygons when the clipped fill area can be described with a single vertex list.

The general strategy in this algorithm is to send the pair of endpoints for each successive polygon line segment through the series of clippers (left, right, bottom, and top). As soon as a clipper completes the processing of one pair of vertices, the clipped coordinate values, if any, for that edge are sent to the next clipper. Then the first clipper processes the next pair of endpoints. In this way, the individual boundary clippers can be operating in parallel.

There are four possible cases that need to be considered when processing a polygon edge against one of the clipping boundaries. One possibility is that the first edge endpoint is outside the clipping boundary and the second endpoint is inside. Or, both endpoints could be inside this clipping boundary. Another possibility is that the first endpoint is inside the clipping boundary and the second endpoint is outside. And, finally, both endpoints could be outside the clipping boundary.

To facilitate the passing of vertices from one clipping stage to the next, the output from each clipper can be formulated as shown in Figure 24. As each successive pair of endpoints is passed to one of the four clippers, an output is generated for the next clipper according to the results of the following tests:

1. If the first input vertex is outside this clipping-window border and the second vertex is inside, both the intersection point of the polygon edge with the window border and the second vertex are sent to the next clipper.
Two-Dimensional Viewing

FIGURE 24
The four possible outputs generated by the left clipper, depending on the position of a pair of endpoints relative to the left boundary of the clipping window.

FIGURE 25
Processing a set of polygon vertices, \{1, 2, 3\}, through the boundary clippers using the Sutherland-Hodgman algorithm. The final set of clipped vertices is \{1', 2', 2''\}.

2. If both input vertices are inside this clipping-window border, only the second vertex is sent to the next clipper.
3. If the first vertex is inside this clipping-window border and the second vertex is outside, only the polygon edge-intersection position with the clipping-window border is sent to the next clipper.
4. If both input vertices are outside this clipping-window border, no vertices are sent to the next clipper.

The last clipper in this series generates a vertex list that describes the final clipped fill area.

Figure 25 provides an example of the Sutherland-Hodgman polygon-clipping algorithm for a fill area defined with the vertex set \{1, 2, 3\}. As soon as a clipper receives a pair of endpoints, it determines the appropriate output using the tests illustrated in Figure 24. These outputs are passed in succession from the left clipper to the right, bottom, and top clippers. The output from the
void clipPoint (wcPt2D p, Boundary winEdge, wcPt2D wMin, wcPt2D wMax,
wcPt2D * pOut, int * cnt, wcPt2D * first[], wcPt2D * s)
{
    wcPt2D iPt;

    /* If no previous point exists for this clipping boundary, *
     * save this point. */
    if (!first[winEdge])
        first[winEdge] = &p;
    else /* Previous point exists. If p and previous point cross *
        * this clipping boundary, find intersection. Clip against *
        * next boundary, if any. If no more clip boundaries, add *
        * intersection to output list. */
        if (cross (p, s[winEdge], winEdge, wMin, wMax)) {
            iPt = intersect (p, s[winEdge], winEdge, wMin, wMax);
            if (winEdge < Top)
                clipPoint (iPt, b+1, wMin, wMax, pOut, cnt, first, s);
            else {
                pOut[*cnt] = iPt; (*cnt)++;
            }
        }
    /* Save p as most recent point for this clip boundary. */
    s[winEdge] = p;

    /* For all, if point inside, proceed to next boundary, if any. */
    if (inside (p, winEdge, wMin, wMax))
        if (winEdge < Top)
            clipPoint (p, winEdge + 1, wMin, wMax, pOut, cnt, first, s);
        else {
            pOut[*cnt] = p; (*cnt)++;
        }
}

void closeClip (wcPt2D wMin, wcPt2D wMax, wcPt2D * pOut,
GLint * cnt, wcPt2D * first [], wcPt2D * s)
{
    wcPt2D pt;
    Boundary winEdge;

    for (winEdge = Left; winEdge <= Top; winEdge++)
    {
        if (cross (s[winEdge], *first[winEdge], winEdge, wMin, wMax))
        {
            pt = intersect (s[winEdge], *first[winEdge], winEdge, wMin, wMax);
            if (winEdge < Top)
                clipPoint (pt, winEdge + 1, wMin, wMax, pOut, cnt, first, s);
            else {
                pOut[*cnt] = pt; (*cnt)++;
            }
        }
    }
top clipper is the set of vertices defining the clipped fill area. For this example, the output vertex list is \( \{1', 2, 2', 2''\} \).

A sequential implementation of the Sutherland-Hodgman polygon-clipping algorithm is demonstrated in the following set of procedures. An input set of vertices is converted to an output vertex list by clipping it against the four edges of the axis-aligned rectangular clipping region.

typedef enum { Left, Right, Bottom, Top } Boundary;
const GLint nClip = 4;

GLint inside (wcPt2D p, Boundary b, wcPt2D wMin, wcPt2D wMax)
{
    switch (b) {
    case Left: if (p.x < wMin.x) return (false); break;
    case Right: if (p.x > wMax.x) return (false); break;
    case Bottom: if (p.y < wMin.y) return (false); break;
    case Top: if (p.y > wMax.y) return (false); break;
    }
    return (true);
}

GLint cross (wcPt2D p1, wcPt2D p2, Boundary winEdge, wcPt2D wMin, wcPt2D wMax)
{
    if (inside (p1, winEdge, wMin, wMax) == inside (p2, winEdge, wMin, wMax))
        return (false);
    else return (true);
}

wcPt2D intersect (wcPt2D p1, wcPt2D p2, Boundary winEdge, wcPt2D wMin, wcPt2D wMax)
{
    wcPt2D iPt;
    GLfloat m;
    if (p1.x != p2.x) m = (p1.y - p2.y) / (p1.x - p2.x);
    switch (winEdge) {
    case Left:
        iPt.x = wMin.x;
        iPt.y = p2.y + (wMin.x - p2.x) * m;
        break;
    case Right:
        iPt.x = wMax.x;
        iPt.y = p2.y + (wMax.x - p2.x) * m;
        break;
    case Bottom:
        iPt.y = wMin.y;
        if (p1.x != p2.x) iPt.x = p2.x + (wMin.y - p2.y) / m;
        else iPt.x = p2.x;
        break;
    case Top:
        iPt.y = wMax.y;
        if (p1.x != p2.x) iPt.x = p2.x + (wMax.y - p2.y) / m;
        else iPt.x = p2.x;
        break;
    }
GLint polygonClipSuthHodg (wcPt2D wMin, wcPt2D wMax, GLint n, wcPt2D * pIn, wcPt2D * pOut)
{
    /* Parameter "first" holds pointer to first point processed for
    * a boundary; "s" holds most recent point processed for boundary.
    */
    wcPt2D * first[nClip] = { 0, 0, 0, 0 }, s[nClip];
    GLint k, cnt = 0;

    for (k = 0; k < n; k++)
        clipPoint (pIn[k], Left, wMin, wMax, pOut, &cnt, first, s);

    closeClip (wMin, wMax, pOut, &cnt, first, s);
    return (cnt);
}

When a concave polygon is clipped with the Sutherland-Hodgman algorithm, extraneous lines may be displayed. An example of this effect is demonstrated in Figure 26. This occurs when the clipped polygon should have two or more separate sections. But since there is only one output vertex list, the last vertex in the list is always joined to the first vertex.

There are several things we can do to display clipped concave polygons correctly. For one, we could split a concave polygon into two or more convex polygons and process each convex polygon separately using the Sutherland-Hodgman algorithm. Another possibility is to modify the Sutherland-Hodgman method so that the final vertex list is checked for multiple intersection points along any clipping-window boundary. If we find more than two vertex positions along any clipping boundary, we can separate the list of vertices into two or more lists that correctly identify the separate sections of the clipped fill area. This may require extensive analysis to determine whether some points along the clipping boundary should be paired or whether they represent single vertex points that have been clipped. A third possibility is to use a more generalized polygon clipper that has been designed to process concave polygons correctly.

**Weiler-Atherton Polygon Clipping**

This algorithm provides a general polygon-clipping approach that can be used to clip a fill area that is either a convex polygon or a concave polygon. Moreover, the method was developed as a means for identifying visible surfaces in a three-dimensional scene. Therefore, we could also use this approach to clip any polygon fill area against a clipping window with any polygon shape.

![Clipping Window](image)

**Figure 26**

Clipping the concave polygon in (a) using the Sutherland-Hodgman algorithm produces the two connected areas in (b).
Instead of simply clipping the fill-area edges as in the Sutherland-Hodgman method, the Weiler-Atherton algorithm traces around the perimeter of the fill polygon searching for the borders that enclose a clipped fill region. In this way, multiple fill regions, as in Figure 26(b), can be identified and displayed as separate, unconnected polygons. To find the edges for a clipped fill area, we follow a path (either counterclockwise or clockwise) around the fill area that detours along a clipping-window boundary whenever a polygon edge crosses to the outside of that boundary. The direction of a detour at a clipping-window border is the same as the processing direction for the polygon edges.

We can usually determine whether the processing direction is counterclockwise or clockwise from the ordering of the vertex list that defines a polygon fill area. In most cases, the vertex list is specified in a counterclockwise order as a means for defining the front face of the polygon. Thus, the cross-product of two successive edge vectors that form a convex angle determines the direction for the normal vector, which is in the direction from the back face to the front face of the polygon. If we do not know the vertex ordering, we could calculate the normal vector, or we can locate the interior of the fill area from any reference position. Then, if we sequentially process the edges so that the polygon interior is always on our left, we obtain a counterclockwise traversal. Otherwise, with the interior to our right, we have a clockwise traversal.

For a counterclockwise traversal of the polygon fill-area vertices, we apply the following Weiler-Atherton procedures:

1. Process the edges of the polygon fill area in a counterclockwise order until an inside-outside pair of vertices is encountered for one of the clipping boundaries; that is, the first vertex of the polygon edge is inside the clip region and the second vertex is outside the clip region.
2. Follow the window boundaries in a counterclockwise direction from the exit-intersection point to another intersection point with the polygon. If this is a previously processed point, proceed to the next step. If this is a new intersection point, continue processing polygon edges in a counterclockwise order until a previously processed vertex is encountered.
3. Form the vertex list for this section of the clipped fill area.
4. Return to the exit-intersection point and continue processing the polygon edges in a counterclockwise order.

Figure 27 illustrates the Weiler-Atherton clipping of a concave polygon against a standard, rectangular clipping window for a counterclockwise traversal of the polygon edges. For a clockwise edge traversal, we would use a clockwise clipping-window traversal.

Starting from the vertex labeled 1 in Figure 27(a), the next polygon vertex to process in a counterclockwise order is labeled 2. Thus, this edge exits the clipping window at the top boundary. We calculate this intersection position (point 1′) and make a left turn there to process the window borders in a counterclockwise direction. Proceeding along the top border of the clipping window, we do not intersect a polygon edge before reaching the left window boundary. Therefore, we label this position as vertex 1″ and follow the left boundary to the intersection position 1‴. We then follow this polygon edge in a counterclockwise direction, which returns us to vertex 1. This completes a circuit of the window boundaries and identifies the vertex list \{1, 1′, 1″, 1‴\} as a clipped region of the original fill area. Processing of the polygon edges is then resumed at point 1′. The edge defined by points 2 and 3 crosses to the outside of the left boundary, but points 2 and 2′
FIGURE 27  A concave polygon (a), defined with the vertex list \{1, 2, 3, 4, 5, 6\}, is clipped using the Weiler-Atherton algorithm to generate the two lists \{1, 1′, 1″, 1‴\} and \{4′, 5, 5′\}, which represent the separate polygon fill areas shown in (b).

FIGURE 28  Clipping a polygon fill area against a concave-polygon clipping window using the Weiler-Atherton algorithm.

are above the top clipping-window border and points 2′ and 3 are to the left of the clipping region. Also, the edge with endpoints 3 and 4 is outside the left clipping boundary, but the next edge (from endpoint 4 to endpoint 5) reenters the clipping region and we pick up intersection point 4′. The edge with endpoints 5 and 6 exits the window at intersection position 5′, so we detour down the left clipping boundary to obtain the closed vertex list \{4′, 5, 5′\}. We resume the polygon edge processing at position 5′, which returns us to the previously processed point 1″′.

At this point, all polygon vertices and edges have been processed, so the fill area is completely clipped.

Polygon Clipping Using Nonrectangular Polygon Clip Windows

The Liang-Barsky algorithm and other parametric line-clipping methods are particularly well suited for processing polygon fill areas against convex-polygon clipping windows. In this approach, we use a parametric representation for the edges of both the fill area and the clipping window, and both polygons are represented with a vertex list. We first compare the positions of the bounding rectangles for the fill area and the clipping polygon. If we cannot identify the fill area as completely outside the clipping polygon, we can use inside-outside tests to process the parametric edge equations. After completing all the region tests, we solve pairs of simultaneous parametric line equations to determine the window intersection positions.

We can also process any polygon fill area against any polygon-shaped clipping window (convex or concave), as in Figure 28, using the edge-traversal approach of the Weiler-Atherton algorithm. In this case, we need to maintain a vertex list for the clipping window as well as for the fill area, with both lists arranged in a counterclockwise (or clockwise) order. In addition, we need to apply
inside-outside tests to determine whether a fill-area vertex is inside or outside a particular clipping-window boundary. As in the previous examples, we follow the window boundaries whenever a fill-area edge exits a clipping boundary. This clipping method can also be used when either the fill area or the clipping window contains holes that are defined with polygon borders. In addition, we can use this basic approach in constructive solid-geometry applications to identify the result of a union, intersection, or difference operation on two polygons. In fact, locating the clipped region of a fill area is equivalent to determining the intersection of two planar areas.

**Polygon Clipping Using Nonlinear Clipping-Window Boundaries**

One method for processing a clipping window with curved boundaries is to approximate the boundaries with straight-line sections and use one of the algorithms for clipping against a general polygon-shaped clipping window. Alternatively, we could use the same general procedures that we discussed for line segments. First, we can compare the coordinate extents of the fill area to the coordinate extents of the clipping window. Depending on the shape of the clipping window, we may also be able to perform some other region tests based on symmetric considerations. For fill areas that cannot be identified as completely inside or completely outside the clipping window, we ultimately need to calculate the window intersection positions with the fill area.

**9 Curve Clipping**

Areas with curved boundaries can be clipped with methods similar to those discussed in the previous sections. If the objects are approximated with straight-line boundary sections, we use a polygon-clipping method. Otherwise, the clipping procedures involve nonlinear equations, and this requires more processing than for objects with linear boundaries.

We can first test the coordinate extents of an object against the clipping boundaries to determine whether it is possible to accept or reject the entire object trivially. If not, we could check for object symmetries that we might be able to exploit in the initial accept/reject tests. For example, circles have symmetries between quadrants and octants, so we could check the coordinate extents of these individual circle regions. We cannot reject the complete circular fill area in Figure 29 just by checking its overall coordinate extents. But half of the circle is outside the right clipping border (or outside the top border), the upper-left quadrant is above the top clipping border, and the remaining two octants can be similarly eliminated.

An intersection calculation involves substituting a clipping-boundary position \((x_{w_{\min}}, x_{w_{\max}}, y_{w_{\min}}, \text{or } y_{w_{\max}})\) in the nonlinear equation for the object boundary and solving for the other coordinate value. Once all intersection positions have been evaluated, the defining positions for the object can be stored for later use by the scan-line fill procedures. Figure 30 illustrates circle clipping against a rectangular window. For this example, the circle radius and the endpoints of the clipped arc can be used to fill the clipped region, by invoking the circle algorithm to locate positions along the arc between the intersection endpoints.

Similar procedures can be applied when clipping a curved object against a general polygon clipping region. On the first pass, we could compare the bounding rectangle of the object with the bounding rectangle of the clipping region. If this does not save or eliminate the entire object, we next solve the simultaneous line-curve equations to determine the clipping intersection points.
10 Text Clipping

Several techniques can be used to provide text clipping in a graphics package. In a particular application, the choice of clipping method depends on how characters are generated and what requirements we have for displaying character strings.

The simplest method for processing character strings relative to the limits of a clipping window is to use the \textit{all-or-none string-clipping} strategy shown in Figure 31. If all of the string is inside the clipping window, we display the entire string. Otherwise, the entire string is eliminated. This procedure is implemented by examining the coordinate extents of the text string. If the coordinate limits of this bounding rectangle are not completely within the clipping window, the string is rejected.

An alternative is to use the \textit{all-or-none character-clipping} strategy. Here we eliminate only those characters that are not completely inside the clipping window (Figure 32). In this case, the coordinate extents of individual characters are compared to the window boundaries. Any character that is not completely within the clipping-window boundary is eliminated.

A third approach to text clipping is to clip the components of individual characters. This provides the most accurate display of clipped character strings, but it requires the most processing. We now treat characters in much the same way that we treated lines or polygons. If an individual character overlaps a clipping window, we clip off only the parts of the character that are outside the window (Figure 33). Outline character fonts defined with line segments are processed in this way using a polygon-clipping algorithm. Characters defined with bit maps are clipped by comparing the relative position of the individual pixels in the character grid patterns to the borders of the clipping region.
11 Summary

The two-dimensional viewing-transformation pipeline is a series of operations that result in the display of a world-coordinate picture that has been defined in the \(xy\) plane. After we construct the scene, it can be mapped to a viewing-coordinate reference frame, then to a normalized coordinate system where clipping routines can be applied. Finally, the scene is transferred to device coordinates for display. Normalized coordinates can be specified in the range from 0 to 1 or in the range from \(-1\) to 1, and they are used to make graphics packages independent of the output-device requirements.

We select part of a scene for display on an output device using a clipping window, which can be described in the world-coordinate system or in a viewing-coordinate frame defined relative to world coordinates. The contents of the clipping window are transferred to a viewport for display on an output device. In some systems, a viewport is specified within normalized coordinates. Other systems specify the viewport in device coordinates. Typically, the clipping window and viewport are rectangles whose edges are parallel to the coordinate axes. An object is mapped to the viewport so that it has the same relative position in the viewport as it has in the clipping window. To maintain the relative proportions of an object, the viewport must have the same aspect ratio as the corresponding clipping window. In addition, we can set up any number of clipping windows and viewports for a scene.

Clipping algorithms are usually implemented in normalized coordinates, so that all geometric transformations and viewing operations that are independent of device coordinates can be concatenated into one transformation matrix. With the viewport specified in device coordinates, we can clip a two-dimensional scene against a normalized, symmetric square, with normalized coordinates varying from \(-1\) to 1, before transferring the contents of the normalized, symmetric square to the viewport.

All graphics packages include routines for clipping straight-line segments and polygon fill areas. Packages that contain functions for specifying single point positions or text strings also include clipping routines for those graphics primitives. Because the clipping calculations are time-consuming, the development of improved clipping algorithms continues to be an area of major concern in computer graphics. Cohen and Sutherland developed a line-clipping algorithm that uses a region code to identify the position of a line endpoint relative to the clipping-window boundaries. Endpoint region codes are used to identify quickly those lines that are completely inside the clipping window and some of the lines that are completely outside. For the remaining lines, intersection positions at the window boundaries must be calculated. Liang and Barsky developed a faster line-clipping algorithm that represents line segments with parametric equations, similar to the Cyrus-Beck algorithm. This approach allows more testing to be accomplished before proceeding to the intersection calculations. The Nicholl-Lee-Nicholl (NLN) algorithm further reduces intersection calculations by using more region testing in the \(xy\) plane. Parametric line-clipping methods are extended easily to convex clipping windows and to three-dimensional scenes. However, the NLN approach applies only to two-dimensional line segments.

Algorithms for clipping straight-line segments against concave-polygon clipping windows have also been developed. One approach is to split a concave clipping window into a set of convex polygons and apply the parametric line-clipping methods. Another approach is to add edges to the concave window to modify it to a convex shape. Then a series of exterior and interior clipping operations can be performed to obtain the clipped line segment.
## Summary of OpenGL Two-Dimensional Viewing Functions

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<td>Renews the contents of the current display window.</td>
</tr>
<tr>
<td>glutMainLoop</td>
<td>Executes the computer-graphics program.</td>
</tr>
<tr>
<td>glutIdleFunc</td>
<td>Specifies a function to execute when the system is idle.</td>
</tr>
<tr>
<td>glutGet</td>
<td>Queries the system about a specified state parameter.</td>
</tr>
</tbody>
</table>
Although clipping windows with curved boundaries are rarely used, we can apply similar line-clipping methods. However, intersection calculations now involve nonlinear equations.

A polygon fill area is defined with a vertex list, and polygon-clipping procedures must retain information about how the clipped edges are to be connected as the polygon proceeds through the various processing stages. In the Sutherland-Hodgman algorithm, pairs of fill-area vertices are processed by each boundary clipper in turn, and clipping information for that edge is passed immediately to the next clipper, which allows the four clipping routines (left, right, bottom, and top) to be operating in parallel. This algorithm provides an efficient method for clipping convex-polygon fill areas. However, when a clipped concave polygon contains disjoint sections, the Sutherland-Hodgman algorithm produces extraneous connecting line segments. Extensions of parametric line clippers, such as the Liang-Barsky method, can also be used to clip convex polygon fill areas. Both convex and concave fill areas can be clipped correctly with the Weiler-Atherton algorithm, which uses a boundary-traversal approach.

Fill areas can be clipped against convex clipping windows using an extension of the parametric line-representation approach. And the Weiler-Atherton method can clip any polygon fill area using any polygon-shaped clipping window. Fill areas can be clipped against windows with nonlinear boundaries by using a polygon approximation for the window or by processing the fill area against the curved window boundaries.

The fastest text-clipping method is the all-or-none strategy, which completely clips a text string if any part of the string is outside any clipping-window boundary. Alternatively, we could clip a text string by eliminating only those characters in the string that are not completely inside the clipping window. And the most accurate text-clipping method is to apply either point, line, polygon, or curve clipping to the individual characters in a string, depending on whether characters are defined as point grids or outline fonts.

Although OpenGL is designed for three-dimensional applications, a two-dimensional GLU function is provided for specifying a standard, rectangular clipping window in world coordinates. In OpenGL, the clipping-window coordinates are parameters for the projection transformation. Therefore, we first need to invoke the projection matrix mode. Next we can specify the viewport, using a function in the basic OpenGL library, and a display window, using GLUT functions. A wide range of GLUT functions are available for setting various display-window parameters. Table 1 summarizes the OpenGL two-dimensional viewing functions. In addition, the table lists some viewing-related functions.

REFERENCES


Basic polygon-clipping methods are presented in Sutherland and Hodgman (1974) and in Liang and Barsky (1983). General techniques for clipping arbitrarily shaped polygons against each other are given in Weiler and Atherton (1977) and in Weiler (1980).

Viewing operations in OpenGL are discussed in Woo, et al. (1999). Display-window GLUT routines are discussed in Kilgard (1996), and additional information on GLUT can be obtained online at http://reality.sgi.com/opengl/glut3/glut3.html.
EXERCISES

1. Write a procedure to calculate the elements of matrix 1 for transforming two-dimensional world coordinates to viewing coordinates, given the viewing coordinate origin $P_0$ and the view up vector $V$.

2. Derive matrix 8 for transferring the contents of a clipping window to a viewport by first scaling the window to the size of the viewport, then translating the scaled window to the viewport position. Use the center of the clipping window as the reference point for the scaling and translation operations.

3. Write a procedure to calculate the elements of matrix 9 for transforming a clipping window to the symmetric normalized square.

4. Write a set of procedures to implement the two-dimensional viewing pipeline without clipping operations. Your program should allow a scene to be constructed with modeling-coordinate transformations, a specified viewing system, and a transformation to the symmetric normalized square. As an option, a viewing table could be implemented to store different sets of viewing transformation parameters.

5. Write a complete program to implement the Cohen-Sutherland line-clipping algorithm.

6. Modify the program in the previous exercise to produce an animation of a single line whose length is longer than the diagonal length of the viewing window. The midpoint of the line should be placed at the center of the viewing window and the line should rotate clockwise by a small amount in each frame. The clipping algorithm implemented in the previous exercise should clip the line appropriately in each frame.

7. Carefully discuss the rationale behind the various tests and methods for calculating the intersection parameters $u_1$ and $u_2$ in the Liang-Barsky line-clipping algorithm.

8. Compare the number of arithmetic operations performed in the Cohen-Sutherland and the Liang-Barsky line-clipping algorithms for several different line orientations relative to a clipping window.

9. Write a complete program to implement the Liang-Barsky line-clipping algorithm.

10. Modify the program in the previous exercise to produce an animation similar to the one described in Exercise 6. The clipping algorithm implemented in the previous exercise should clip the line appropriately in each frame.

11. Devise symmetry transformations for mapping the intersection calculations for the three regions in Figure 14 to the other six regions of the $xy$ plane.

12. Set up a detailed algorithm for the Nicholl-Lee-Nicholl approach to line clipping for any input pair of line endpoints.

13. Compare the number of arithmetic operations performed in the NLN algorithm to both the Cohen-Sutherland and Liang-Barsky line-clipping algorithms, for several different line orientations relative to a clipping window.

14. Adapt the Liang-Barsky line-clipping algorithm to polygon clipping.

15. Use the implementation of Liang-Barsky polygon clipping developed in the previous exercise to write a program that displays an animation of a moving hexagon in the display window. The hexagon should be displayed as moving into the window from the top-left corner of the window diagonally towards the bottom-right corner and off the screen. Once the hexagon has exited the window completely the animation should repeat.

16. Set up a detailed algorithm for Weiler-Atherton polygon clipping, assuming that the clipping window is a rectangle in standard position.

17. Use the implementation of Weiler-Atherton polygon clipping developed in the previous exercise to write a program that produces a similar animation to the one described in Exercise 14.

18. Devise an algorithm for Weiler-Atherton polygon clipping, where the clipping window can be any convex polygon.

19. Devise an algorithm for Weiler-Atherton polygon clipping, where the clipping window can be any specified polygon (convex or concave).

20. Write a routine to clip an ellipse in standard position against a rectangular window.

21. Assuming that all characters in a text string have the same width, develop a text-clipping algorithm that clips a string according to the all-or-none character-clipping strategy.

22. Use the implementation of text clipping developed in the previous exercise to write a program that displays an animation of a moving marquee in the display window. That is, a sequence of characters should be displayed as moving into the window from the left side, across the window horizontally, and out of the window on the right side. Once all of the characters have exited the viewport on the right side the animation should repeat.

23. Develop a text-clipping algorithm that clips individual characters, assuming that the characters are defined in a pixel grid of a specified size.

24. Use the implementation of text clipping developed in the previous exercise to write a program that performs the same behavior as that in Exercise 21.
IN MORE DEPTH
1 Implement both the Sutherland-Hodgman and Weiler-Atherton polygon-clipping algorithms in two separate routines. Use them to clip the objects in the current snapshot of your scene against a sub-rectangle of the full scene extents. Compare the performance of the two algorithms. Make any modifications necessary to handle clipping of concave polygons in your scene using the Sutherland-Hodgman algorithm. The routines should take in the position and size of a rectangular clipping window and clip the objects in the scene against it.

2 Use the GLUT commands discussed in this chapter to set up a display window in which you will display a portion of the animated scene that you developed. More specifically, define a rectangle whose size is moderately smaller than the coordinate extents of all the objects in your scene. This rectangle will act as a clipping window against which you will employ the clipping algorithms you implemented in the previous exercise. The animation should be run continuously, but the objects in the scene should be clipped in each frame against the clipping window, and only this portion of the scene displayed in the display window. Additionally, add the ability to move the clipping window around the scene via keyboard input by using the directional arrows. Each keystroke should move the clipping window by a small amount in the appropriate direction. Run the animation using both the Sutherland-Hodgman and Weiler-Atherton algorithms and note any differences in performance.
Methods for geometric transformations in three dimensions are extended from two-dimensional methods by including considerations for the z coordinate. In most cases, this extension is relatively straightforward. However, in some cases—particularly, rotation—the extension to three dimensions is less obvious.

When we discussed two-dimensional rotations in the $xy$ plane, we needed to consider only rotations about axes that were perpendicular to the $xy$ plane. In three-dimensional space, we can now select any spatial orientation for the rotation axis. Some graphics packages handle three-dimensional rotation as a composite of three rotations, one for each of the three Cartesian axes. Alternatively, we can set up general rotation equations, given the orientation of a rotation axis and the required rotation angle.

A three-dimensional position, expressed in homogeneous coordinates, is represented as a four-element column vector. Thus, each geometric transformation operator is now a $4 \times 4$ matrix, which

premultiplies a coordinate column vector. In addition, as in two dimensions, any sequence of transformations is represented as a single matrix, formed by concatenating the matrices for the individual transformations in the sequence. Each successive matrix in a transformation sequence is concatenated to the left of previous transformation matrices.

**1 Three-Dimensional Translation**

A position \( \mathbf{P} = (x, y, z) \) in three-dimensional space is translated to a location \( \mathbf{P}' = (x', y', z') \) by adding translation distances \( t_x, t_y, \) and \( t_z \) to the Cartesian coordinates of \( \mathbf{P} \):

\[
x' = x + t_x, \quad y' = y + t_y, \quad z' = z + t_z
\]

(1)

Figure 1 illustrates three-dimensional point translation.

We can express these three-dimensional translation operations in matrix form. But now the coordinate positions, \( \mathbf{P} \) and \( \mathbf{P}' \), are represented in homogeneous coordinates with four-element column matrices, and the translation operator \( \mathbf{T} \) is a \( 4 \times 4 \) matrix:

\[
\begin{bmatrix} x' \\ y' \\ z' \\ 1 \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 & t_x \\ 0 & 1 & 0 & t_y \\ 0 & 0 & 1 & t_z \\ 0 & 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} x \\ y \\ z \\ 1 \end{bmatrix}
\]

(2)

or

\[
\mathbf{P}' = \mathbf{T} \cdot \mathbf{P}
\]

(3)

An object is translated in three dimensions by transforming each of the defining coordinate positions for the object, then reconstructing the object at the new location. For an object represented as a set of polygon surfaces, we translate each vertex for each surface (Figure 2) and redisplay the polygon facets at the translated positions.

The following program segment illustrates construction of a translation matrix, given an input set of translation parameters.

![Figure 1](image-url)

*Figure 1*

Moving a coordinate position with translation vector \( \mathbf{T} = (t_x, t_y, t_z) \).
Shifting the position of a three-dimensional object using translation vector $\mathbf{T}$.

>An inverse of a three-dimensional translation matrix is obtained using the same procedures that we applied in a two-dimensional translation. That is, we negate the translation distances $t_x$, $t_y$, and $t_z$. This produces a translation in the opposite direction, and the product of a translation matrix and its inverse is the identity matrix.

## 2 Three-Dimensional Rotation

We can rotate an object about any axis in space, but the easiest rotation axes to handle are those that are parallel to the Cartesian-coordinate axes. Also, we can use combinations of coordinate-axis rotations (along with appropriate translations) to specify a rotation about any other line in space. Therefore, we first consider the operations involved in coordinate-axis rotations, then we discuss the calculations needed for other rotation axes.

By convention, positive rotation angles produce counterclockwise rotations about a coordinate axis, assuming that we are looking in the negative direction along that coordinate axis (Figure 3). This agrees with our earlier discussion of
rotations in two dimensions, where positive rotations in the \( xy \) plane are counterclockwise about a pivot point (an axis that is parallel to the \( z \) axis).

**Three-Dimensional Coordinate-Axis Rotations**

The two-dimensional \( z \)-axis rotation equations are easily extended to three dimensions, as follows:

\[
\begin{align*}
  x' &= x \cos \theta - y \sin \theta \\
  y' &= x \sin \theta + y \cos \theta \\
  z' &= z
\end{align*}
\]  

Parameter \( \theta \) specifies the rotation angle about the \( z \) axis, and \( z \)-coordinate values are unchanged by this transformation. In homogeneous-coordinate form, the three-dimensional \( z \)-axis rotation equations are

\[
\begin{bmatrix}
  x' \\
  y' \\
  z' \\
  1
\end{bmatrix} =
\begin{bmatrix}
  \cos \theta & -\sin \theta & 0 & 0 \\
  \sin \theta & \cos \theta & 0 & 0 \\
  0 & 0 & 1 & 0 \\
  0 & 0 & 0 & 1
\end{bmatrix}
\begin{bmatrix}
  x \\
  y \\
  z \\
  1
\end{bmatrix}
\]  

\[ (5) \]
which we can write more compactly as

$$P' = R_z(\theta) \cdot P \quad (6)$$

Figure 4 illustrates rotation of an object about the z axis.

Transformation equations for rotations about the other two coordinate axes can be obtained with a cyclic permutation of the coordinate parameters $x, y,$ and $z$ in Equations 4:

$$x \rightarrow y \rightarrow z \rightarrow x \quad (7)$$

Thus, to obtain the $x$-axis and $y$-axis rotation transformations, we cyclically replace $x$ with $y$, $y$ with $z$, and $z$ with $x$, as illustrated in Figure 5.

Substituting permutations 7 into Equations 4, we get the equations for an $x$-axis rotation:

$$y' = y \cos \theta - z \sin \theta$$

$$z' = y \sin \theta + z \cos \theta$$

$$x' = x \quad (8)$$

Rotation of an object around the $x$ axis is demonstrated in Figure 6.

A cyclic permutation of coordinates in Equations 8 gives us the transformation equations for a $y$-axis rotation:

$$z' = z \cos \theta - x \sin \theta$$

$$x' = z \sin \theta + x \cos \theta$$

$$y' = y \quad (9)$$

An example of $y$-axis rotation is shown in Figure 7.

An inverse three-dimensional rotation matrix is obtained in the same way as the inverse rotations in two dimensions. We just replace the angle $\theta$ with $-\theta$. 

$$F I G U R E \ 4$$
Rotation of an object about the $z$ axis.

$$F I G U R E \ 5$$
Cyclic permutation of the Cartesian-coordinate axes to produce the three sets of coordinate-axis rotation equations.
General Three-Dimensional Rotations

A rotation matrix for any axis that does not coincide with a coordinate axis can be set up as a composite transformation involving combinations of translations and the coordinate-axis rotations. We first move the designated rotation axis onto one of the coordinate axes. Then we apply the appropriate rotation matrix for that coordinate axis. The last step in the transformation sequence is to return the rotation axis to its original position.

In the special case where an object is to be rotated about an axis that is parallel to one of the coordinate axes, we attain the desired rotation with the following transformation sequence:

1. Translate the object so that the rotation axis coincides with the parallel coordinate axis.
2. Perform the specified rotation about that axis.
3. Translate the object so that the rotation axis is moved back to its original position.

The steps in this sequence are illustrated in Figure 8. A coordinate position \( P \) is transformed with the sequence shown in this figure as

\[
P' = T^{-1} \cdot R_x(\theta) \cdot T \cdot P
\]

where the composite rotation matrix for the transformation is

\[
R(\theta) = T^{-1} \cdot R_x(\theta) \cdot T
\]

This composite matrix is of the same form as the two-dimensional transformation sequence for rotation about an axis that is parallel to the \( z \) axis (a pivot point that is not at the coordinate origin).

When an object is to be rotated about an axis that is not parallel to one of the coordinate axes, we must perform some additional transformations. In this
case, we also need rotations to align the rotation axis with a selected coordinate axis and then to bring the rotation axis back to its original orientation. Given the specifications for the rotation axis and the rotation angle, we can accomplish the required rotation in five steps:

1. Translate the object so that the rotation axis passes through the coordinate origin.
2. Rotate the object so that the axis of rotation coincides with one of the coordinate axes.
3. Perform the specified rotation about the selected coordinate axis.
4. Apply inverse rotations to bring the rotation axis back to its original orientation.
5. Apply the inverse translation to bring the rotation axis back to its original spatial position.

We can transform the rotation axis onto any one of the three coordinate axes. The z-axis is often a convenient choice, and we next consider a transformation sequence using the z-axis rotation matrix (Figure 9).

A rotation axis can be defined with two coordinate positions, as in Figure 10, or with one coordinate point and direction angles (or direction cosines) between the rotation axis and two of the coordinate axes. We assume that the rotation axis is defined by two points, as illustrated, and that the direction of rotation is to be counterclockwise when looking along the axis from \( P_2 \) to \( P_1 \). The components of the rotation-axis vector are then computed as

\[
V = P_2 - P_1 \\
= (x_2 - x_1, y_2 - y_1, z_2 - z_1)
\]  

(12)

The unit rotation-axis vector \( \mathbf{u} \) is

\[
\mathbf{u} = \frac{V}{|V|} = (a, b, c)
\]  

(13)
where the components $a$, $b$, and $c$ are the direction cosines for the rotation axis:

$$
a = \frac{x_2 - x_1}{|V|}, \quad b = \frac{y_2 - y_1}{|V|}, \quad c = \frac{z_2 - z_1}{|V|} \quad (14)$$

If the rotation is to be in the opposite direction (clockwise when viewing from $P_2$ to $P_1$), then we would reverse axis vector $V$ and unit vector $u$ so that they point in the direction from $P_2$ to $P_1$.

The first step in the rotation sequence is to set up the translation matrix that repositions the rotation axis so that it passes through the coordinate origin. Because we want a counterclockwise rotation when viewing along the axis from $P_2$ to $P_1$ (Figure 10), we move the point $P_1$ to the origin. (If the rotation had been specified in the opposite direction, we would move $P_2$ to the origin.) This translation matrix is

$$T = \begin{bmatrix} 1 & 0 & 0 & -x_1 \\ 0 & 1 & 0 & -y_1 \\ 0 & 0 & 1 & -z_1 \\ 0 & 0 & 0 & 1 \end{bmatrix} \quad (15)$$

which repositions the rotation axis and the object as shown in Figure 11.

Next, we formulate the transformations that will put the rotation axis onto the $z$ axis. We can use the coordinate-axis rotations to accomplish this alignment in two steps, and there are a number of ways to perform these two steps. For this example, we first rotate about the $x$ axis, then rotate about the $y$ axis. The $x$-axis rotation gets vector $u$ into the $xz$ plane, and the $y$-axis rotation swings $u$ around to the $z$ axis. These two rotations are illustrated in Figure 12 for one possible orientation of vector $u$.

Because rotation calculations involve sine and cosine functions, we can use standard vector operations to obtain elements of the two rotation matrices. A vector dot product can be used to determine the cosine term, and a vector cross product can be used to calculate the sine term.
We establish the transformation matrix for rotation around the $x$ axis by determining the values for the sine and cosine of the rotation angle necessary to get $u$ into the $xz$ plane. This rotation angle is the angle between the projection of $u$ in the $yz$ plane and the positive $z$ axis (Figure 13). If we represent the projection of $u$ in the $yz$ plane as the vector $u' = (0, b, c)$, then the cosine of the rotation angle $\alpha$ can be determined from the dot product of $u'$ and the unit vector $u_z$ along the $z$ axis:

$$\cos \alpha = \frac{u' \cdot u_z}{|u'| |u_z|} = \frac{c}{d}$$

(16)

where $d$ is the magnitude of $u'$:

$$d = \sqrt{b^2 + c^2}$$

(17)

Similarly, we can determine the sine of $\alpha$ from the cross-product of $u'$ and $u_z$. The coordinate-independent form of this cross-product is

$$u' \times u_z = u_z |u'| |u_z| \sin \alpha$$

(18)

and the Cartesian form for the cross-product gives us

$$u' \times u_z = u_z \cdot b$$

(19)

Equating the right sides of Equations 18 and 19, and noting that $|u| = 1$, and $|u'| = d$, we have

$$d \sin \alpha = b$$

or

$$\sin \alpha = \frac{b}{d}$$

(20)

Now that we have determined the values for $\cos \alpha$ and $\sin \alpha$ in terms of the components of vector $u$, we can set up the matrix elements for rotation of this vector about the $x$ axis and into the $xz$ plane:

$$R_x(\alpha) = \begin{bmatrix} 1 & 0 & 0 & 0 \\ \frac{c}{d} & \frac{b}{d} & -\frac{b}{d} & 0 \\ 0 & \frac{b}{d} & \frac{c}{d} & 0 \\ 0 & \frac{0}{d} & \frac{0}{d} & 1 \end{bmatrix}$$

(21)

The next step in the formulation of the transformation sequence is to determine the matrix that will swing the unit vector in the $xz$ plane counterclockwise around the $y$ axis onto the positive $z$ axis. Figure 14 shows the orientation of
the unit vector in the \(xz\) plane, resulting from the rotation about the \(x\) axis. This vector, labeled \(u''\), has the value \(a\) for its \(x\) component, because rotation about the \(x\) axis leaves the \(x\) component unchanged. Its \(z\) component is \(d\) (the magnitude of \(u'\)), because vector \(u'\) has been rotated onto the \(z\) axis. Also, the \(y\) component of \(u''\) is 0 because it now lies in the \(xz\) plane. Again, we can determine the cosine of rotation angle \(\beta\) from the dot product of unit vectors \(u''\) and \(u_z\). Thus,

\[
\cos \beta = \frac{u'' \cdot u_z}{|u''| |u_z|} = d
\]

because \(|u_z| = |u''| = 1\). Comparing the coordinate-independent form of the cross-product

\[
u'' \times u_z = u_y |u''| |u_z| \sin \beta
\]

with the Cartesian form

\[
u'' \times u_z = u_y \cdot (-a)
\]

we find that

\[
\sin \beta = -a
\]

Therefore, the transformation matrix for rotation of \(u''\) about the \(y\) axis is

\[
R_y(\beta) = \begin{bmatrix}
d & 0 & -a \\
0 & 1 & 0 \\
a & 0 & d \\
0 & 0 & 1
\end{bmatrix}
\]

(26)

With transformation matrices 15, 21, and 26, we have aligned the rotation axis with the positive \(z\) axis. The specified rotation angle \(\theta\) can now be applied as a rotation about the \(z\) axis as follows:

\[
R_z(\theta) = \begin{bmatrix}
\cos \theta & -\sin \theta & 0 & 0 \\
\sin \theta & \cos \theta & 0 & 0 \\
0 & 0 & 1 & 0 \\
0 & 0 & 0 & 1
\end{bmatrix}
\]

(27)

To complete the required rotation about the given axis, we need to transform the rotation axis back to its original position. This is done by applying the inverse of transformations 15, 21, and 26. The transformation matrix for rotation about an arbitrary axis can then be expressed as the composition of these seven individual transformations:

\[
R(\theta) = T^{-1} \cdot R_x^{-1}(\alpha) \cdot R_y^{-1}(\beta) \cdot R_z(\theta) \cdot R_y(\beta) \cdot R_z(\alpha) \cdot T
\]

(28)

A somewhat quicker, but perhaps less intuitive, method for obtaining the composite rotation matrix \(R_y(\beta) \cdot R_z(\alpha)\) is to use the fact that the composite matrix for any sequence of three-dimensional rotations is of the form

\[
R = \begin{bmatrix}
r_{11} & r_{12} & r_{13} & 0 \\
r_{21} & r_{22} & r_{23} & 0 \\
r_{31} & r_{32} & r_{33} & 0 \\
0 & 0 & 0 & 1
\end{bmatrix}
\]

(29)

The upper-left \(3 \times 3\) submatrix of this matrix is orthogonal. This means that the rows (or the columns) of this submatrix form a set of orthogonal unit vectors that
are rotated by matrix $\mathbf{R}$ onto the $x$, $y$, and $z$ axes, respectively:

$$
\mathbf{R} \cdot \begin{bmatrix}
  r_{11} \\
  r_{12} \\
  r_{13} \\
  1
\end{bmatrix} = \begin{bmatrix}
  1 \\
  0 \\
  0 \\
  1
\end{bmatrix}, \\
\mathbf{R} \cdot \begin{bmatrix}
  r_{21} \\
  r_{22} \\
  r_{23} \\
  1
\end{bmatrix} = \begin{bmatrix}
  0 \\
  1 \\
  0 \\
  1
\end{bmatrix}, \\
\mathbf{R} \cdot \begin{bmatrix}
  r_{31} \\
  r_{32} \\
  r_{33} \\
  1
\end{bmatrix} = \begin{bmatrix}
  0 \\
  0 \\
  1 \\
  1
\end{bmatrix}
$$

(30)

Therefore, we can set up a local coordinate system with one of its axes aligned on the rotation axis. Then the unit vectors for the three coordinate axes are used to construct the columns of the rotation matrix. Assuming that the rotation axis is not parallel to any coordinate axis, we could form the following set of local unit vectors (Figure 15).

$$
\begin{align*}
\mathbf{u}_x' &= \mathbf{u} \\
\mathbf{u}_y' &= \frac{\mathbf{u} \times \mathbf{u}_z}{|\mathbf{u} \times \mathbf{u}_z|} \\
\mathbf{u}_z' &= \mathbf{u}_y' \times \mathbf{u}_x'
\end{align*}
$$

(31)

If we express the elements of the unit local vectors for the rotation axis as

$$
\begin{align*}
\mathbf{u}_x' &= (u'_x, u'_y, u'_z) \\
\mathbf{u}_y' &= (u''_x, u''_y, u''_z) \\
\mathbf{u}_z' &= (u'^'_x, u'^'_y, u'^'_z)
\end{align*}
$$

(32)

then the required composite matrix, which is equal to the product $\mathbf{R}_y(\beta) \cdot \mathbf{R}_x(\alpha)$, is

$$
\mathbf{R} = \begin{bmatrix}
  u'_{x1} & u'_{x2} & u'_{x3} & 0 \\
  u'_{y1} & u'_{y2} & u'_{y3} & 0 \\
  u'_{z1} & u'_{z2} & u'_{z3} & 0 \\
  0 & 0 & 0 & 1
\end{bmatrix}
$$

(33)

This matrix transforms the unit vectors $\mathbf{u}_x'$, $\mathbf{u}_y'$, and $\mathbf{u}_z'$ onto the $x$, $y$, and $z$ axes, respectively. This aligns the rotation axis with the $z$ axis, because $\mathbf{u}_z' = \mathbf{u}$.

**Quaternion Methods for Three-Dimensional Rotations**

A more efficient method for generating a rotation about an arbitrarily selected axis is to use a quaternion representation for the rotation transformation. Quaternions, which are extensions of two-dimensional complex numbers, are useful in a number of computer-graphics procedures, including the generation of fractal objects. They require less storage space than $4 \times 4$ matrices, and it is simpler to write quaternion procedures for transformation sequences. This is particularly important in animations, which often require complicated motion sequences and motion interpolations between two given positions of an object.

One way to characterize a quaternion is as an ordered pair, consisting of a scalar part and a vector part:

$$
q = (s, \mathbf{v})
$$

We can also think of a quaternion as a higher-order complex number with one real part (the scalar part) and three complex parts (the elements of vector $\mathbf{v}$). A rotation about any axis passing through the coordinate origin is accomplished by first setting up a unit quaternion with the scalar and vector parts as follows:

$$
\begin{align*}
s &= \cos \frac{\theta}{2}, \\
\mathbf{v} &= \mathbf{u} \sin \frac{\theta}{2}
\end{align*}
$$

(34)
where \( \mathbf{u} \) is a unit vector along the selected rotation axis and \( \theta \) is the specified rotation angle about this axis (Figure 16). Any point position \( \mathbf{P} \) that is to be rotated by this quaternion can be represented in quaternion notation as

\[
\mathbf{P} = (0, \mathbf{p})
\]

with the coordinates of the point as the vector part \( \mathbf{p} = (x, y, z) \). The rotation of the point is then carried out with the quaternion operation

\[
\mathbf{P}' = q \mathbf{P} q^{-1}
\]

where \( q^{-1} = (s, -\mathbf{v}) \) is the inverse of the unit quaternion \( q \) with the scalar and vector parts given in Equations 34. This transformation produces the following new quaternion:

\[
\mathbf{P}' = (0, \mathbf{p}')
\]

The second term in this ordered pair is the rotated point position \( \mathbf{p}' \), which is evaluated with vector dot and cross-products as

\[
\mathbf{p}' = s^2 \mathbf{p} + \mathbf{v} (\mathbf{p} \cdot \mathbf{v}) + 2s (\mathbf{v} \times \mathbf{p}) + (\mathbf{v} \times (\mathbf{v} \times \mathbf{p}))
\]

Values for parameters \( s \) and \( \mathbf{v} \) are obtained from the expressions in 34. Many computer graphics systems use efficient hardware implementations of these vector calculations to perform rapid three-dimensional object rotations.

Transformation 35 is equivalent to rotation about an axis that passes through the coordinate origin. This is the same as the sequence of rotation transformations in Equation 28 that aligns the rotation axis with the \( z \) axis, rotates about \( z \), and then returns the rotation axis to its original orientation at the coordinate origin.

We can evaluate the terms in Equation 37 using the definition for quaternion multiplication. Also, designating the components of the vector part of \( q \) as \( \mathbf{v} = (a, b, c) \), we obtain the elements for the composite rotation matrix \( R_z^{-1}(\alpha) \cdot R_y^{-1}(\beta) \cdot R_z(\theta) \cdot R_y(\beta) \cdot R_z(\alpha) \) in a \( 3 \times 3 \) form as

\[
\begin{bmatrix}
2b^2 - 2c^2 & 2ab - 2sc & a(2c^2 - 2b^2) \\
2ab - 2sc & 2c^2 - 2a^2 & a(1 - 2a^2 - 2c^2) \\
2a^2 - 2sb & 2bc + 2sa & 2a^2 - 2b^2
\end{bmatrix}
\]

The calculations involved in this matrix can be greatly reduced by substituting explicit values for parameters \( a, b, c, \) and \( s \), and then using the following trigonometric identities to simplify the terms:

\[
\cos^2 \frac{\theta}{2} - \sin^2 \frac{\theta}{2} = 1 - 2 \sin^2 \frac{\theta}{2} = \cos \theta, \quad 2 \cos \theta \sin \frac{\theta}{2} = \sin \theta
\]

Thus, we can rewrite Matrix 38 as

\[
M_R(\theta) =
\begin{bmatrix}
u_x (1 - \cos \theta) + \cos \theta & u_x u_y (1 - \cos \theta) - u_z \sin \theta & u_x u_z (1 - \cos \theta) + u_y \sin \theta \\
u_y (1 - \cos \theta) + \cos \theta & u_y u_z (1 - \cos \theta) + \cos \theta & u_y u_x (1 - \cos \theta) - u_z \sin \theta \\
u_z (1 - \cos \theta) - \sin \theta & u_z u_y (1 - \cos \theta) + \sin \theta & u_z u_x (1 - \cos \theta) + \cos \theta
\end{bmatrix}
\]

where \( u_x, u_y, \) and \( u_z \) are the components of the unit axis vector \( \mathbf{u} \).

To complete the transformation sequence for rotating about an arbitrarily placed rotation axis, we need to include the translations that move the rotation axis to the coordinate axis and return it to its original position. Thus, the complete quaternion rotation expression, corresponding to Equation 28, is

\[
R(\theta) = T^{-1} \cdot M_R \cdot T
\]
For example, we can perform a rotation about the z axis by setting rotation-axis vector \( \mathbf{u} \) to the unit z-axis vector \((0, 0, 1)\). Substituting the components of this vector into Matrix 39, we get the \( 3 \times 3 \) version of the z-axis rotation matrix \( \mathbf{R}_z(\theta) \) in Equation 5. Similarly, substituting the unit-quaternion rotation values into Equation 35 produces the rotated coordinate values in Equations 4.

In the following code, we give examples of procedures that could be used to construct a three-dimensional rotation matrix. The quaternion representation in Equation 40 is used to set up the matrix elements for a general three-dimensional rotation.
/* Concatenate translation matrix with matRot. */
    matrix4x4PreMultiply (matTransl3D, matRot);
}

void rotate3D (wcPt3D p1, wcPt3D p2, GLfloat radianAngle)
{
    Matrix4x4 matQuaternionRot;

    GLfloat axisVectLength = sqrt ((p2.x - p1.x) * (p2.x - p1.x) +
            (p2.y - p1.y) * (p2.y - p1.y) +
            (p2.z - p1.z) * (p2.z - p1.z));
    GLfloat cosA = cos (radianAngle);
    GLfloat oneC = 1 - cosA;
    GLfloat sinA = sin (radianAngle);
    GLfloat ux = (p2.x - p1.x) / axisVectLength;
    GLfloat uy = (p2.y - p1.y) / axisVectLength;
    GLfloat uz = (p2.z - p1.z) / axisVectLength;

    /* Set up translation matrix for moving p1 to origin. */
    translate3D (-p1.x, -p1.y, -p1.z);

    /* Initialize matQuaternionRot to identity matrix. */
    matrix4x4SetIdentity (matQuaternionRot);
    matQuaternionRot [0][0] = ux*ux*oneC + cosA;
    matQuaternionRot [0][1] = ux*uy*oneC - uz*sinA;
    matQuaternionRot [0][2] = ux*uz*oneC + uy*sinA;
    matQuaternionRot [1][0] = uy*ux*oneC + uz*sinA;
    matQuaternionRot [1][1] = uy*uy*oneC + cosA;
    matQuaternionRot [1][2] = uy*uz*oneC - ux*sinA;
    matQuaternionRot [2][0] = uz*ux*oneC - uy*sinA;
    matQuaternionRot [2][1] = uz*uy*oneC + ux*sinA;
    matQuaternionRot [2][2] = uz*uz*oneC + cosA;

    /* Combine matQuaternionRot with translation matrix. */
    matrix4x4PreMultiply (matQuaternionRot, matRot);

    /* Set up inverse matTransl3D and concatenate with */
    /* product of previous two matrices. */
    translate3D (p1.x, p1.y, p1.z);
}

void displayFcn (void)
{
    /* Input rotation parameters. */

    /* Initialize matRot to identity matrix: */
    matrix4x4SetIdentity (matRot);

    /* Pass rotation parameters to procedure rotate3D. */

    /* Display rotated object. */
}
3 Three-Dimensional Scaling

The matrix expression for the three-dimensional scaling transformation of a position \( P = (x, y, z) \) relative to the coordinate origin is a simple extension of two-dimensional scaling. We just include the parameter for \( z \)-coordinate scaling in the transformation matrix:

\[
\begin{bmatrix}
  x' \\
  y' \\
  z'
\end{bmatrix}
= \begin{bmatrix}
  s_x & 0 & 0 \\
  0 & s_y & 0 \\
  0 & 0 & s_z
\end{bmatrix}
\begin{bmatrix}
  x \\
  y \\
  z
\end{bmatrix}
\]  

(41)

The three-dimensional scaling transformation for a point position can be represented as

\[ P' = S \cdot P \]  

(42)

where scaling parameters \( s_x, s_y, \) and \( s_z \) are assigned any positive values. Explicit expressions for the scaling transformation relative to the origin are

\[ x' = x \cdot s_x, \quad y' = y \cdot s_y, \quad z' = z \cdot s_z \]  

(43)

Scaling an object with transformation 41 changes the position of the object relative to the coordinate origin. A parameter value greater than 1 moves a point farther from the origin in the corresponding coordinate direction. Similarly, a parameter value less than 1 moves a point closer to the origin in that coordinate direction. Also, if the scaling parameters are not all equal, relative dimensions of a transformed object are changed. We preserve the original shape of an object with a uniform scaling: \( s_x = s_y = s_z \). The result of scaling an object uniformly, with each scaling parameter set to 2, is illustrated in Figure 17.

Because some graphics packages provide only a routine that scales relative to the coordinate origin, we can always construct a scaling transformation with respect to any selected fixed position \((x_f, y_f, z_f)\) using the following transformation sequence:

1. Translate the fixed point to the origin.
2. Apply the scaling transformation relative to the coordinate origin using Equation 41.
3. Translate the fixed point back to its original position.

This sequence of transformations is demonstrated in Figure 18. The matrix representation for an arbitrary fixed-point scaling can then be expressed as the
concatenation of these translate-scale-translate transformations:

\[
\begin{bmatrix}
s_x & 0 & 0 & (1 - s_x)x_f \\
0 & s_y & 0 & (1 - s_y)y_f \\
0 & 0 & s_z & (1 - s_z)z_f \\
0 & 0 & 0 & 1
\end{bmatrix}
\]  

(44)

We can set up programming procedures for constructing a three-dimensional scaling matrix using either a translate-scale-translate sequence or a direct incorporation of the fixed-point coordinates. In the following code example, we demonstrate a direct construction of a three-dimensional scaling matrix relative to a selected fixed point using the calculations in Equation 44:

```cpp
class wcPt3D {
private:
    GLfloat x, y, z;

public:
    /* Default Constructor: */
    wcPt3D ( ) {
        x = y = z = 0.0;
    }

    void setCoords (GLfloat xCoord, GLfloat yCoord, GLfloat zCoord) {
        x = xCoord;
        y = yCoord;
        z = zCoord;
    }

    GLfloat getx ( ) const {
        return x;
    }

    GLfloat gety ( ) const {
        return y;
    }

    GLfloat getz ( ) const {
        return z;
    }
};

typedef float Matrix4x4 [4][4];

void scale3D (GLfloat sx, GLfloat sy, GLfloat sz, wcPt3D fixedPt) {
    Matrix4x4 matScale3D;

    /* Initialize scaling matrix to identity. */
    matrix4x4SetIdentity (matScale3D);
```

**FIGURE 18**
A sequence of transformations for scaling an object relative to a selected fixed point, using Equation 41.
An inverse, three-dimensional scaling matrix is set up for either Equation 41 or Equation 44 by replacing each scaling parameter \((s_x, s_y, s_z)\) with its reciprocal. However, this inverse transformation is undefined if any scaling parameter is assigned the value 0. The inverse matrix generates an opposite scaling transformation, and the concatenation of a three-dimensional scaling matrix with its inverse yields the identity matrix.

### 4 Composite Three-Dimensional Transformations

As with two-dimensional transformations, we form a composite three-dimensional transformation by multiplying the matrix representations for the individual operations in the transformation sequence. Any of the two-dimensional transformation sequences, such as scaling in noncoordinate directions, can be carried out in three-dimensional space.

We can implement a transformation sequence by concatenating the individual matrices from right to left or from left to right, depending on the order in which the matrix representations are specified. Of course, the rightmost term in a matrix product is always the first transformation to be applied to an object and the leftmost term is always the last transformation. We need to use this ordering for the matrix product because coordinate positions are represented as four-element column vectors, which are premultiplied by the composite \(4 \times 4\) transformation matrix.

The following program provides example routines for constructing a three-dimensional composite transformation matrix. The three basic geometric transformations are combined in a selected order to produce a single composite matrix, which is initialized to the identity matrix. For this example, we first rotate, then scale, then translate. We choose a left-to-right evaluation of the composite matrix so that the transformations are called in the order that they are to be applied. Thus, as each matrix is constructed, it is concatenated on the left of the current composite matrix to form the updated product matrix.

```cpp
class wcPt3D {
    public:
        GLfloat x, y, z;
};
typedef GLfloat Matrix4x4 [4][4];
Matrix4x4 matComposite;

/* Construct the 4 x 4 identity matrix. */
void matrix4x4SetIdentity (Matrix4x4 matIdent4x4)
```
{ GLint row, col;
  for (row = 0; row < 4; row++)
    for (col = 0; col < 4; col++)
      matIdent4x4[row][col] = (row == col);
}

/* Premultiply matrix m1 by matrix m2, store result in m2. */
void matrix4x4PreMultiply (Matrix4x4 m1, Matrix4x4 m2)
{
  GLint row, col;
  Matrix4x4 matTemp;
  for (row = 0; row < 4; row++)
    for (col = 0; col < 4; col++)
      matTemp[row][col] = m1[row][0] * m2[0][col] + m1[row][1] * m2[1][col] + m1[row][2] * m2[2][col] + m1[row][3] * m2[3][col];
  for (row = 0; row < 4; row++)
    for (col = 0; col < 4; col++)
      m2[row][col] = matTemp[row][col];
}

/* Procedure for generating 3-D translation matrix. */
void translate3D (GLfloat tx, GLfloat ty, GLfloat tz)
{
  Matrix4x4 matTransl3D;
  /* Initialize translation matrix to identity. */
  matrix4x4SetIdentity (matTransl3D);
  matTransl3D[0][3] = tx;
  matTransl3D[1][3] = ty;
  matTransl3D[2][3] = tz;
  /* Concatenate matTransl3D with composite matrix. */
  matrix4x4PreMultiply (matTransl3D, matComposite);
}

/* Procedure for generating a quaternion rotation matrix. */
void rotate3D (wcPt3D p1, wcPt3D p2, GLfloat radianAngle)
{
  Matrix4x4 matQuatRot;
  float axisVectLength = sqrt ((p2.x - p1.x) * (p2.x - p1.x) +
                              (p2.y - p1.y) * (p2.y - p1.y) +
                              (p2.z - p1.z) * (p2.z - p1.z));
  float cosA = cosf (radianAngle);
  float oneC = 1 - cosA;
  float sinA = sinf (radianAngle);
  float ux = (p2.x - p1.x) / axisVectLength;
  float uy = (p2.y - p1.y) / axisVectLength;
  float uz = (p2.z - p1.z) / axisVectLength;
  /* Set up translation matrix for moving p1 to origin,
and concatenate translation matrix with matComposite.
*/
translate3D (-p1.x, -p1.y, -p1.z);

/* Initialize matQuatRot to identity matrix. */
matrix4x4SetIdentity (matQuatRot);

matQuatRot [0][0] = ux*ux*oneC + cosA;
matQuatRot [0][1] = ux*uy*oneC - uz*sinA;
matQuatRot [0][2] = ux*uz*oneC + uy*sinA;
matQuatRot [1][0] = uy*ux*oneC + uz*sinA;
matQuatRot [1][1] = uy*uy*oneC + cosA;
matQuatRot [1][2] = uy*uz*oneC - ux*sinA;
matQuatRot [2][0] = uz*ux*oneC - uy*sinA;
matQuatRot [2][1] = uz*uy*oneC + ux*sinA;
matQuatRot [2][2] = uz*uz*oneC + cosA;

/* Concatenate matQuatRot with composite matrix. */
matrix4x4PreMultiply (matQuatRot, matComposite);

/* Construct inverse translation matrix for p1 and
* concatenate with composite matrix. */
translate3D (p1.x, p1.y, p1.z);

/* Procedure for generating a 3-D scaling matrix. */
void scale3D (GLfloat sx, GLfloat sy, GLfloat sz, wcPt3D fixedPt)
{
    Matrix4x4 matScale3D;

    /* Initialize scaling matrix to identity. */
    matrix4x4SetIdentity (matScale3D);

    matScale3D [0][0] = sx;
    matScale3D [0][3] = (1 - sx) * fixedPt.x;
    matScale3D [1][1] = sy;
    matScale3D [1][3] = (1 - sy) * fixedPt.y;
    matScale3D [2][2] = sz;
    matScale3D [2][3] = (1 - sz) * fixedPt.z;

    /* Concatenate matScale3D with composite matrix. */
    matrix4x4PreMultiply (matScale3D, matComposite);
}

void displayFcn (void)
{
    /* Input object description. */
    /* Input translation, rotation, and scaling parameters. */

    /* Set up 3-D viewing-transformation routines. */

    /* Initialize matComposite to identity matrix. */
    matrix4x4SetIdentity (matComposite);

    /* Invoke transformation routines in the order they
Three-Dimensional Geometric Transformations
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5 Other Three-Dimensional Transformations

In addition to translation, rotation, and scaling, the other transformations discussed for two-dimensional applications are also useful in many three-dimensional situations. These additional transformations include reflection, shear, and transformations between coordinate-reference frames.

Three-Dimensional Reflections

A reflection in a three-dimensional space can be performed relative to a selected reflection axis or with respect to a reflection plane. In general, three-dimensional reflection matrices are set up similarly to those for two dimensions. Reflections relative to a given axis are equivalent to 180° rotations about that axis. Reflections with respect to a plane are similar; when the reflection plane is a coordinate plane (xy, xz, or yz), we can think of the transformation as a 180° rotation in four-dimensional space with a conversion between a left-handed frame and a right-handed frame.

An example of a reflection that converts coordinate specifications from a right-handed system to a left-handed system (or vice versa) is shown in Figure 19. This transformation changes the sign of z coordinates, leaving the values for the x and y coordinates unchanged. The matrix representation for this reflection relative to the xy plane is

\[
M_{\text{reflect}} = \begin{bmatrix}
1 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 \\
0 & 0 & -1 & 0 \\
0 & 0 & 0 & 1
\end{bmatrix}
\] (45)

Transformation matrices for inverting x coordinates or y coordinates are defined similarly, as reflections relative to the yz plane or to the xz plane, respectively. Reflections about other planes can be obtained as a combination of rotations and coordinate-plane reflections.
Three-Dimensional Shears

These transformations can be used to modify object shapes, just as in two-
dimensional applications. They are also applied in three-dimensional viewing
transformations for perspective projections. For three-dimensional applications,
we can also generate shears relative to the z axis.

A general z-axis shearing transformation relative to a selected reference
position is produced with the following matrix:

\[
M_{\text{shear}} = \begin{bmatrix}
1 & 0 & \text{sh}_{zx} & -\text{sh}_{zx} \cdot z_{\text{ref}} \\
0 & 1 & \text{sh}_{zy} & -\text{sh}_{zy} \cdot z_{\text{ref}} \\
0 & 0 & 1 & 0 \\
0 & 0 & 0 & 1
\end{bmatrix}
\] (46)

Shearing parameters \(\text{sh}_{zx}\) and \(\text{sh}_{zy}\) can be assigned any real values. The effect of
this transformation matrix is to alter the values for the \(x\) and \(y\) coordinates by an
amount that is proportional to the distance from \(z_{\text{ref}}\), while leaving the \(z\) coordinate
unchanged. Plane areas that are perpendicular to the \(z\) axis are thus shifted by an
amount equal to \(z - z_{\text{ref}}\). An example of the effect of this shearing matrix on a unit
cube is shown in Figure 20 for shearing values \(\text{sh}_{zx} = \text{sh}_{zy} = 1\) and a reference
position \(z_{\text{ref}} = 0\). Three-dimensional transformation matrices for an \(x\)-axis shear
and a \(y\)-axis shear are similar to the two-dimensional matrices. We just need to
add a row and a column for the \(z\)-coordinate shearing parameters.

6 Transformations between
Three-Dimensional Coordinate Systems

Coordinate-system transformations are employed in computer-graphics pack-
ages to construct (model) scenes and to implement viewing routines for both
two-dimensional and three-dimensional applications. A transformation matrix for
transferring a two-dimensional scene description from one coordinate system to
another is constructed with operations for superimposing the coordinate axes of
the two systems. The same procedures apply to three-dimensional scene transfers.

We again consider only Cartesian reference frames, and we assume that an
\(x' y' z'\) system is defined with respect to an \(xyz\) system. To transfer the \(xyz\) coor-
dinate descriptions to the \(x' y' z'\) system, we first set up a translation that brings
the \(x' y' z'\) coordinate origin to the position of the \(xyz\) origin. This is followed by a
sequence of rotations that align corresponding coordinate axes. If different scales
are used in the two coordinate systems, a scaling transformation may also be
necessary to compensate for the differences in coordinate intervals.

Figure 21 shows an \(x' y' z'\) coordinate system with origin \((x_0, y_0, z_0)\) and unit
axis vectors defined relative to an \(xyz\) reference frame. The coordinate origin of

F I G U R E 2 0
A unit cube (a) is sheared relative to the origin (b) by Matrix 46, with
\(\text{sh}_{zx} = \text{sh}_{zy} = 1\).

F I G U R E 2 1
An \(x' y' z'\) coordinate system defined within an \(xyz\) system. A scene description is
transferred to the new coordinate reference using a transformation sequence that
superimposes the \(x' y' z'\) frame on the \(xyz\) axes.
the \(x'y'z'\) system is brought into coincidence with the \(xyz\) origin using the translation matrix \(T(-x_0, -y_0, -z_0)\). Also, we can use the unit axis vectors to form the coordinate-axis rotation matrix

\[
R = \begin{bmatrix}
  u'_{x1} & u'_{x2} & u'_{x3} & 0 \\
  u'_{y1} & u'_{y2} & u'_{y3} & 0 \\
  u'_{z1} & u'_{z2} & u'_{z3} & 0 \\
  0 & 0 & 0 & 1
\end{bmatrix}
\]

(47)

which transforms unit vectors \(u'_x, u'_y,\) and \(u'_z\) onto the \(x, y,\) and \(z\) axes, respectively. The complete coordinate transformation sequence is then given by the composite matrix \(R \cdot T\). This matrix correctly transforms coordinate descriptions from one Cartesian system to another, even if one system is left-handed and the other is right-handed.

### 7 Affine Transformations

A coordinate transformation of the form

\[
\begin{align*}
x' &= a_{xx}x + a_{xy}y + a_{xz}z + b_x \\
y' &= a_{yx}x + a_{yy}y + a_{yz}z + b_y \\
z' &= a_{zx}x + a_{zy}y + a_{zz}z + b_z
\end{align*}
\]

(48)

is called an **affine transformation**. Each of the transformed coordinates \(x', y',\) and \(z'\) is a linear function of the original coordinates \(x, y,\) and \(z\), and parameters \(a_{ij}\) and \(b_k\) are constants determined by the transformation type. Affine transformations (in two dimensions, three dimensions, or higher dimensions) have the general properties that parallel lines are transformed into parallel lines, and finite points map to finite points.

Translation, rotation, scaling, reflection, and shear are examples of affine transformations. We can always express any affine transformation as some composition of these five transformations. Another example of an affine transformation is the conversion of coordinate descriptions for a scene from one reference system to another because this transformation can be described as a combination of translation and rotation. An affine transformation involving only translation, rotation, and reflection preserves angles and lengths, as well as parallel lines. For each of these three transformations, line lengths and the angle between any two lines remain the same after the transformation.

### 8 OpenGL Geometric-Transformation Functions

The basic OpenGL functions for performing geometric transformations are the same functions used to perform transformations in three dimensions. For convenience, those functions are listed in Table 1 at the end of the chapter.

**OpenGL Matrix Stacks**

You are already familiar with the OpenGL modelview mode. This mode is selected with the `glMatrixMode` routine and is used to select the modelview...
For each of the four modes (modelview, projection, texture, and color) that we can select with the glMatrixMode function, OpenGL maintains a matrix stack. Initially, each stack contains only the identity matrix. At any time during the processing of a scene, the top matrix on each stack is called the “current matrix” for that mode. After we specify the viewing and geometric transformations, the top of the modelview matrix stack is the $4 \times 4$ composite matrix that combines the viewing transformations and the various geometric transformations that we want to apply to a scene. In some cases, we may want to create multiple views and transformation sequences, and then save the composite matrix for each. Therefore, OpenGL supports a modelview stack depth of at least 32, and some implementations may allow more than 32 matrices to be saved on the modelview stack. We can determine the number of positions available in the modelview stack for a particular implementation of OpenGL with

$$\text{glGetIntegerv (GL_MAX_MODELVIEW_STACK_DEPTH, stackSize);}$$

which returns a single integer value to array stackSize. The other three matrix modes have a minimum stack depth of 2, and we can determine the maximum available depth of each for a particular implementation using one of the following OpenGL symbolic constants: GL_MAX_PROJECTION_STACK_DEPTH, GL_MAX_TEXTURE_STACK_DEPTH, or GL_MAX_COLOR_STACK_DEPTH.

We can also find out how many matrices are currently in the stack with

$$\text{glGetIntegerv (GL_MODELVIEW_STACK_DEPTH, numMats);}$$

Initially, the modelview stack contains only the identity matrix, so the value 1 is returned by this function if we issue the query before any stack processing has occurred. Similar symbolic constants are available for determining the number of matrices currently in the other three stacks.

We have two functions available in OpenGL for processing the matrices in a stack. These stack-processing functions are more efficient than manipulating the stack matrices individually, particularly when the stack functions are implemented in hardware. For example, a hardware implementation can copy multiple matrix elements simultaneously. And we can maintain an identity matrix on the stack, so that initializations of the current matrix can be performed faster than by using repeated calls to glLoadIdentity.

With the following function, we copy the current matrix at the top of the active stack and store that copy in the second stack position:

$$\text{glPushMatrix ( );}$$

This gives us duplicate matrices at the top two positions of the stack. The other stack function is

$$\text{glPopMatrix ( );}$$

which destroys the matrix at the top of the stack, and the second matrix in the stack becomes the current matrix. To “pop” the top of the stack, there must be at least two matrices in the stack. Otherwise, we generate an error.
Usually, it is more efficient to use the stack-processing functions than to use the matrix-manipulation functions. This is particularly true when we want to make several changes in the viewing or geometric transformations. In the following code, we perform rectangle transformations using stack processing instead of the glLoadIdentity function:

```c
glMatrixMode (GL_MODELVIEW);

glColor3f (0.0, 0.0, 1.0); // Set current color to blue.
glRecti (50, 100, 200, 150); // Display blue rectangle.

glPushMatrix ( ); // Make copy of identity (top) matrix.

glColor3f (1.0, 0.0, 0.0); // Set current color to red.

glTranslatef (-200.0, -50.0, 0.0); // Set translation parameters.

glRecti (50, 100, 200, 150); // Display red, translated rectangle.

glPopMatrix ( ); // Throw away the translation matrix.

glPushMatrix ( ); // Make copy of identity (top) matrix.

glRotatef (90.0, 0.0, 0.0, 1.0); // Set 90-deg. rotation about z axis.

glRecti (50, 100, 200, 150); // Display red, rotated rectangle.

glPopMatrix ( ); // Throw away the rotation matrix.

glScalef (-0.5, 1.0, 1.0); // Set scale-reflection parameters.

glRecti (50, 100, 200, 150); // Display red, transformed rectangle.
```

For our next geometric-transformation programming example, we give an OpenGL version of the three-dimensional, composite-transformation code in Section 4. Because OpenGL postmultiplies transformation matrices as they are called, we must now invoke the transformations in the opposite order from which they are to be applied. Thus, each subsequent transformation call concatenates the designated transformation matrix on the right of the composite matrix. Because we have not yet explored the three-dimensional OpenGL viewing routines, this program could be completed using two-dimensional OpenGL viewing operations and applying the geometric transformations to objects in the xy plane.

```c
class wcPt3D {
public:
    GLfloat x, y, z;
};

/* Procedure for generating a matrix for rotation about
```
void rotate3D (wcPt3D p1, wcPt3D p2, GLfloat thetaDegrees)
{
    /* Set up components for rotation-axis vector. */
    float vx = (p2.x - p1.x);
    float vy = (p2.y - p1.y);
    float vz = (p2.z - p1.z);

    /* Specify translate-rotate-translate sequence in reverse order: */
    glTranslatef (p1.x, p1.y, p1.z); // Move p1 back to original position.
    /* Rotate about axis through origin: */
    glRotatef (thetaDegrees, vx, vy, vz);
    glTranslatef (-p1.x, -p1.y, -p1.z); // Translate p1 to origin.
}

void scale3D (GLfloat sx, GLfloat sy, GLfloat sz, wcPt3D fixedPt)
{
    /* Specify translate-scale-translate sequence in reverse order: */
    /* (3) Translate fixed point back to original position: */
    glTranslatef (fixedPt.x, fixedPt.y, fixedPt.z);
    glScalef (sx, sy, sz); // (2) Scale with respect to origin.
    /* (1) Translate fixed point to coordinate origin: */
    glTranslatef (-fixedPt.x, -fixedPt.y, -fixedPt.z);
}

void displayFcn (void)
{
    /* Input object description. */
    /* Set up 3D viewing-transformation routines. */
    /* Display object. */
    glMatrixMode (GL_MODELVIEW);

    /* Input translation parameters tx, ty, tz. */
    /* Input the defining points, p1 and p2, for the rotation axis. */
    /* Input rotation angle in degrees. */
    /* Input scaling parameters: sx, sy, sz, and fixedPt. */

    /* Invoke geometric transformations in reverse order: */
    glTranslatef (tx, ty, tz); // Final transformation: Translate.
    scale3D (sx, sy, sz, fixedPt); // Second transformation: Scale.
    rotate3D (p1, p2, thetaDegrees); // First transformation: Rotate.

    /* Call routines for displaying transformed objects. */
}

10 Summary

We can express three-dimensional transformations as $4 \times 4$ matrix operators, so that sequences of transformations can be concatenated into a single composite
Three-Dimensional Geometric Transformations

### TABLE 1
Summary of OpenGL Geometric Transformation Functions

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glTranslate*</td>
<td>Specifies translation parameters.</td>
</tr>
<tr>
<td>glRotate*</td>
<td>Specifies parameters for rotation about any axis through the origin.</td>
</tr>
<tr>
<td>glScale*</td>
<td>Specifies scaling parameters with respect to coordinate origin.</td>
</tr>
<tr>
<td>glMatrixMode</td>
<td>Specifies current matrix for geometric-viewing transformations, projection transformations, texture transformations, or color transformations.</td>
</tr>
<tr>
<td>glLoadIdentity</td>
<td>Sets current matrix to identity.</td>
</tr>
<tr>
<td>glLoadMatrix* (elems);</td>
<td>Sets elements of current matrix.</td>
</tr>
<tr>
<td>glMultMatrix* (elems);</td>
<td>Postmultiplies the current matrix by the specified matrix.</td>
</tr>
<tr>
<td>glGetIntegerv</td>
<td>Gets max stack depth or current number of matrices in the stack for the selected matrix mode.</td>
</tr>
<tr>
<td>glPushMatrix</td>
<td>Copies the top matrix in the stack and store copy in the second stack position.</td>
</tr>
<tr>
<td>glPopMatrix</td>
<td>Erases the top matrix in the stack and moves the second matrix to the top of the stack.</td>
</tr>
<tr>
<td>glPixelZoom</td>
<td>Specifies two-dimensional scaling parameters for raster operations.</td>
</tr>
</tbody>
</table>

We use a four-element column matrix (vector) representation for three-dimensional coordinate points, representing them using a homogeneous coordinate representation.

We can create composite transformations through matrix multiplications of translation, rotation, scaling, and other transformations. In general, matrix multiplications are not commutative. The upper-left $3 \times 3$ submatrix of a rigid-body transformation is an orthogonal matrix. Thus, rotation matrices can be formed by setting the upper-left, $3 \times 3$ submatrix equal to the elements of two orthogonal unit vectors. When the angle is small, we can reduce rotation computations by using first-order approximations for the sine and cosine functions. Over many rotational steps, however, the approximation error can accumulate to a significant value.

Transformations between Cartesian coordinate systems in three dimensions are accomplished with a sequence of translate-rotate transformations that brings the two systems into coincidence. We specify the coordinate origin and axis vectors for one reference frame relative to the original coordinate reference frame. The transfer of object descriptions from the original coordinate system to the second system is calculated as the matrix product of a translation that moves the new origin to the old coordinate origin and a rotation to align the two sets of axes. The rotation needed to align the two frames can be obtained from the orthonormal set of axis vectors for the new system.
Three-Dimensional Geometric Transformations

The OpenGL library provides functions for applying individual translate, rotate, and scale transformations to coordinate positions. Each function generates a matrix that is premultiplied by the modelview matrix. Transformation matrices are applied to subsequently defined objects. In addition to accumulating transformation sequences in the modelview matrix, we can set this matrix to the identity or some other matrix, and can also form products with the modelview matrix and any specified matrices. All matrices are stored in stacks, and OpenGL maintains four stacks for the various types of transformations that we use in graphics applications. We can use an OpenGL query function to determine the current stack size or the maximum allowable stack depth for a system. Two stack-processing routines are available: one for copying the top matrix in a stack to the second position, and one for removing the top matrix.

Table 1 summarizes the OpenGL geometric-transformation functions and matrix routines discussed in this chapter.

REFERENCES

Additional programming examples using OpenGL geometric-transformation functions are given in Woo, et al. (1999). Programming examples for the OpenGL geometric-transformation functions are also available at Nate Robins’s tutorial website: http://www.xmission.com/~nate/opengl.html. Finally, a complete listing of OpenGL geometric-transformation functions is provided in Shreiner (2000).

EXERCISES
1 Show that rotation matrix 33 is equal to the composite matrix \( R_y(\beta) \cdot R_z(\alpha) \).
2 By evaluating the terms in Equation 37, derive the elements for the general rotation matrix given in Equation 38.
3 Prove that the quaternion rotation matrix 38 reduces to the matrix representation in Equation 5 when the rotation axis is the coordinate z axis.
4 Prove that Equation 40 is equivalent to the general rotation transformation given in Equation 28.
5 Using trigonometric identities, derive the elements of the quaternion-rotation matrix 39 from 38.
6 Develop a procedure for animating a three-dimensional object by incrementally rotating it about any specified axis. Use appropriate approximations to the trigonometric equations to speed up the calculations, and reset the object to its initial position after each complete revolution about the axis.
7 Derive the three-dimensional transformation matrix for scaling an object by a scaling factor \( s \) in a direction defined by the direction cosines \( \alpha, \beta, \) and \( \gamma \).
8 Develop a routine to reflect a three-dimensional object about an arbitrarily selected plane.
9 Write a procedure to shear a three-dimensional object with respect to any specified axis, using input values for the shearing parameters.
10 Develop a procedure for converting an object definition in one three-dimensional coordinate reference to any other coordinate system defined relative to the first system.
11 Develop a routine to scale an object by a given factor in each dimension relative to a given point contained within the object.
12 Write a program to perform a series of transformations on a 30 × 30 square whose centroid lies at (−20, −20, 0) and that is contained in the xy plane. Use three-dimensional OpenGL matrix operations to perform the transformations. The square should first be reflected in the x axis, then rotated counterclockwise by 45° about its center, then sheared in the x direction by a value of 2.

13 Modify the program from the previous exercise so that the transformation sequence can be applied to any two-dimensional polygon, with vertices specified as user input.

14 Modify the example program in the previous exercise so that the order of the geometric transformation sequence can be specified as user input.

15 Modify the example program from the previous exercise so that the geometric transformation parameters are specified as user input.

**IN MORE DEPTH**

1 You have not yet been exposed to the material necessary to construct three-dimensional representations of the objects in your application, so you can instead embed the two-dimensional polygonal approximations to those objects in a three-dimensional scene and perform three-dimensional transformations on those approximations using the techniques in this chapter. In this exercise, you will set up a set of transformations to produce an animation. Define the three-dimensional transformation matrices to do this using homogeneous coordinate representations. If two or more objects act as a single “unit” in certain behaviors that are easier to model in terms of relative positions, you can use the techniques in Section 6 to convert the local transformations of the objects relative to each other (in their own coordinate frame) into transformations in the world coordinate frame.

2 Use the matrices you designed in the previous exercise to produce an animation. You should employ the OpenGL matrix operations for three-dimensional transformations and have the matrices produce small changes in position for each of the objects in the scene. Since you haven’t covered the material necessary for generating views of a three-dimensional scene, simply display the animation using a two-dimensional orthogonal projection, with all of the polygons in the scene being contained in the xy plane. The transformations themselves, however, are still three-dimensional.
For two-dimensional graphics applications, viewing operations transfer positions from the world-coordinate plane to pixel positions in the plane of the output device. Using the rectangular boundaries for the clipping window and the viewport, a two-dimensional package clips a scene and maps it to device coordinates. Three-dimensional viewing operations, however, are more involved, because we now have many more choices as to how we can construct a scene and how we can generate views of the scene on an output device.
1 Overview of Three-Dimensional Viewing Concepts

When we model a three-dimensional scene, each object in the scene is typically defined with a set of surfaces that form a closed boundary around the object interior. And, for some applications, we may need also to specify information about the interior structure of an object. In addition to procedures that generate views of the surface features of an object, graphics packages sometimes provide routines for displaying internal components or cross-sectional views of a solid object. Viewing functions process the object descriptions through a set of procedures that ultimately project a specified view of the objects onto the surface of a display device. Many processes in three-dimensional viewing, such as the clipping routines, are similar to those in the two-dimensional viewing pipeline. But three-dimensional viewing involves some tasks that are not present in two-dimensional viewing. For example, projection routines are needed to transfer the scene to a view on a planar surface, visible parts of a scene must be identified, and, for a realistic display, lighting effects and surface characteristics must be taken into account.

Viewing a Three-Dimensional Scene

To obtain a display of a three-dimensional world-coordinate scene, we first set up a coordinate reference for the viewing, or “camera,” parameters. This coordinate reference defines the position and orientation for a view plane (or projection plane) that corresponds to a camera film plane (Figure 1). Object descriptions are then transferred to the viewing reference coordinates and projected onto the view plane. We can generate a view of an object on the output device in wireframe (outline) form, or we can apply lighting and surface-rendering techniques to obtain a realistic shading of the visible surfaces.

Projections

Unlike a camera picture, we can choose different methods for projecting a scene onto the view plane. One method for getting the description of a solid object onto a view plane is to project points on the object surface along parallel lines. This technique, called parallel projection, is used in engineering and architectural drawings to represent an object with a set of views that show accurate dimensions of the object, as in Figure 2.

Another method for generating a view of a three-dimensional scene is to project points to the view plane along converging paths. This process, called perspective projection, causes objects farther from the viewing position to be displayed smaller than objects of the same size that are nearer to the viewing position. A scene that is generated using a perspective projection appears more realistic, because this is the way that our eyes and a camera lens form images. Parallel lines along the viewing direction appear to converge to a distant point in the background, and objects in the background appear to be smaller than objects in the foreground.
Depth Cueing

With few exceptions, depth information is important in a three-dimensional scene so that we can easily identify, for a particular viewing direction, which is the front and which is the back of each displayed object. Figure 3 illustrates the ambiguity that can result when a wire-frame object is displayed without depth information. There are several ways in which we can include depth information in the two-dimensional representation of solid objects.

A simple method for indicating depth with wire-frame displays is to vary the brightness of line segments according to their distances from the viewing position. Figure 4 shows a wire-frame object displayed with depth cueing. The lines closest to the viewing position are displayed with the highest intensity, and lines farther away are displayed with decreasing intensities. Depth cueing is applied by choosing a maximum and a minimum intensity value and a range of distances over which the intensity is to vary.

Another application of depth cueing is modeling the effect of the atmosphere on the perceived intensity of objects. More distant objects appear dimmer to us than nearer objects due to light scattering by dust particles, haze, and smoke. Some atmospheric effects can even change the perceived color of an object, and we can model these effects with depth cueing.

Identifying Visible Lines and Surfaces

We can also clarify depth relationships in a wire-frame display using techniques other than depth cueing. One approach is simply to highlight the visible lines or to display them in a different color. Another technique, commonly used for engineering drawings, is to display the nonvisible lines as dashed lines. Or we could remove the nonvisible lines from the display, as in Figures 3(b) and 3(c). But removing the hidden lines also removes information about the shape of the back surfaces of an object, and wire-frame representations are generally used to get an indication of an object’s overall appearance, front and back.

When a realistic view of a scene is to be produced, back parts of the objects are completely eliminated so that only the visible surfaces are displayed. In this case, surface-rendering procedures are applied so that screen pixels contain only the color patterns for the front surfaces.

Surface Rendering

Added realism is attained in displays by rendering object surfaces using the lighting conditions in the scene and the assigned surface characteristics. We set the lighting conditions by specifying the color and location of the light sources, and we can also set background illumination effects. Surface properties of objects include whether a surface is transparent or opaque and whether the surface is smooth or rough. We set values for parameters to model surfaces such as glass, plastic, wood-grain patterns, and the bumpy appearance of an orange. In Color Plate 9 surface-rendering methods are combined with perspective and visible-surface identification to generate a degree of realism in a displayed scene.

Exploded and Cutaway Views

Many graphics packages allow objects to be defined as hierarchical structures, so that internal details can be stored. Exploded and cutaway views of such objects can then be used to show the internal structure and relationship of the object parts. An alternative to exploding an object into its component parts is a cutaway view, which removes part of the visible surfaces to show internal structure.
Three-Dimensional and Stereoscopic Viewing

Other methods for adding a sense of realism to a computer-generated scene include three-dimensional displays and stereoscopic views. Three-dimensional views can be obtained by reflecting a raster image from a vibrating, flexible mirror. The vibrations of the mirror are synchronized with the display of the scene on the cathode ray tube (CRT). As the mirror vibrates, the focal length varies so that each point in the scene is reflected to a spatial position corresponding to its depth.

Stereoscopic devices present two views of a scene: one for the left eye and the other for the right eye. The viewing positions correspond to the eye positions of the viewer. These two views are typically displayed on alternate refresh cycles of a raster monitor. When we view the monitor through special glasses that alternately darken first one lens and then the other, in synchronization with the monitor refresh cycles, we see the scene displayed with a three-dimensional effect.

2 The Three-Dimensional Viewing Pipeline

Procedures for generating a computer-graphics view of a three-dimensional scene are somewhat analogous to the processes involved in taking a photograph. First of all, we need to choose a viewing position corresponding to where we would place a camera. We choose the viewing position according to whether we want to display a front, back, side, top, or bottom view of the scene. We could also pick a position in the middle of a group of objects or even inside a single object, such as a building or a molecule. Then we must decide on the camera orientation (Figure 5). Which way do we want to point the camera from the viewing position, and how should we rotate it around the line of sight to set the “up” direction for the picture? Finally, when we snap the shutter, the scene is cropped to the size of a selected clipping window, which corresponds to the aperture or lens type of a camera, and light from the visible surfaces is projected onto the camera film.

We need to keep in mind, however, that the camera analogy can be carried only so far, because we have more flexibility and many more options for generating views of a scene with a computer-graphics program than we do with a real camera. We can choose to use either a parallel projection or a perspective projection, we can selectively eliminate parts of a scene along the line of sight, we can move the projection plane away from the “camera” position, and we can even get a picture of objects in back of our synthetic camera.

Some of the viewing operations for a three-dimensional scene are the same as, or similar to, those used in the two-dimensional viewing pipeline. A two-dimensional viewport is used to position a projected view of the three-dimensional scene on the output device, and a two-dimensional clipping window is used to

![Figure 5](image-url)

*Photographing a scene involves selection of the camera position and orientation.*
select a view that is to be mapped to the viewport. In addition, we set up a display window in screen coordinates, just as we do in a two-dimensional application. Clipping windows, viewports, and display windows are usually specified as rectangles with their edges parallel to the coordinate axes. In three-dimensional viewing, however, the clipping window is positioned on a selected view plane, and scenes are clipped against an enclosing volume of space, which is defined by a set of clipping planes. The viewing position, view plane, clipping window, and clipping planes are all specified within the viewing-coordinate reference frame.

Figure 6 shows the general processing steps for creating and transforming a three-dimensional scene to device coordinates. Once the scene has been modeled in world coordinates, a viewing-coordinate system is selected and the description of the scene is converted to viewing coordinates. The viewing coordinate system defines the viewing parameters, including the position and orientation of the projection plane (view plane), which we can think of as the camera film plane. A two-dimensional clipping window, corresponding to a selected camera lens, is defined on the projection plane, and a three-dimensional clipping region is established. This clipping region is called the view volume, and its shape and size depends on the dimensions of the clipping window, the type of projection we choose, and the selected limiting positions along the viewing direction. Projection operations are performed to convert the viewing-coordinate description of the scene to coordinate positions on the projection plane. Objects are mapped to normalized coordinates, and all parts of the scene outside the view volume are clipped off. The clipping operations can be applied after all device-independent coordinate transformations (from world coordinates to normalized coordinates) are completed. In this way, the coordinate transformations can be concatenated for maximum efficiency.

As in two-dimensional viewing, the viewport limits could be given in normalized coordinates or in device coordinates. In developing the viewing algorithms, we will assume that the viewport is to be specified in device coordinates and that normalized coordinates are transferred to viewport coordinates, following the clipping operations. There are also a few other tasks that must be performed, such as identifying visible surfaces and applying the surface-rendering procedures. The final step is to map viewport coordinates to device coordinates within a selected display window. Scene descriptions in device coordinates are sometimes expressed in a left-handed reference frame so that positive distances from the display screen can be used to measure depth values in the scene.
3 Three-Dimensional Viewing-Coordinate Parameters

Establishing a three-dimensional viewing reference frame is similar to setting up the two-dimensional viewing reference frame. We first select a world-coordinate position $P_0 = (x_0, y_0, z_0)$ for the viewing origin, which is called the view point or viewing position. (Sometimes the view point is also referred to as the eye position or the camera position.) And we specify a view-up vector $V$, which defines the $y_{view}$ direction. For three-dimensional space, we also need to assign a direction for one of the remaining two coordinate axes. This is typically accomplished with a second vector that defines the $z_{view}$ axis, with the viewing direction along this axis. Figure 7 illustrates the positioning of a three-dimensional viewing-coordinate frame within a world system.

The View-Plane Normal Vector

Because the viewing direction is usually along the $z_{view}$ axis, the view plane, also called the projection plane, is normally assumed to be perpendicular to this axis. Thus, the orientation of the view plane, as well as the direction for the positive $z_{view}$ axis, can be defined with a view-plane normal vector $N$, as shown in Figure 8.

An additional scalar parameter is used to set the position of the view plane at some coordinate value $z_{vp}$ along the $z_{view}$ axis, as illustrated in Figure 9. This parameter value is usually specified as a distance from the viewing origin along the direction of viewing, which is often taken to be in the negative $z_{view}$ direction. Thus, the view plane is always parallel to the $x_{view}y_{view}$ plane, and the projection of objects to the view plane corresponds to the view of the scene that will be displayed on the output device.

Vector $N$ can be specified in various ways. In some graphics systems, the direction for $N$ is defined to be along the line from the world-coordinate origin to a selected point position. Other systems take $N$ to be in the direction from a reference point $P_{ref}$ to the viewing origin $P_0$, as in Figure 10. In this case, the reference point is often referred to as a look-at point within the scene, with the viewing direction opposite to the direction of $N$.

We could also define the view-plane normal vector, and other vector directions, using direction angles. These are the three angles, $\alpha$, $\beta$, and $\gamma$, that a spatial line makes with the $x$, $y$, and $z$ axes, respectively. But it is usually much easier to specify a vector direction with two point positions in a scene than with direction angles.
Three-Dimensional Viewing

The View-Up Vector

Once we have chosen a view-plane normal vector $N$, we can set the direction for the view-up vector $V$. This vector is used to establish the positive direction for the $y_{\text{view}}$ axis.

Usually, $V$ is defined by selecting a position relative to the world-coordinate origin, so that the direction for the view-up vector is from the world origin to this selected position. Because the view-plane normal vector $N$ defines the direction for the $z_{\text{view}}$ axis, vector $V$ should be perpendicular to $N$. But, in general, it can be difficult to determine a direction for $V$ that is precisely perpendicular to $N$. Therefore, viewing routines typically adjust the user-defined orientation of vector $V$, as shown in Figure 11, so that $V$ is projected onto a plane that is perpendicular to the view-plane normal vector.

We can choose any direction for the view-up vector $V$, so long as it is not parallel to $N$. A convenient choice is often in a direction parallel to the world $y_w$ axis; that is, we could set $V = (0, 1, 0)$.

The $uvw$ Viewing-Coordinate Reference Frame

Left-handed viewing coordinates are sometimes used in graphics packages, with the viewing direction in the positive $z_{\text{view}}$ direction. With a left-handed system, increasing $z_{\text{view}}$ values are interpreted as being farther from the viewing position along the line of sight. But right-handed viewing systems are more common, because they have the same orientation as the world-reference frame. This allows a graphics package to deal with only one coordinate orientation for both world and viewing references. Although some early graphics packages defined viewing coordinates within a left-handed frame, right-handed viewing coordinates are now used by the graphics standards. However, left-handed coordinate references are often used to represent screen coordinates and for the normalization transformation.

Because the view-plane normal $N$ defines the direction for the $z_{\text{view}}$ axis and the view-up vector $V$ is used to obtain the direction for the $y_{\text{view}}$ axis, we need only determine the direction for the $x_{\text{view}}$ axis. Using the input values for $N$ and $V$, we can compute a third vector, $U$, that is perpendicular to both $N$ and $V$. Vector $U$ then defines the direction for the positive $x_{\text{view}}$ axis. We determine the correct direction for $U$ by taking the vector cross product of $V$ and $N$ so as to form a right-handed viewing frame. The vector cross product of $N$ and $U$ also produces the adjusted value for $V$, perpendicular to both $N$ and $U$, along the positive $y_{\text{view}}$ axis. Following these procedures, we obtain the following set of unit axis vectors for a right-handed viewing coordinate system.

\[
\begin{align*}
  n &= \frac{N}{|N|} = (n_x, n_y, n_z) \\
  u &= \frac{V \times n}{|V \times n|} = (u_x, u_y, u_z) \\
  v &= n \times u = (v_x, v_y, v_z)
\end{align*}
\]
The coordinate system formed with these unit vectors is often described as a \textbf{uvn viewing-coordinate reference frame} (Figure 12).

### Generating Three-Dimensional Viewing Effects

By varying the viewing parameters, we can obtain different views of objects in a scene. For instance, from a fixed viewing position, we could change the direction of \( \mathbf{N} \) to display objects at positions around the viewing-coordinate origin. We could also vary \( \mathbf{N} \) to create a composite display consisting of multiple views from a fixed camera position. We can simulate a wide viewing angle by producing seven views of the scene from the same viewing position, but with slight shifts in the viewing direction; the views are then combined to form a composite display. Similarly, we generate stereoscopic views by shifting the viewing direction as well as shifting the view point slightly to simulate the two eye positions.

In interactive applications, the normal vector \( \mathbf{N} \) is the viewing parameter that is most often changed. Of course, when we change the direction for \( \mathbf{N} \), we also have to change the other axis vectors to maintain a right-handed viewing-coordinate system.

If we want to simulate an animation panning effect, as when a camera moves through a scene or follows an object that is moving through a scene, we can keep the direction for \( \mathbf{N} \) fixed as we move the view point, as illustrated in Figure 13. And to display different views of an object, such as a side view and a front view, we could move the view point around the object, as in Figure 14. Alternatively, different views of an object or group of objects can be generated using geometric transformations without changing the viewing parameters.

### 4 Transformation from World to Viewing Coordinates

In the three-dimensional viewing pipeline, the first step after a scene has been constructed is to transfer object descriptions to the viewing-coordinate reference frame. This conversion of object descriptions is equivalent to a sequence of transformations that superimposes the viewing reference frame onto the world frame. We can accomplish this conversion using the methods for transforming between
coordinate systems:

1. Translate the viewing-coordinate origin to the origin of the world-coordinate system.

2. Apply rotations to align the \( x_{\text{view}} \), \( y_{\text{view}} \), and \( z_{\text{view}} \) axes with the world \( x_w \), \( y_w \), and \( z_w \) axes, respectively.

The viewing-coordinate origin is at world position \( P_0 = (x_0, y_0, z_0) \). Therefore, the matrix for translating the viewing origin to the world origin is

\[
T = \begin{bmatrix}
1 & 0 & 0 & -x_0 \\
0 & 1 & 0 & -y_0 \\
0 & 0 & 1 & -z_0 \\
0 & 0 & 0 & 1
\end{bmatrix}
\] (2)

For the rotation transformation, we can use the unit vectors \( u \), \( v \), and \( n \) to form the composite rotation matrix that superimposes the viewing axes onto the world frame. This transformation matrix is

\[
R = \begin{bmatrix}
ux & uy & uz & 0 \\
vx & vy & vz & 0 \\
nx & ny & nz & 0 \\
0 & 0 & 0 & 1
\end{bmatrix}
\] (3)

where the elements of matrix \( R \) are the components of the \( uvn \) axis vectors.

The coordinate transformation matrix is then obtained as the product of the preceding translation and rotation matrices:

\[
M_{WC, VC} = R \cdot T = \begin{bmatrix}
ux & uy & uz & -u \cdot P_0 \\
vx & vy & vz & -v \cdot P_0 \\
nx & ny & nz & -n \cdot P_0 \\
0 & 0 & 0 & 1
\end{bmatrix}
\] (4)
Translation factors in this matrix are calculated as the vector dot product of each of the $u$, $v$, and $n$ unit vectors with $P_0$, which represents a vector from the world origin to the viewing origin. In other words, the translation factors are the negative projections of $P_0$ on each of the viewing-coordinate axes (the negative components of $P_0$ in viewing coordinates). These matrix elements are evaluated as

$$
-u \cdot P_0 = -x_0 u_x - y_0 u_y - z_0 u_z \\
-v \cdot P_0 = -x_0 v_x - y_0 v_y - z_0 v_z \\
-n \cdot P_0 = -x_0 n_x - y_0 n_y - z_0 n_z
$$

(5)

Matrix 4 transfers world-coordinate object descriptions to the viewing reference frame.

5 Projection Transformations

In the next phase of the three-dimensional viewing pipeline, after the transformation to viewing coordinates, object descriptions are projected to the view plane. Graphics packages generally support both parallel and perspective projections.

In a parallel projection, coordinate positions are transferred to the view plane along parallel lines. Figure 15 illustrates a parallel projection for a straight-line segment defined with endpoint coordinates $P_1$ and $P_2$. A parallel projection preserves relative proportions of objects, and this is the method used in computer-aided drafting and design to produce scale drawings of three-dimensional objects. All parallel lines in a scene are displayed as parallel when viewed with a parallel projection. There are two general methods for obtaining a parallel-projection view of an object: We can project along lines that are perpendicular to the view plane, or we can project at an oblique angle to the view plane.

For a perspective projection, object positions are transformed to projection coordinates along lines that converge to a point behind the view plane. An example of a perspective projection for a straight-line segment, defined with endpoint coordinates $P_1$ and $P_2$, is given in Figure 16. Unlike a parallel projection, a perspective projection does not preserve relative proportions of objects. But perspective views of a scene are more realistic because distant objects in the projected display are reduced in size.

6 Orthogonal Projections

A transformation of object descriptions to a view plane along lines that are all parallel to the view-plane normal vector $N$ is called an orthogonal projection (or,
equivalently, an orthographic projection). This produces a parallel-projection transformation in which the projection lines are perpendicular to the view plane. Orthogonal projections are most often used to produce the front, side, and top views of an object, as shown in Figure 17. Front, side, and rear orthogonal projections of an object are called elevations; and a top orthogonal projection is called a plan view. Engineering and architectural drawings commonly employ these orthographic projections, because lengths and angles are accurately depicted and can be measured from the drawings.

Axonometric and Isometric Orthogonal Projections
We can also form orthogonal projections that display more than one face of an object. Such views are called axonometric orthogonal projections. The most commonly used axonometric projection is the isometric projection, which is generated by aligning the projection plane (or the object) so that the plane intersects each coordinate axis in which the object is defined, called the principal axes, at the same distance from the origin. Figure 18 shows an isometric projection for a cube. We can obtain the isometric projection shown in this figure by aligning the view-plane normal vector along a cube diagonal. There are eight positions, one in each octant, for obtaining an isometric view. All three principal axes are foreshortened equally in an isometric projection, so that relative proportions are maintained. This is not the case in a general axonometric projection, where scaling factors may be different for the three principal directions.

Orthogonal Projection Coordinates
With the projection direction parallel to the $z_{\text{view}}$ axis, the transformation equations for an orthogonal projection are trivial. For any position $(x, y, z)$ in viewing coordinates, as in Figure 19, the projection coordinates are

$$x_p = x, \quad y_p = y$$

(6)
Three-Dimensional Viewing

The $z$-coordinate value for any projection transformation is preserved for use in the visibility determination procedures. And each three-dimensional coordinate point in a scene is converted to a position in normalized space.

Clipping Window and Orthogonal-Projection View Volume

In the camera analogy, the type of lens is one factor that determines how much of the scene is transferred to the film plane. A wide-angle lens takes in more of the scene than a regular lens. For computer-graphics applications, we use the rectangular clipping window for this purpose. As in two-dimensional viewing, graphics packages typically require that clipping rectangles be placed in specific positions.

In OpenGL, we set up a clipping window for three-dimensional viewing just as we did for two-dimensional viewing, by choosing two-dimensional coordinate positions for its lower-left and upper-right corners. For three-dimensional viewing, the clipping window is positioned on the view plane with its edges parallel to the $x_{\text{view}}$ and $y_{\text{view}}$ axes, as shown in Figure 20. If we want to use some other shape or orientation for the clipping window, we must develop our own viewing procedures.

The edges of the clipping window specify the $x$ and $y$ limits for the part of the scene that we want to display. These limits are used to form the top, bottom, and two sides of a clipping region called the orthogonal-projection view volume. Because projection lines are perpendicular to the view plane, these four boundaries are planes that are also perpendicular to the view plane and that pass...
Three-Dimensional Viewing

A clipping window on the view plane, with minimum and maximum coordinates given in the viewing reference system.

FIGURE 20

Infinite orthogonal-projection view volume.

FIGURE 21

through the edges of the clipping window to form an infinite clipping region, as in Figure 21.

We can limit the extent of the orthogonal view volume in the $z_{\text{view}}$ direction by selecting positions for one or two additional boundary planes that are parallel to the view plane. These two planes are called the near-far clipping planes, or the front-back clipping planes. The near and far planes allow us to exclude objects that are in front of or behind the part of the scene that we want to display. With the viewing direction along the negative $z_{\text{view}}$ axis, we usually have $z_{\text{far}} < z_{\text{near}}$, so that the far plane is farther out along the negative $z_{\text{view}}$ axis. Some graphics libraries provide these two planes as options, and other libraries require them. When the near and far planes are specified, we obtain a finite orthogonal view volume that is a rectangular parallelepiped, as shown in Figure 22 along with one possible placement for the view plane. Our view of the scene will then contain only those objects within the view volume, with all parts of the scene outside the view volume eliminated by the clipping algorithms.

Graphics packages provide varying degrees of flexibility in the positioning of the near and far clipping planes, including options for specifying additional clipping planes at other positions in the scene. In general, the near and far planes can be in any relative position to each other to achieve various viewing effects, including positions that are on opposite sides of the view point. Similarly, the view plane can sometimes be placed in any position relative to the near and far clipping planes, although it is often taken to be coincident with the near clipping plane. However, providing numerous positioning options for the clipping and view planes usually results in less efficient processing of a three-dimensional scene.
Normalization Transformation for an Orthogonal Projection

Using an orthogonal transfer of coordinate positions onto the view plane, we obtain the projected position of any spatial point \((x, y, z)\) as simply \((x, y)\). Thus, once we have established the limits for the view volume, coordinate descriptions inside this rectangular parallelepiped are the projection coordinates, and they can be mapped into a normalized view volume without any further projection processing. Some graphics packages use a unit cube for this normalized view volume, with each of the \(x\), \(y\), and \(z\) coordinates normalized in the range from 0 to 1. Another normalization-transformation approach is to use a symmetric cube, with coordinates in the range from \(-1\) to 1.

Because screen coordinates are often specified in a left-handed reference frame (Figure 23), normalized coordinates also are often specified in a left-handed system. This allows positive distances in the viewing direction to be directly interpreted as distances from the screen (the viewing plane). Thus, we can convert projection coordinates into positions within a left-handed normalized-coordinate reference frame, and these coordinate positions will then be transferred to left-handed screen coordinates by the viewport transformation.

To illustrate the normalization transformation, we assume that the orthogonal-projection view volume is to be mapped into the symmetric...
normalization cube within a left-handed reference frame. Also, z-coordinate positions for the near and far planes are denoted as $z_{\text{near}}$ and $z_{\text{far}}$, respectively. Figure 24 illustrates this normalization transformation. Position $(x_{\text{min}}, y_{\text{min}}, z_{\text{near}})$ is mapped to the normalized position $(-1, -1, -1)$, and position $(x_{\text{max}}, y_{\text{max}}, z_{\text{far}})$ is mapped to $(1, 1, 1)$.

Transforming the rectangular-parallelepiped view volume to a normalized cube is similar to the methods for converting the clipping window into the normalized symmetric square. The normalization transformation for the orthogonal view volume is

$$
M_{\text{ortho\_norm}} = \begin{bmatrix}
\frac{2}{x_{\text{max}} - x_{\text{min}}} & 0 & 0 & -\frac{x_{\text{wmax}} + x_{\text{wmin}}}{x_{\text{max}} - x_{\text{min}}} \\
0 & \frac{2}{y_{\text{max}} - y_{\text{min}}} & 0 & -\frac{y_{\text{wmax}} + y_{\text{wmin}}}{y_{\text{max}} - y_{\text{min}}} \\
0 & 0 & \frac{-2}{z_{\text{near}} - z_{\text{far}}} & \frac{z_{\text{near}} + z_{\text{far}}}{z_{\text{near}} - z_{\text{far}}} \\
0 & 0 & 0 & 1
\end{bmatrix}
$$

(7)

This matrix is multiplied on the right by the composite viewing transformation $RT$ (Section 4) to produce the complete transformation from world coordinates to normalized orthogonal-projection coordinates.

At this stage of the viewing pipeline, all device-independent coordinate transformations are completed and can be concatenated into a single composite matrix. Thus, the clipping procedures are most efficiently performed following the normalization transformation. After clipping, procedures for visibility testing, surface rendering, and the viewport transformation can be applied to generate the final screen display of the scene.

7 Oblique Parallel Projections

In general, a parallel-projection view of a scene is obtained by transferring object descriptions to the view plane along projection paths that can be in any selected
direction relative to the view-plane normal vector. When the projection path is not perpendicular to the view plane, this mapping is called an oblique parallel projection. Using this projection, we can produce combinations such as a front, side, and top view of an object, as in Figure 25. Oblique parallel projections are defined by a vector direction for the projection lines, and this direction can be specified in various ways.

**Oblique Parallel Projections in Drafting and Design**

For applications in engineering and architectural design, an oblique parallel projection is often specified with two angles, $\alpha$ and $\phi$, as shown in Figure 26. A spatial position $(x, y, z)$, in this illustration, is projected to $(x_p, y_p, z_{vp})$ on a view plane, which is at location $z_{vp}$ along the viewing $z$ axis. Position $(x, y, z_{vp})$ is the corresponding orthogonal-projection point. The oblique parallel projection line from $(x, y, z)$ to $(x_p, y_p, z_{vp})$ has an intersection angle $\alpha$ with the line on the projection plane that joins $(x_p, y_p, z_{vp})$ and $(x, y, z_{vp})$. This view-plane line, with length $L$, is at an angle $\phi$ with the horizontal direction in the projection plane. Angle $\alpha$ can be assigned a value between 0 and $90^\circ$, and angle $\phi$ can vary from 0 to $360^\circ$. We can express the projection coordinates in terms of $x$, $y$, $L$, and $\phi$ as

$$
\begin{align*}
    x_p &= x + L \cos \phi \\
    y_p &= y + L \sin \phi
\end{align*}
$$

(Figure 26)

An oblique parallel projection of coordinate position $(x, y, z)$ to position $(x_p, y_p, z_{vp})$ on a projection plane at position $z_{vp}$ along the $z_{view}$ axis.
Length $L$ depends on the angle $\alpha$ and the perpendicular distance of the point $(x, y, z)$ from the view plane:

$$\tan \alpha = \frac{z_{vp} - z}{L}$$

Thus

$$L = \frac{z_{vp} - z}{\tan \alpha} = L_1(z_{vp} - z)$$

where $L_1 = \cot \alpha$, which is also the value of $L$ when $z_{vp} - z = 1$. We can then write the oblique parallel projection equations (8) as

$$x_p = x + L_1(z_{vp} - z) \cos \phi$$

$$y_p = y + L_1(z_{vp} - z) \sin \phi$$

Equations 11 represent a $z$-axis shearing transformation. In fact, the effect of an oblique parallel projection is to shear planes of constant $z$ and project them onto the view plane. The $(x, y)$ positions on each plane of constant $z$ are shifted by an amount proportional to the distance of the plane from the view plane, so that angles, distances, and parallel lines in the plane are projected accurately. This effect is shown in Figure 27, where the view plane is positioned at the front face of a cube. The back plane of the cube is sheared and overlapped with the front plane in the projection to the viewing surface. A side edge of the cube connecting the front and back planes is projected into a line of length $L_1$ that makes an angle $\phi$ with a horizontal line in the projection plane.

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**Figure 27**

An oblique parallel projection (a) of a cube (top view) onto a view plane that is coincident with the front face of the cube produces the combination front, side, and top view shown in (b).

**Cavalier and Cabinet Oblique Parallel Projections**

Typical choices for angle $\phi$ are $30^\circ$ and $45^\circ$, which display a combination view of the front, side, and top (or front, side, and bottom) of an object. Two commonly used values for $\alpha$ are those for which $\tan \alpha = 1$ and $\tan \alpha = 2$. For the first case, $\alpha = 45^\circ$ and the views obtained are called **cavalier** projections. All lines perpendicular to the projection plane are projected with no change in length. Examples of cavalier projections for a cube are given in Figure 28.

When the projection angle $\alpha$ is chosen so that $\tan \alpha = 2$, the resulting view is called a **cabinet** projection. For this angle ($\approx 63.4^\circ$), lines perpendicular to the viewing surface are projected at half their length. Cabinet projections appear more realistic than cavalier projections because of this reduction in the length of perpendiculars. Figure 29 shows examples of cabinet projections for a cube.
Cavalier projections of a cube onto a
view plane for two values of angle $\phi$.
The depth of the cube is projected with
a length equal to that of the width and
height.

(a) $\phi = 45^\circ$
(b) $\phi = 30^\circ$

Cabinet projections of a cube onto a
view plane for two values of angle $\phi$.
The depth is projected with a length
that is one half that of the width and
height of the cube.

(a) $\phi = 45^\circ$
(b) $\phi = 30^\circ$

Oblique Parallel-Projection Vector

In graphics programming libraries that support oblique parallel projections, the
direction of projection to the view plane is specified with a parallel-projection vector, $V_p$. This direction vector can be designated with a reference position relative to
the view point, as we did with the view-plane normal vector, or with any other two
points. Some packages use a reference point relative to the center of the clipping
window to define the direction for a parallel projection. If the projection vector is
specified in world coordinates, it must first be transformed to viewing coordinates
using the rotation matrix discussed in Section 4. (The projection vector is unaffected by the translation, because it is simply a direction with no fixed position.)

Once the projection vector $V_p$ is established in viewing coordinates, all points
in the scene are transferred to the view plane along lines that are parallel to this
vector. Figure 30 illustrates an oblique parallel projection of a spatial point to
the view plane. We can denote the components of the projection vector relative
to the viewing-coordinate frame as $V_p = (V_{px}, V_{py}, V_{pz})$, where $V_{py}/V_{px} = \tan \phi$.

Then, comparing similar triangles in Figure 30, we have

\[
\frac{x_p - x}{z_{vp} - z} = \frac{V_{px}}{V_{pz}}
\]

\[
\frac{y_p - y}{z_{vp} - z} = \frac{V_{py}}{V_{pz}}
\]

And we can write the equivalent of the oblique parallel-projection equations 11
in terms of the projection vector as

\[
x_p = x + (z_{vp} - z) \frac{V_{px}}{V_{pz}}
\]

\[
y_p = y + (z_{vp} - z) \frac{V_{py}}{V_{pz}}
\]

(12)

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The oblique parallel-projection coordinates in 12 reduce to the orthogonal-projection coordinates 6 when \( V_{px} = V_{py} = 0 \).

**Clipping Window and Oblique Parallel-Projection View Volume**

A view volume for an oblique parallel projection is set up using the same procedures as in an orthogonal projection. We select a clipping window on the view plane with coordinate positions \((x_{w_{\text{min}}}, y_{w_{\text{min}}})\) and \((x_{w_{\text{max}}}, y_{w_{\text{max}}})\), for the lower-left and upper-right corners of the clipping rectangle. The top, bottom, and sides of the view volume are then defined by the direction of projection and the edges of the clipping window. In addition, we can limit the extent of the view volume by adding a near plane and a far plane, as in Figure 31. The finite oblique parallel-projection view volume is an oblique parallelepiped.

Oblique parallel projections may be affected by changes in the position of the view plane, depending on how the projection direction is to be specified. In some systems, the oblique parallel-projection direction is parallel to the line connecting a reference point to the center of the clipping window. Therefore, moving the position of the view plane or clipping window without adjusting the reference point changes the shape of the view volume.

**Oblique Parallel-Projection Transformation Matrix**

Using the projection-vector parameters from the equations in 12, we can express the elements of the transformation matrix for an oblique parallel
projection as

\[
M_{\text{oblique}} = \begin{bmatrix}
1 & 0 & \frac{V_{px}}{V_{pz}} & z_{vp} \frac{V_{pz}}{V_{pz}} \\
0 & 1 & \frac{V_{py}}{V_{pz}} & z_{vp} \frac{V_{pz}}{V_{pz}} \\
0 & 0 & 1 & 0 \\
0 & 0 & 0 & 1
\end{bmatrix}
\] (13)

This matrix shifts the values of the \(x\) and \(y\) coordinates by an amount proportional to the distance from the view plane, which is at position \(z_{vp}\) on the \(z\) view axis. The \(z\) values of spatial positions are unchanged. If \(V_{px} = V_{py} = 0\), we have an orthogonal projection and matrix 13 is reduced to the identity matrix.

For a general oblique parallel projection, matrix 13 represents a \(z\)-axis shearing transformation. All coordinate positions within the oblique view volume are sheared by an amount proportional to their distance from the view plane. The effect is to shear the oblique view volume into a rectangular parallelepiped, as illustrated in Figure 32. Thus, positions inside the view volume are sheared into orthogonal-projection coordinates by the oblique parallel-projection transformation.

Normalization Transformation for an Oblique Parallel Projection

Because the oblique parallel-projection equations convert object descriptions to orthogonal-coordinate positions, we can apply the normalization procedures following this transformation. The oblique view volume has been converted to a rectangular parallelepiped, so we use the same procedures as in Section 6.

Following the normalization example in Section 6, we again map to the symmetric normalized cube within a left-handed coordinate frame. Thus, the complete transformation, from viewing coordinates to normalized coordinates, for an oblique parallel projection is

\[
M_{\text{oblique, norm}} = M_{\text{ortho, norm}} \cdot M_{\text{oblique}}
\] (14)

\[\text{Clipping Window} \quad \text{View Plane} \quad \text{Clipping Window} \]

\[\text{Near Plane} \quad \text{Shear Transformation} \quad \text{Far Plane} \]

(a) Oblique-Projection View Volume

(b) Transformed Oblique View Volume

\textbf{Figure 32}  
Top view of an oblique parallel-projection transformation. The oblique view volume is converted into a rectangular parallelepiped, and objects in the view volume, such as the green block, are mapped to orthogonal-projection coordinates.
Transformation $M_{oblique}$ is matrix 13, which converts the scene description to orthogonal-projection coordinates; and transformation $M_{ortho,norm}$ is matrix 7, which maps the contents of the orthogonal view volume to the symmetric normalization cube.

To complete the viewing transformations (with the exception of the mapping to viewport screen coordinates), we concatenate matrix 14 to the left of the transformation $M_{WC, VC}$ from Section 4. Clipping routines can then be applied to the normalized view volume, followed by the determination of visible objects, the surface-rendering procedures, and the viewport transformation.

8 Perspective Projections

Although a parallel-projection view of a scene is easy to generate and preserves relative proportions of objects, it does not provide a realistic representation. To simulate a camera picture, we need to consider that reflected light rays from the objects in a scene follow converging paths to the camera film plane. We can approximate this geometric-optics effect by projecting objects to the view plane along converging paths to a position called the projection reference point (or center of projection). Objects are then displayed with foreshortening effects, and projections of distant objects are smaller than the projections of objects of the same size that are closer to the view plane (Figure 33).

Perspective-Projection Transformation Coordinates

We can sometimes select the projection reference point as another viewing parameter in a graphics package, but some systems place this convergence point at a fixed position, such as at the view point. Figure 34 shows the projection path of a spatial position $(x, y, z)$ to a general projection reference point at $(x_{prp}, y_{prp}, z_{prp})$. The projection line intersects the view plane at the coordinate position $(x_p, y_p, z_{vp})$, where $z_{vp}$ is some selected position for the view plane on the $z_{view}$ axis. We can write equations describing coordinate positions along this perspective-projection line in parametric form as

$$
x' = x - (x - x_{prp})u \quad 0 \leq u \leq 1
\]
$$

$$
y' = y - (y - y_{prp})u
\]

$$
z' = z - (z - z_{prp})u
\]

(15)

Coordinate position $(x', y', z')$ represents any point along the projection line. When $u = 0$, we are at position $P = (x, y, z)$. At the other end of the line, $u = 1$ and...
we have the projection reference-point coordinates \((x_{prp}, y_{prp}, z_{prp})\). On the view plane, \(z' = z_v\) and we can solve the \(z'\) equation for parameter \(u\) at this position along the projection line:

\[
u = \frac{z_v - z}{z_{prp} - z}
\]

Substituting this value of \(u\) into the equations for \(x'\) and \(y'\), we obtain the general perspective-transformation equations

\[
\begin{align*}
x_p &= x \left( \frac{z_{prp} - z_v}{z_{prp} - z} \right) + x_{prp} \left( \frac{z_v - z}{z_{prp} - z} \right) \\
y_p &= y \left( \frac{z_{prp} - z_v}{z_{prp} - z} \right) + y_{prp} \left( \frac{z_v - z}{z_{prp} - z} \right)
\end{align*}
\]

Calculations for a perspective mapping are more complex than the parallel-projection equations, because the denominators in the perspective calculations 17 are functions of the \(z\) coordinate of the spatial position. Therefore, we now need to formulate the perspective-transformation procedures a little differently so that this mapping can be concatenated with the other viewing transformations. But first we take a look at some of the properties of Equations 17.

**Perspective-Projection Equations: Special Cases**

Various restrictions are often placed on the parameters for a perspective projection. Depending on a particular graphics package, positioning for either the projection reference point or the view plane may not be completely optional.

To simplify the perspective calculations, the projection reference point could be limited to positions along the \(z_{\text{view}}\) axis, then

1. \(x_{prp} = y_{prp} = 0\):

\[
x_p = x \left( \frac{z_{prp} - z_v}{z_{prp} - z} \right), \quad y_p = y \left( \frac{z_{prp} - z_v}{z_{prp} - z} \right)
\]

Sometimes the projection reference point is fixed at the coordinate origin, and

2. \((x_{prp}, y_{prp}, z_{prp}) = (0, 0, 0)\):

\[
x_p = x \left( \frac{z_v}{z} \right), \quad y_p = y \left( \frac{z_v}{z} \right)
\]

If the view plane is the \(uv\) plane and there are no restrictions on the placement of the projection reference point, then we have
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3. $z_{vp} = 0$:

$$
x_p = x \left( \frac{z_{prp}}{z_{prp} - z} \right) - x_{prp} \left( \frac{z}{z_{prp} - z} \right)
$$

$$
y_p = y \left( \frac{z_{prp}}{z_{prp} - z} \right) - y_{prp} \left( \frac{z}{z_{prp} - z} \right)
$$

(20)

With the $uv$ plane as the view plane and the projection reference point on the $z_{view}$ axis, the perspective equations are

4. $x_{prp} = y_{prp} = z_{vp} = 0$:

$$
x_p = x \left( \frac{z_{prp}}{z_{prp} - z} \right), \quad y_p = y \left( \frac{z_{prp}}{z_{prp} - z} \right)
$$

(21)

Of course, we cannot have the projection reference point on the view plane. In that case, the entire scene would project to a single point. The view plane is usually placed between the projection reference point and the scene, but, in general, the view plane could be placed anywhere except at the projection point. If the projection reference point is between the view plane and the scene, objects are inverted on the view plane (Figure 35). With the scene between the view plane and the projection point, objects are simply enlarged as they are projected away from the viewing position onto the view plane.

Perspective effects also depend on the distance between the projection reference point and the view plane, as illustrated in Figure 36. If the projection

![Figure 35](image1)

**Figure 35**
A perspective-projection view of an object is upside down when the projection reference point is between the object and the view plane.

![Figure 36](image2)

**Figure 36**
Changing perspective effects by moving the projection reference point away from the view plane.
reference point is close to the view plane, perspective effects are emphasized; that is, closer objects will appear much larger than more distant objects of the same size. Similarly, as the projection reference point moves farther from the view plane, the difference in the size of near and far objects decreases. When the projection reference point is very far from the view plane, a perspective projection approaches a parallel projection.

**Vanishing Points for Perspective Projections**

When a scene is projected onto a view plane using a perspective mapping, lines that are parallel to the view plane are projected as parallel lines. But any parallel lines in the scene that are not parallel to the view plane are projected into converging lines. The point at which a set of projected parallel lines appears to converge is called a **vanishing point**. Each set of projected parallel lines has a separate vanishing point.

For a set of lines that are parallel to one of the principal axes of an object, the vanishing point is referred to as a **principal vanishing point**. We control the number of principal vanishing points (one, two, or three) with the orientation of the projection plane, and perspective projections are accordingly classified as one-point, two-point, or three-point projections. The number of principal vanishing points in a projection is equal to the number of principal axes that intersect the view plane. Figure 37 illustrates the appearance of one-point and two-point perspective projections for a cube. In the projected view (b), the view plane is aligned parallel to the $xy$ object plane so that only the object $z$ axis is intersected. This orientation produces a one-point perspective projection with a $z$-axis

![Diagram](image-url)
vanishing point. For the view shown in (c), the projection plane intersects both the \(x\) and \(z\) axes but not the \(y\) axis. The resulting two-point perspective projection contains both \(x\)-axis and \(z\)-axis vanishing points. There is not much increase in the realism of a three-point perspective projection compared to a two-point projection, so three-point projections are not used as often in architectural and engineering drawings.

**Perspective-Projection View Volume**

We again create a view volume by specifying the position of a rectangular clipping window on the view plane. But now the bounding planes for the view volume are not parallel, because the projection lines are not parallel. The bottom, top, and sides of the view volume are planes through the window edges that all intersect at the projection reference point. This forms a view volume that is an infinite rectangular pyramid with its apex at the center of projection (Figure 38). All objects outside this pyramid are eliminated by the clipping routines. A perspective-projection view volume is often referred to as a **pyramid of vision** because it approximates the **cone of vision** of our eyes or a camera. The displayed view of a scene includes only those objects within the pyramid, just as we cannot see objects beyond our peripheral vision, which are outside the cone of vision.

By adding near and far clipping planes that are perpendicular to the \(z_{\text{view}}\) axis (and parallel to the view plane), we chop off parts of the infinite, perspective-projection view volume to form a truncated pyramid, or **frustum**, view volume. Figure 39 illustrates the shape of a finite, perspective-projection view volume with a view plane that is placed between the near clipping plane and the projection reference point. Sometimes the near and far planes are required in a graphics package, and sometimes they are optional.

Usually, both the near and far clipping planes are on the same side of the projection reference point, with the far plane farther from the projection point than the near plane along the viewing direction. And, as in a parallel projection, we can use the near and far planes simply to enclose the scene to be viewed. But with a perspective projection, we could also use the near clipping plane to take out large objects close to the view plane that could project into unrecognizable shapes within the clipping window. Similarly, the far clipping plane could be used to cut out objects far from the projection reference point that might project to small blots on the view plane. Some systems restrict the placement of the view plane relative to the near and far planes, and other systems allow it to be placed anywhere except...
at the position of the projection reference point. If the view plane is “behind” the projection reference point, objects are inverted, as shown in Figure 35.

**Perspective-Projection Transformation Matrix**

Unlike a parallel projection, we cannot directly use the coefficients of the $x$ and $y$ coordinates in equations 17 to form the perspective-projection matrix elements, because the denominators of the coefficients are functions of the $z$ coordinate. But we can use a three-dimensional, homogeneous-coordinate representation to express the perspective-projection equations in the form

$$x_p = \frac{x_h}{h}, \quad y_p = \frac{y_h}{h}$$

(22)

where the homogeneous parameter has the value

$$h = z_{pp} - z$$

(23)

The numerators in 22 are the same as in equations 17:

$$x_h = x(z_{pp} - z_{vp}) + x_{prp}(z_{vp} - z)$$

$$y_h = y(z_{pp} - z_{vp}) + y_{prp}(z_{vp} - z)$$

(24)

Thus, we can set up a transformation matrix to convert a spatial position to homogeneous coordinates so that the matrix contains only the perspective parameters and not coordinate values. The perspective-projection transformation of a viewing-coordinate position is then accomplished in two steps. First, we calculate the homogeneous coordinates using the perspective-transformation matrix:

$$P_h = M_{pers} \cdot P$$

(25)

where $P_h$ is the column-matrix representation of the homogeneous point $(x_h, y_h, z_h, h)$ and $P$ is the column-matrix representation of the coordinate position $(x, y, z, 1)$. (Actually, the perspective matrix would be concatenated with the other viewing-transformation matrices, and then the composite matrix would be applied to the world-coordinate description of a scene to produce homogeneous coordinates.) Second, after other processes have been applied, such as the normalization transformation and clipping routines, homogeneous coordinates are divided by parameter $h$ to obtain the true transformation-coordinate positions.
Setting up matrix elements for obtaining the homogeneous-coordinate \( x_h \) and \( y_h \) values in 24 is straightforward, but we must also structure the matrix to preserve depth (z-value) information. Otherwise, the z coordinates are distorted by the homogeneous-division parameter \( h \). We can do this by setting up the matrix elements for the z transformation so as to normalize the perspective-projection \( z_p \) coordinates. There are various ways that we could choose the matrix elements to produce the homogeneous coordinates 24 and the normalized \( z_p \) value for a spatial position \((x, y, z)\). The following matrix gives one possible way to formulate a perspective-projection matrix.

\[
M_{\text{pers}} = \begin{bmatrix}
z_{prp} - z_{vp} & 0 & -x_{prp} & x_{prp}z_{prp} \\
0 & z_{prp} - z_{vp} & -y_{prp} & y_{prp}z_{prp} \\
0 & 0 & s_z & t_z \\
0 & 0 & -1 & z_{prp}
\end{bmatrix}
\]  

(26)

Parameters \( s_z \) and \( t_z \) are the scaling and translation factors for normalizing the projected values of \( z \)-coordinates. Specific values for \( s_z \) and \( t_z \) depend on the normalization range we select.

Matrix 26 converts the description of a scene into homogeneous parallel-projection coordinates. However, the frustum view volume can have any orientation, so that these transformed coordinates could correspond to an oblique parallel projection. This occurs if the frustum view volume is not symmetric. If the frustum view volume for the perspective projection is symmetric, the resulting parallel-projection coordinates correspond to an orthogonal projection. We next consider these two possibilities.

**Symmetric Perspective-Projection Frustum**

The line from the projection reference point through the center of the clipping window and on through the view volume is the centerline for a perspective-projection frustum. If this centerline is perpendicular to the view plane, we have a symmetric frustum (with respect to its centerline) as in Figure 40.

Because the frustum centerline intersects the view plane at the coordinate location \((x_{prp}, y_{prp}, z_{vp})\), we can express the corner positions for the clipping window.
Therefore, we could specify a symmetric perspective-projection view of a scene using the width and height of the clipping window instead of the window coordinates. This uniquely establishes the position of the clipping window, because it is symmetric about the x and y coordinates of the projection reference point.

Another way to specify a symmetric perspective projection is to use parameters that approximate the properties of a camera lens. A photograph is produced with a symmetric perspective projection of a scene onto the film plane. Reflected light rays from the objects in a scene are collected on the film plane from within the “cone of vision” of the camera. This cone of vision can be referenced with a field-of-view angle, which is a measure of the size of the camera lens. A large field-of-view angle, for example, corresponds to a wide-angle lens. In computer graphics, the cone of vision is approximated with a symmetric frustum, and we can use a field-of-view angle to specify an angular size for the frustum. Typically, the field-of-view angle is the angle between the top clipping plane and the bottom clipping plane of the frustum, as shown in Figure 41.

For a given projection reference point and view-plane position, the field-of-view angle determines the height of the clipping window (Figure 42), but not the width. We need an additional parameter to define completely the clipping-window dimensions, and this second parameter could be either the window width or the aspect ratio (width/height) of the clipping window. From the right triangles in the diagram of Figure 42, we see that

$$\tan\left(\frac{\theta}{2}\right) = \frac{\text{height}/2}{z_{prp} - z_{vp}}$$

so that the clipping-window height can be calculated as

$$\text{height} = 2(z_{prp} - z_{vp}) \tan\left(\frac{\theta}{2}\right)$$

\[\text{(27)}\]
Therefore, the diagonal elements with the value $z_{prp} - z_{vp}$ in matrix 26 could be replaced by either of the following two expressions.

$$z_{prp} - z_{vp} = \frac{\text{height}}{2} \cot \left( \frac{\theta}{2} \right)$$

$$= \frac{\text{width} \cdot \cot(\theta/2)}{2 \cdot \text{aspect}}$$

(29)

In some graphics libraries, fixed positions are used for the view plane and the projection reference point, so that a symmetric perspective projection is completely specified by the field-of-view angle, the aspect ratio of the clipping window, and the distances from the viewing position to the near and far clipping planes. The same aspect ratio is usually applied to the specification of the viewport.

If the field-of-view angle is decreased in a particular application, the foreshortening effects of a perspective projection are also decreased. This is comparable to moving the projection reference point farther from the view plane. Also, decreasing the field-of-view angle decreases the height of the clipping window, and this provides a method for zooming in on small regions of a scene. Similarly, a large field-of-view angle results in a large clipping-window height (a zoom out), and it increases perspective effects, which is what we achieve when we set the projection reference point close to the view plane. Figure 43 illustrates the effects of various field-of-view angles for a fixed-width clipping window.

When the perspective-projection view volume is a symmetric frustum, the perspective transformation maps locations inside the frustum to orthogonal-projection coordinates within a rectangular parallelepiped. The centerline of the parallelepiped is the frustum centerline, because this line is already perpendicular to the view plane (Figure 44). This is a consequence of the fact that all positions along a projection line within the frustum map to the same point $(x_p, y_p)$ on the view plane. Thus, each projection line is converted by the perspective transformation to a line that is perpendicular to the view plane and, thus, parallel to the frustum centerline. With the symmetric frustum converted to an orthogonal-projection view volume, we can next apply the normalization transformation.

**Oblique Perspective-Projection Frustum**

If the centerline of a perspective-projection view volume is not perpendicular to the view plane, we have an oblique frustum. Figure 45 illustrates the general
Increasing the size of the field-of-view angle increases the height of the clipping window and increases the perspective-projection foreshortening.

A symmetric frustum view volume is mapped to an orthogonal parallelepiped by a perspective-projection transformation.
appearance of an oblique perspective-projection view volume. In this case, we can first transform the view volume to a symmetric frustum and then to a normalized view volume.

An oblique perspective-projection view volume can be converted to a symmetric frustum by applying a z-axis shearing-transformation matrix. This transformation shifts all positions on any plane that is perpendicular to the z axis by an amount that is proportional to the distance of the plane from a specified z-axis reference position. In this case, the reference position is \( z_{prp} \), which is the z coordinate of the projection reference point. And we need to shift by an amount that will move the center of the clipping window to position \((x_{prp}, y_{prp})\) on the view plane. Because the frustum centerline passes through the center of the clipping window, this shift adjusts the centerline so that it is perpendicular to the view plane, as in Figure 40.

The computations for the shearing transformation, as well as for the perspective and normalization transformations, are greatly reduced if we take the projection reference point to be the viewing-coordinate origin. We could do this with no loss in generality by translating all coordinate positions in a scene so that our selected projection reference point is shifted to the coordinate origin. Or we could have initially set up the viewing-coordinate reference frame so that its origin is at the projection point that we want for a scene. And, in fact, some graphics libraries do fix the projection reference point at the coordinate origin.

Taking the projection reference point as \((x_{prp}, y_{prp}, z_{prp}) = (0, 0, 0)\), we obtain the elements of the required shearing matrix as

\[
M_{z\text{shear}} = \begin{bmatrix}
1 & 0 & \text{sh}_{zx} & 0 \\
0 & 1 & \text{sh}_{zy} & 0 \\
0 & 0 & 1 & 0 \\
0 & 0 & 0 & 1
\end{bmatrix}
\] (30)

We can also simplify the elements of the perspective-projection matrix a bit more if we place the view plane at the position of the near clipping plane. And, because we now want to move the center of the clipping window to coordinates \((0, 0)\)
on the view plane, we need to choose values for the shearing parameters such that

$$\begin{bmatrix}
0 \\
0 \\
z_{\text{near}} \\
1
\end{bmatrix} = M_{z,\text{shear}} \cdot \begin{bmatrix}
x w_{\text{min}} + x w_{\text{max}} \\
y w_{\text{min}} + y w_{\text{max}} \\
z_{\text{near}} \\
1
\end{bmatrix}$$

(31)

Therefore, the parameters for this shearing transformation are

$$sh_{xx} = -\frac{x w_{\text{min}} + x w_{\text{max}}}{2 z_{\text{near}}}$$

$$sh_{yy} = -\frac{y w_{\text{min}} + y w_{\text{max}}}{2 z_{\text{near}}}$$

(32)

Similarly, with the projection reference point at the viewing-coordinate origin and with the near clipping plane as the view plane, the perspective-projection matrix 26 is simplified to

$$M_{\text{pers}} = \begin{bmatrix}
-z_{\text{near}} & 0 & 0 & 0 \\
0 & -z_{\text{near}} & 0 & 0 \\
0 & 0 & s_z & t_z \\
0 & 0 & -1 & 0
\end{bmatrix}$$

(33)

Expressions for the z-coordinate scaling and translation parameters will be determined by the normalization requirements.

Concatenating the simplified perspective-projection matrix 33 with the shear matrix 30, we obtain the following oblique perspective-projection matrix for converting coordinate positions in a scene to homogeneous orthogonal-projection coordinates. The projection reference point for this transformation is the viewing-coordinate origin, and the near clipping plane is the view plane.

$$M_{\text{obliquepers}} = M_{\text{pers}} \cdot M_{z,\text{shear}}$$

$$= \begin{bmatrix}
-z_{\text{near}} & 0 & x w_{\text{min}} + x w_{\text{max}} & 0 \\
0 & -z_{\text{near}} & \frac{y w_{\text{min}} + y w_{\text{max}}}{2} & 0 \\
0 & 0 & s_z & t_z \\
0 & 0 & -1 & 0
\end{bmatrix}$$

(34)

Although we no longer have options for the placement of the projection reference point and the view plane, this matrix provides an efficient method for generating a perspective-projection view of a scene without sacrificing a great deal of flexibility.

If we choose the clipping-window coordinates so that $x w_{\text{max}} = -x w_{\text{min}}$ and $y w_{\text{max}} = -y w_{\text{min}}$, the frustum view volume is symmetric and matrix 34 reduces to matrix 33. This is because the projection reference point is now at the origin of the viewing-coordinate frame. We could also use Equations 29, with $z_{\text{prp}} = 0$ and $z_{\text{vp}} = z_{\text{near}}$, to express the first two diagonal elements of this matrix in terms of the field-of-view angle and the clipping-window dimensions.

### Normalized Perspective-Projection Transformation Coordinates

Matrix 34 transforms object positions in viewing coordinates to perspective-projection homogeneous coordinates. When we divide the homogeneous coordinates by the homogeneous parameter $h$, we obtain the actual projection coordinates, which are orthogonal-projection coordinates. Thus, this perspective
Projection transforms all points within the frustum view volume to positions within a rectangular parallelepiped view volume. The final step in the perspective transformation process is to map this parallelepiped to a *normalized view volume*.

We follow the same normalization procedure that we used for a parallel projection. The transformed frustum view volume, which is a rectangular parallelepiped, is mapped to a symmetric normalized cube within a left-handed reference frame (Figure 46). We have already included the normalization parameters for $z$ coordinates in the perspective-projection matrix 34, but we still need to determine the values for these parameters when we transform to the symmetric normalization cube. Also, we need to determine the normalization transformation parameters for $x$ and $y$ coordinates. Because the centerline of the rectangular parallelepiped view volume is now the $z_{\text{view}}$ axis, no translation is needed in the $x$ and $y$ normalization transformations: We require only the $x$ and $y$ scaling parameters relative to the coordinate origin. The scaling matrix for accomplishing the $xy$ normalization is

$$
M_{xy\text{scale}} = \begin{bmatrix}
    s_x & 0 & 0 & 0 \\
    0 & s_y & 0 & 0 \\
    0 & 0 & 1 & 0 \\
    0 & 0 & 0 & 1
\end{bmatrix} \tag{35}
$$

Concatenating the $xy$-scaling matrix with matrix 34 produces the following normalization matrix for a perspective-projection transformation.

$$
M_{\text{normpers}} = M_{xy\text{scale}} \cdot M_{\text{obliquepers}} = \begin{bmatrix}
    -z_{\text{near}}s_x & 0 & s_x \frac{x w_{\text{min}} + x w_{\text{max}}}{2} & 0 \\
    0 & -z_{\text{near}}s_y & s_y \frac{y w_{\text{min}} + y w_{\text{max}}}{2} & 0 \\
    0 & 0 & s_z & t_z \\
    0 & 0 & -1 & 0
\end{bmatrix} \tag{36}
$$

From this transformation, we obtain the homogeneous coordinates:

$$
\begin{bmatrix}
    x_h \\
    y_h \\
    z_h \\
    h
\end{bmatrix} = M_{\text{normpers}} \cdot \begin{bmatrix}
    x \\
    y \\
    z \\
    1
\end{bmatrix} \tag{37}
$$
And the projection coordinates are

\[
\begin{align*}
    x_p &= \frac{x_h}{h} = \frac{-z_{\text{near}} s_x x + s_z (x_{w_{\text{min}}} + x_{w_{\text{max}}})}{-z} \\
    y_p &= \frac{y_h}{h} = \frac{-z_{\text{near}} s_y y + s_y (y_{w_{\text{min}}} + y_{w_{\text{max}}})}{-z} \\
    z_p &= \frac{z_h}{h} = \frac{s_z z + t_z}{-z}
\end{align*}
\]  

(38)

To normalize this perspective transformation, we want the projection coordinates to be \((x_p, y_p, z_p) = (-1, -1, -1)\) when the input coordinates are \((x, y, z) = (x_{w_{\text{min}}}, y_{w_{\text{min}}}, z_{\text{near}})\), and we want the projection coordinates to be \((x_p, y_p, z_p) = (1, 1, 1)\) when the input coordinates are \((x, y, z) = (x_{w_{\text{max}}}, y_{w_{\text{max}}}, z_{\text{far}})\). Therefore, when we solve equations 38 for the normalization parameters using these conditions, we obtain

\[
\begin{align*}
    s_x &= \frac{2}{x_{w_{\text{max}}} - x_{w_{\text{min}}}} \\
    s_y &= \frac{2}{y_{w_{\text{max}}} - y_{w_{\text{min}}}} \\
    s_z &= \frac{z_{\text{near}} + z_{\text{far}}}{z_{\text{near}} - z_{\text{far}}} \\
    t_z &= \frac{2 z_{\text{near}} z_{\text{far}}}{z_{\text{near}} - z_{\text{far}}}
\end{align*}
\]  

(39)

And the elements of the normalized transformation matrix for a general perspective-projection are

\[
M_{\text{normpers}} = \begin{bmatrix}
-2z_{\text{near}} & 0 & x_{w_{\text{max}}} + x_{w_{\text{min}}} & 0 \\
0 & -2z_{\text{near}} & y_{w_{\text{max}}} + y_{w_{\text{min}}} & 0 \\
0 & 0 & z_{\text{near}} + z_{\text{far}} & 0 \\
0 & 0 & 2z_{\text{near}} z_{\text{far}} & -1
\end{bmatrix}
\]  

(40)

If the perspective-projection view volume was originally specified as a symmetric frustum, we can express the elements of the normalized perspective transformation in terms of the field-of-view angle and the dimensions of the clipping window. Thus, using Equations 29, with the projection reference point at the origin and the view plane at the position of the near clipping plane, we have

\[
M_{\text{normsympers}} = \begin{bmatrix}
\cot \left( \frac{\theta}{2} \right) & 0 & 0 & 0 \\
0 & \cot \left( \frac{\theta}{2} \right) & 0 & 0 \\
0 & 0 & z_{\text{near}} + z_{\text{far}} & 0 \\
0 & 0 & 2z_{\text{near}} z_{\text{far}} & -1
\end{bmatrix}
\]  

(41)

The complete transformation from world coordinates to normalized perspective-projection coordinates is the composite matrix formed by concatenating this perspective matrix on the left of the viewing-transformation product \(R \cdot T\). Next, the clipping routines can be applied to the normalized view volume. The remaining tasks are visibility determination, surface rendering, and the transformation to the viewport.
9 The Viewport Transformation and Three-Dimensional Screen Coordinates

Once we have completed the transformation to normalized projection coordinates, clipping can be applied efficiently to the symmetric cube (or the unit cube). Following the clipping procedures, the contents of the normalized view volume can be transferred to screen coordinates. For the \( x \) and \( y \) positions in the normalized clipping window, this procedure is the same as the two-dimensional viewport transformation. But positions throughout the three-dimensional view volume also have a depth (\( z \) coordinate), and we need to retain this depth information for the visibility testing and surface-rendering algorithms. So we can now think of the viewport transformation as a mapping to \textbf{three-dimensional screen coordinates}.

The \( x \) and \( y \) transformation equations from the normalized clipping window to positions within a rectangular viewport are given in matrix 8-10. We can adapt that matrix to three-dimensional applications by including parameters for the transformation of \( z \) values to screen coordinates. Often the normalized \( z \) values within the symmetric cube are renormalized on the range from 0 to 1.0. This allows the video screen to be referenced as \( z = 0 \), and depth processing can be conveniently carried out over the unit interval from 0 to 1. If we include this \( z \) renormalization, the transformation from the normalized view volume to three-dimensional screen coordinates is

\[
M_{\text{normviewvol,3D screen}} = \begin{bmatrix}
\frac{xv_{\text{max}} - xv_{\text{min}}}{2} & 0 & 0 & \frac{xv_{\text{max}} + xv_{\text{min}}}{2} \\
0 & \frac{yv_{\text{max}} - yv_{\text{min}}}{2} & 0 & \frac{yv_{\text{max}} + yv_{\text{min}}}{2} \\
0 & 0 & 1 & 1 \\
0 & 0 & \frac{2}{2} & \frac{2}{2}
\end{bmatrix}
\]

In normalized coordinates, the \( z_{\text{norm}} = -1 \) face of the symmetric cube corresponds to the clipping-window area. And this face of the normalized cube is mapped to the rectangular viewport, which is now referenced at \( z_{\text{screen}} = 0 \). Thus, the lower-left corner of the viewport screen area is at position \( (xv_{\text{min}}, yv_{\text{min}}, 0) \) and the upper-right corner is at position \( (xv_{\text{max}}, yv_{\text{max}}, 0) \).

Each \( xy \) position on the viewport corresponds to a position in the refresh buffer, which contains the color information for that point on the screen. And the depth value for each screen point is stored in another buffer area, called the \textit{depth buffer}. In later chapters, we explore the algorithms for determining the visible surface positions and their colors.

We position the rectangular viewport on the screen just as we did for two-dimensional applications. The lower-left corner of the viewport is usually placed at a coordinate position specified relative to the lower-left corner of the display window. And object proportions are maintained if we set the aspect ratio of this viewport area to be the same as the clipping window.

10 OpenGL Three-Dimensional Viewing Functions

The OpenGL Utility library (GLU) includes a function for specifying the three-dimensional viewing parameters and another function for setting up a symmetric
Hubjustification: perspective-projection transformation. Other functions, such as those for an orthogonal projection, an oblique perspective projection, and the viewport transformation, are contained in the basic OpenGL library. In addition, GLUT functions are available for defining and manipulating display windows.

OpenGL Viewing-Transformation Function

When we designate the viewing parameters in OpenGL, a matrix is formed and concatenated with the current modelview matrix. Consequently, this viewing matrix is combined with any geometric transformations we may have also specified. This composite matrix is then applied to transform object descriptions in world coordinates to viewing coordinates. We set the modelview mode with the statement:

```c
glMatrixMode (GL_MODELVIEW);
```

Viewing parameters are specified with the following GLU function, which is in the OpenGL Utility library because it invokes the translation and rotation routines in the basic OpenGL library.

```c
gluLookAt (x0, y0, z0, xref, yref, zref, Vx, Vy, Vz);
```

Values for all parameters in this function are to be assigned double-precision, floating-point values. This function designates the origin of the viewing reference frame as the world-coordinate position \( P_0 = (x_0, y_0, z_0) \), the reference position as \( P_{\text{ref}} = (x_{\text{ref}}, y_{\text{ref}}, z_{\text{ref}}) \), and the view-up vector as \( V = (V_x, V_y, V_z) \). The positive \( z_{\text{view}} \) axis for the viewing frame is in the direction \( N = P_0 - P_{\text{ref}} \), and the unit axis vectors for the viewing reference frame are calculated with Equations 1.

Because the viewing direction is along the \(-z_{\text{view}}\) axis, the reference position \( P_{\text{ref}} \) is also referred to as the “look-at point.” This is usually taken to be some position in the center of the scene that we can use as a reference for specifying the projection parameters. And we can think of the reference position as the point at which we want to aim a camera that is located at the viewing origin. The up orientation for the camera is designated with vector \( V \), which is adjusted to a direction perpendicular to \( N \).

Viewing parameters specified with the `gluLookAt` function are used to form the viewing-transformation matrix \( M \) that we derived in Section 4. This matrix is formed as a combination of a translation, which shifts the viewing origin to the world origin, and a rotation, which aligns the viewing axes with the world axes.

If we do not invoke the `gluLookAt` function, the default OpenGL viewing parameters are

\[
\begin{align*}
P_0 &= (0, 0, 0) \\
P_{\text{ref}} &= (0, 0, -1) \\
V &= (0, 1, 0)
\end{align*}
\]

For these default values, the viewing reference frame is the same as the world frame, with the viewing direction along the negative \( z_{\text{world}} \) axis. In many applications, we can conveniently use the default values for the viewing parameters.

OpenGL Orthogonal-Projection Function

Projection matrices are stored in the OpenGL projection mode. So, to set up a projection-transformation matrix, we must first invoke that mode with the
Then, when we issue any transformation command, the resulting matrix will be concatenated with the current projection matrix.

Orthogonal-projection parameters are chosen with the function

```
glOrtho (xwmin, xwmax, ywmin, ywmax, dnear, dfar);
```

All parameter values in this function are to be assigned double-precision, floating-point numbers. We use `glOrtho` to select the clipping-window coordinates and the distances to the near and far clipping planes from the viewing origin. There is no option in OpenGL for the placement of the view plane. The near clipping plane is always also the view plane, and therefore the clipping window is always on the near plane of the view volume.

Function `glOrtho` generates a parallel projection that is perpendicular to the view plane (the near clipping plane). Thus, this function creates a finite orthogonal-projection view volume for the specified clipping planes and clipping window. In OpenGL, the near and far clipping planes are not optional; they must always be specified for any projection transformation.

Parameters `dnear` and `dfar` denote distances in the negative `zview` direction from the viewing-coordinate origin. For example, if `dfar` = 55.0, then the far clipping plane is at the coordinate position `zfar` = −55.0. A negative value for either parameter denotes a distance “behind” the viewing origin, along the positive `zview` axis. We can assign any values (positive, negative, or zero) to these parameters, so long as `dnear` < `dfar`.

The resulting view volume for this projection transformation is a rectangular parallelepiped. Coordinate positions within this view volume are transformed to locations within the symmetric normalized cube in a left-handed reference frame using matrix 7, with `z_near` = −`dnear` and `z_far` = −`dfar`.

Default parameter values for the OpenGL orthogonal-projection function are ±1, which produce a view volume that is a symmetric normalized cube in the right-handed viewing-coordinate system. This default is equivalent to issuing the statement

```
glOrtho (-1.0, 1.0, -1.0, 1.0, -1.0, 1.0);
```

The default clipping window is thus a symmetric normalized square, and the default view volume is a symmetric normalized cube with `z_near` = 1.0 (behind the viewing position) and `z_far` = −1.0. Figure 47 shows the appearance and position of the default orthogonal-projection view volume.

For two-dimensional applications, we used the `gluOrtho2D` function to set up the clipping window. We could also have used the `glOrtho` function to specify the clipping window, as long as parameters `dnear` and `dfar` were assigned values that were on opposite sides of the coordinate origin. In fact, a call to `gluOrtho2D` is equivalent to a call to `glOrtho` with `dnear` = −1.0 and `dfar` = 1.0.

There is no OpenGL function for generating an oblique projection. To produce an oblique-projection view of a scene, we could set up our own projection matrix as in Equation 14. Then we need to make this the current OpenGL projection matrix. Another way to generate an oblique-projection view is to rotate the scene into an appropriate position so that an orthogonal projection in the `zview` direction yields the desired view.
OpenGL Symmetric Perspective-Projection Function

There are two functions available for producing a perspective-projection view of a scene. One of these functions generates a symmetric frustum view volume about the viewing direction (the negative $z_{\text{view}}$ axis). The other function can be used for either a symmetric-perspective projection or an oblique-perspective projection. For both functions, the projection reference point is the viewing-coordinate origin and the near clipping plane is the view plane.

A symmetric, perspective-projection, frustum view volume is set up with the GLU function

\[ \text{gluPerspective} (\theta, \text{aspect}, d_{\text{near}}, d_{\text{far}}); \]

with each of the four parameters assigned a double-precision, floating-point number. The first two parameters define the size and position of the clipping window on the near plane, and the second two parameters specify the distances from the view point (coordinate origin) to the near and far clipping planes. Parameter $\theta$ represents the field-of-view angle, which is the angle between the top and bottom clipping planes (Figure 41). This angle can be assigned any value from $0^\circ$ to $180^\circ$. Parameter aspect is assigned a value for the aspect ratio ($width/height$) of the clipping window.

For a perspective projection in OpenGL, the near and far clipping planes must always be somewhere along the negative $z_{\text{view}}$ axis; neither can be “behind” the viewing position. This restriction does not apply to an orthogonal projection, but it precludes the inverted perspective projection of an object when the view plane is behind the view point. Therefore, both $d_{\text{near}}$ and $d_{\text{far}}$ must be assigned positive numerical values, and the positions of the near and far planes are calculated as $z_{\text{near}} = -d_{\text{near}}$ and $z_{\text{far}} = -d_{\text{far}}$.

If we do not specify a projection function, our scene is displayed using the default orthogonal projection. In this case, the view volume is the symmetric normalized cube shown in Figure 47.

The frustum view volume set up by the \text{gluPerspective} function is symmetric about the negative $z_{\text{view}}$ axis. And the description of a scene is converted to normalized, homogeneous projection coordinates with matrix 41.

OpenGL General Perspective-Projection Function

We can use the following function to specify a perspective projection that has either a symmetric frustum view volume or an oblique frustum view volume.

---

**Figure 47**
Default orthogonal-projection view volume. Coordinate extents for this symmetric cube are from $-1$ to $+1$ in each direction. The near clipping plane is at $z_{\text{near}} = 1$, and the far clipping plane is at $z_{\text{far}} = -1$. 

---

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\[ \text{glFrustum (xwmin, xwmax, ywmin, ywmax, dnear, dfar);} \]

All parameters in this function are assigned double-precision, floating-point numbers. As in the other viewing-projection functions, the near plane is the view plane and the projection reference point is at the viewing position (coordinate origin). This function has the same parameters as the orthogonal, parallel-projection function, but now the near and far clipping-plane distances must be positive. The first four parameters set the coordinates for the clipping window on the near plane, and the last two parameters specify the distances from the coordinate origin to the near and far clipping planes along the negative \( z_{\text{view}} \) axis. Locations for the near and far planes are calculated as \( z_{\text{near}} = -dnear \) and \( z_{\text{far}} = -dfar \).

The clipping window can be specified anywhere on the near plane. If we select the clipping window coordinates so that \( xw_{\text{min}} = -xw_{\text{max}} \) and \( yw_{\text{min}} = -yw_{\text{max}} \), we obtain a symmetric frustum (about the negative \( z_{\text{view}} \) axis as its centerline).

Again, if we do not explicitly invoke a projection command, OpenGL applies the default orthogonal projection to the scene. The view volume in this case is the symmetric cube (Figure 47).

OpenGL Viewports and Display Windows

After the clipping routines have been applied in normalized coordinates, the contents of the normalized clipping window, along with the depth information, are transferred to three-dimensional screen coordinates. The color value for each \( xy \) position on the viewport is stored in the refresh buffer (color buffer), and the depth information for each \( xy \) position is stored in the depth buffer.

A rectangular viewport is defined with the following OpenGL function.

\[ \text{glViewport (xvmin, yvmin, vpWidth, vpHeight);} \]

The first two parameters in this function specify the integer screen position of the lower-left corner of the viewport relative to the lower-left corner of the display window. And the last two parameters give the integer width and height of the viewport. To maintain the proportions of objects in a scene, we set the aspect ratio of the viewport equal to the aspect ratio of the clipping window.

Display windows are created and managed with GLUT routines. The default viewport in OpenGL is the size and position of the current display window.

OpenGL Three-Dimensional Viewing Program Example

A perspective-projection view of a square, as shown in Figure 48, is displayed using the following program example. The square is defined in the \( xy \) plane, and a viewing-coordinate origin is selected to view the front face at an angle. Choosing the center of the square as the look-at point, we obtain a perspective view using the \text{glFrustum} function. If we move the viewing origin around to the other side of the polygon, the back face would be displayed as a wire-frame object.
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FIGURE 48
Output display generated by the
three-dimensional viewing example
program.

```
#include <GL/glut.h>

GLint winWidth = 600, winHeight = 600; // Initial display-window size.
GLfloat x0 = 100.0, y0 = 50.0, z0 = 50.0; // Viewing-coordinate origin.
GLfloat xref = 50.0, yref = 50.0, zref = 0.0; // Look-at point.
GLfloat Vx = 0.0, Vy = 1.0, Vz = 0.0; // View-up vector.

/* Set coordinate limits for the clipping window: */
GLfloat xwMin = -40.0, ywMin = -60.0, xwMax = 40.0, ywMax = 60.0;

/* Set positions for near and far clipping planes: */
GLfloat dnear = 25.0, dfar = 125.0;

void init (void)
{
    glClearColor (1.0, 1.0, 1.0, 0.0);
    glMatrixMode (GL_MODELVIEW);
    gluLookAt (x0, y0, z0, xref, yref, zref, Vx, Vy, Vz);
    glMatrixMode (GL_PROJECTION);
    glFrustum (xwMin, xwMax, ywMin, ywMax, dnear, dfar);
}

void displayFcn (void)
{
    glClear (GL_COLOR_BUFFER_BIT);
```

/* Set parameters for a square fill area. */
glColor3f (0.0, 1.0, 0.0); // Set fill color to green.
glPolygonMode (GL_FRONT, GL_FILL);
glPolygonMode (GL_BACK, GL_LINE); // Wire-frame back face.
gBegin (GL_QUADS);
    glVertex3f (0.0, 0.0, 0.0);
    glVertex3f (100.0, 0.0, 0.0);
    glVertex3f (100.0, 100.0, 0.0);
    glVertex3f (0.0, 100.0, 0.0);
End ( );

flush ( );

void reshapeFcn (GLint newWidth, GLint newHeight)
{  
    glViewport (0, 0, newWidth, newHeight);
    winWidth = newWidth;
    winHeight = newHeight;
}

void main (int argc, char** argv)
{  
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowPosition (50, 50);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Perspective View of A Square");
    init ( );
    glutDisplayFunc (displayFcn);
    glutReshapeFunc (reshapeFcn);
    glutMainLoop ( );
}

11 Three-Dimensional Clipping Algorithms

Previously, we discussed the advantages of using the normalized boundaries of the clipping window in two-dimensional clipping algorithms. Similarly, we can apply three-dimensional clipping algorithms to the normalized boundaries of the view volume. This allows the viewing pipeline and the clipping procedures to be implemented in a highly efficient way. All device-independent transformations (geometric and viewing) are concatenated and applied before executing the clipping routines. And each of the clipping boundaries for the normalized view volume is a plane that is parallel to one of the Cartesian planes, regardless of the projection type and original shape of the view volume. Depending on whether the view volume has been normalized to a unit cube or to a symmetric cube with edge length 2, the clipping planes have coordinate positions either at 0 and 1 or at −1 and 1. For the symmetric cube, the equations for the three-dimensional
clipping planes are

\[
\begin{align*}
x_{\min} &= -1, & x_{\max} &= 1 \\
y_{\min} &= -1, & y_{\max} &= 1 \\
z_{\min} &= -1, & z_{\max} &= 1
\end{align*}
\]  

(43)

The \(x\) and \(y\) clipping boundaries are the normalized limits for the clipping window, and the \(z\) clipping boundaries are the normalized positions for the near and far clipping planes.

Clipping algorithms for three-dimensional viewing identify and save all object sections within the normalized view volume for display on the output device. All parts of objects that are outside the view-volume clipping planes are eliminated. And the algorithms are now extensions of two-dimensional methods, using the normalized boundary planes of the view volume instead of the straight-line boundaries of the normalized clipping window.

**Clipping in Three-Dimensional Homogeneous Coordinates**

Computer-graphics libraries process spatial positions as four-dimensional homogeneous coordinates so that all transformations can be represented as 4 by 4 matrices. As each coordinate position enters the viewing pipeline, it is converted to a four-dimensional representation:

\[(x, y, z) \rightarrow (x, y, z, 1)\]

After a position has passed through the geometric, viewing, and projection transformations, it is now in the homogeneous form

\[
\begin{bmatrix}
x_h \\y_h \\z_h \\h
\end{bmatrix} = M \cdot \begin{bmatrix} x \\y \\z \\
1
\end{bmatrix}
\]

(44)

where matrix \(M\) represents the concatenation of all the various transformations from world coordinates to normalized, homogeneous projection coordinates, and the homogeneous parameter \(h\) may no longer have the value 1. In fact, \(h\) can have any real value, depending on how we represented objects in the scene and the type of projection we used.

If the homogeneous parameter \(h\) does have the value 1, the homogeneous coordinates are the same as the Cartesian projection coordinates. This is often the case for a parallel-projection transformation. But a perspective projection produces a homogeneous parameter that is a function of the \(z\) coordinate for any spatial position. The perspective-projection homogeneous parameter can even be negative. This occurs when coordinate positions are behind the projection reference point. Also, rational spline representations for object surfaces are often formulated in homogeneous coordinates, where the homogeneous parameter can be positive or negative. Therefore, if clipping is performed in projection coordinates after division by the homogeneous parameter \(h\), some coordinate information can be lost and objects may not be clipped correctly.

An effective method for dealing with all possible projection transformations and object representations is to apply the clipping routines to the homogeneous-coordinate representations of spatial positions. And, because all view volumes can be converted to a normalized cube, a single clipping procedure can be implemented in hardware to clip objects in homogeneous coordinates against the normalized clipping planes.
Three-Dimensional Region Codes

We extend the concept of a region code to three dimensions by simply adding a couple of additional bit positions to accommodate the near and far clipping planes. Thus, we now use a six-bit region code, as illustrated in Figure 49. Bit positions in this region-code example are numbered from right to left, referencing the left, right, bottom, top, near, and far clipping planes, in that order.

For a three-dimensional scene, we need to apply the clipping routines to the projection coordinates, which have been transformed to a normalized space. After the projection transformation, each point in a scene has the four-component representation \( P = (x_h, y_h, z_h, h) \). Assuming that we are clipping against the boundaries of the normalized symmetric cube (Eqs. 43), then a point is inside this normalized view volume if the projection coordinates of the point satisfy the following six inequalities:

\[
-1 \leq \frac{x_h}{h} \leq 1, \quad -1 \leq \frac{y_h}{h} \leq 1, \quad -1 \leq \frac{z_h}{h} \leq 1
\] (45)

Unless we have encountered an error, the value of the homogeneous parameter \( h \) is nonzero. Before implementing region-code procedures, we can first check for the possibility of a homogeneous parameter with either a zero value or an extremely small magnitude. Also, the homogeneous parameter can be either positive or negative. Therefore, assuming \( h \neq 0 \), we can write the preceding inequalities in the form

\[
-h \leq x_h \leq h, \quad -h \leq y_h \leq h, \quad -h \leq z_h \leq h \quad \text{if } h > 0
\]

\[
h \leq x_h \leq -h, \quad h \leq y_h \leq -h, \quad h \leq z_h \leq -h \quad \text{if } h < 0
\] (46)

In most cases \( h > 0 \), and we can then assign the bit values in the region code for a coordinate position according to the tests:

- bit 1 = 1 if \( h + x_h < 0 \) (left)
- bit 2 = 1 if \( h - x_h < 0 \) (right)
- bit 3 = 1 if \( h + y_h < 0 \) (bottom)
- bit 4 = 1 if \( h - y_h < 0 \) (top)
- bit 5 = 1 if \( h + z_h < 0 \) (near)
- bit 6 = 1 if \( h - z_h < 0 \) (far)

These bit values can be set using the same approach as in two-dimensional clipping. That is, we simply use the sign bit of one of the calculations \( h \pm x_h \), \( h \pm y_h \), or \( h \pm z_h \) to set the corresponding region-code bit value. Figure 50 lists the 27 region codes for a view volume. In those cases where \( h < 0 \) for some point, we could apply clipping using the second set of inequalities in 46 or we could negate the coordinates and clip using the tests for \( h > 0 \).
Three-Dimensional Point and Line Clipping

For standard point positions and straight-line segments that are defined in a scene that is not behind the projection reference point, all homogeneous parameters are positive and the region codes can be established using the conditions in 47. Then, once we have set up the region code for each position in a scene, we can easily identify a point position as outside the view volume or inside the view volume. For instance, a region code of 101000 tells us that the point is above and directly behind the view volume, while the region code 000000 indicates a point within the volume (Figure 50). Thus, for point clipping, we simply eliminate any individual point whose region code is not 000000. In other words, if any one of the tests in 47 is negative, the point is outside the view volume.

Methods for three-dimensional line clipping are essentially the same as for two-dimensional lines. We can first test the line endpoint region codes for trivial acceptance or rejection of the line. If the region code for both endpoints of a line is 000000, the line is completely inside the view volume. Equivalently, we can trivially accept the line if the logical or operation on the two endpoint region codes produces a value of 0. And we can trivially reject the line if the logical and operation on the two endpoint region codes produces a value that is not 0. This nonzero value indicates that both endpoint region codes have a 1 value in the same bit position, and hence the line is completely outside one of the clipping planes. As an example of this, the line from $P_3$ to $P_4$ in Figure 51 has the endpoint region-code values of 010101 and 100110. So this line is completely below the bottom clipping plane. If a line fails these two tests, we next analyze the line equation to determine whether any part of the line should be saved.
Equations for three-dimensional line segments are conveniently expressed in parametric form, and the clipping methods of Cyrus-Beck or Liang-Barsky can be extended to three-dimensional scenes. For a line segment with endpoints \( P_1 = (x_{h1}, y_{h1}, z_{h1}, h_1) \) and \( P_2 = (x_{h2}, y_{h2}, z_{h2}, h_2) \), we can write the parametric equation describing any point position along the line as

\[
P = P_1 + (P_2 - P_1)u \quad 0 \leq u \leq 1
\] (48)

When the line parameter has the value \( u = 0 \), we are at position \( P_1 \). And \( u = 1 \) brings us to the other end of the line, \( P_2 \). Writing the parametric line equation explicitly, in terms of the homogeneous coordinates, we have

\[
\begin{align*}
x_h &= x_{h1} + (x_{h2} - x_{h1})u \\
y_h &= y_{h1} + (y_{h2} - y_{h1})u \\
z_h &= z_{h1} + (z_{h2} - z_{h1})u \\
h &= h_1 + (h_2 - h_1)u
\end{align*}
\] (49)

Using the endpoint region codes for a line segment, we can first determine which clipping planes are intersected. If one of the endpoint region codes has a 0 value in a certain bit position while the other code has a 1 value in the same bit position, then the line crosses that clipping boundary. In other words, one of the tests in 47 generates a negative value, while the same test for the other endpoint of the line produces a nonnegative value. To find the intersection position with this clipping plane, we first use the appropriate equations in 49 to determine the corresponding value of parameter \( u \). Then we calculate the intersection coordinates.

As an example of the intersection-calculation procedure, we consider the line segment \( P_1P_2 \) in Figure 51. This line intersects the right clipping plane, which can be described with the equation \( x_{\text{max}} = 1 \). Therefore, we determine the intersection value for parameter \( u \) by setting the \( x \)-projection coordinate equal to 1:

\[
x_p = \frac{x_h}{h} = \frac{x_{h1} + (x_{h2} - x_{h1})u}{h_1 + (h_2 - h_1)u} = 1
\] (50)

Solving for parameter \( u \), we obtain

\[
u = \frac{x_{h1} - h_1}{(x_{h1} - h_1) - (x_{h2} - h_2)}
\] (51)
Next, we determine the values $y_p$ and $z_p$ on this clipping plane, using the calculated value for $u$. In this case, the $y_p$ and $z_p$ intersection values are within the $\pm 1$ boundaries of the view volume and the line does cross into the view-volume interior. So we next proceed to locate the intersection position with the top clipping plane. That completes the processing for this line segment, because the intersection points with the top and right clipping planes identify the part of the line that is inside the view volume and all the line sections that are outside the view volume.

When a line intersects a clipping boundary but does not enter the view-volume interior, we continue the line processing as in two-dimensional clipping. The section of the line outside that clipping boundary is eliminated, and we update the region-code information and the values for parameter $u$ for the part of the line inside that boundary. Then we test the remaining section of the line against the other clipping planes for possible rejection or for further intersection calculations.

Line segments in three-dimensional scenes are usually not isolated. They are most often components in the description for the solid objects in the scene, and we need to process the lines as part of the surface-clipping routines.

### Three-Dimensional Polygon Clipping

Graphics packages typically deal only with scenes that contain “graphics objects.” These are objects whose boundaries are described with linear equations, so that each object is composed of a set of surface polygons. Therefore, to clip objects in a three-dimensional scene, we apply the clipping routines to the polygon surfaces. Figure 52, for example, highlights the surface sections of a pyramid that are to be clipped, and the dashed lines show sections of the polygon surfaces that are inside the view volume.

We can first test a polyhedron for trivial acceptance or rejection using its coordinate extents, a bounding sphere, or some other measure of its coordinate limits. If the coordinate limits of the object are inside all clipping boundaries, we save the entire object. If the coordinate limits are all outside any one of the clipping boundaries, we eliminate the entire object.

When we cannot save or eliminate the entire object, we can next process the vertex lists for the set of polygons that define the object surfaces. Applying

---

**Figure 52**

Three-dimensional object clipping. Surface sections that are outside the view-volume clipping planes are eliminated from the object description, and new surface facets may need to be constructed.
methods similar to those in two-dimensional polygon clipping, we can clip edges to obtain new vertex lists for the object surfaces. We may also need to create some new vertex lists for additional surfaces that result from the clipping operations. And the polygon tables are updated to add any new polygon surfaces and to revise the connectivity and shared-edge information about the surfaces.

To simplify the clipping of general polyhedra, polygon surfaces are often divided into triangular sections and described with triangle strips. We can then clip the triangle strips using the Sutherland-Hodgman approach. Each triangle strip is processed in turn against the six clipping planes to obtain the final vertex list for the strip.

For concave polygons, we can apply splitting methods to obtain a set of triangles, for example, and then clip the triangles. Alternatively, we could clip three-dimensional concave polygons using the Weiler-Atherton algorithm.

### Three-Dimensional Curve Clipping

As in polyhedra clipping, we first check to determine whether the coordinate extents of a curved object, such as a sphere or a spline surface, are completely inside the view volume. Then we can check to determine whether the object is completely outside any one of the six clipping planes.

If the trivial rejection-acceptance tests fail, we locate the intersections with the clipping planes. To do this, we solve the simultaneous set of surface equations and the clipping-plane equation. For this reason, most graphics packages do not include clipping routines for curved objects. Instead, curved surfaces are approximated as a set of polygon patches, and the objects are then clipped using polygon-clipping routines. When surface-rendering procedures are applied to polygon patches, they can provide a highly realistic display of a curved surface.

### Arbitrary Clipping Planes

It is also possible, in some graphics packages, to clip a three-dimensional scene using additional planes that can be specified in any spatial orientation. This option is useful in a variety of applications. For example, we might want to isolate or clip off an irregularly shaped object, eliminate part of a scene at an oblique angle for a special effect, or slice off a section of an object along a selected axis to show a cross-sectional view of its interior.

Optional clipping planes can be specified along with the description of a scene, so that the clipping operations can be performed prior to the projection transformation. However, this also means that the clipping routines are implemented in software.

A clipping plane can be specified with the plane parameters \(A, B, C,\) and \(D\). The plane then divides three-dimensional space into two parts, so that all parts of a scene that lie on one side of the plane are clipped off. Assuming that objects behind the plane are to be clipped, then any spatial position \((x, y, z)\) that satisfies the following inequality is eliminated from the scene.

\[
Ax + By + Cz + D < 0
\]  

(52)

As an example, if the plane-parameter array has the values \((A, B, C, D) = (1.0, 0.0, 0.0, 8.0)\), then any coordinate position satisfying \(x + 8.0 < 0.0\) (or, \(x < -8.0\)) is clipped from the scene.

To clip a line segment, we can first test its two endpoints to see if the line is completely behind the clipping plane or completely in front of the plane. We can represent inequality 52 in a vector form using the plane normal vector.
FIGURE 53
Clipping a line segment against a plane with normal vector \( \mathbf{N} \).

\[ \mathbf{N} = (A, B, C) \]. Then, for a line segment with endpoint positions \( \mathbf{P}_1 \) and \( \mathbf{P}_2 \), we clip the entire line if both endpoints satisfy

\[ \mathbf{N} \cdot \mathbf{P}_k + D < 0, \quad k = 1, 2 \]  

(53)

We save the entire line if both endpoints satisfy

\[ \mathbf{N} \cdot \mathbf{P}_k + D \geq 0, \quad k = 1, 2 \]  

(54)

Otherwise, the endpoints are on opposite sides of the clipping plane, as in Figure 53, and we calculate the line intersection point.

To calculate the line-intersection point with the clipping plane, we can use the following parametric representation for the line segment:

\[ \mathbf{P} = \mathbf{P}_1 + (\mathbf{P}_2 - \mathbf{P}_1)u, \quad 0 \leq u \leq 1 \]  

(55)

Point \( \mathbf{P} \) is on the clipping plane if it satisfies the plane equation

\[ \mathbf{N} \cdot \mathbf{P} + D = 0 \]  

(56)

Substituting the expression for \( \mathbf{P} \) from Equation 55, we have

\[ \mathbf{N} \cdot [\mathbf{P}_1 + (\mathbf{P}_2 - \mathbf{P}_1)u] + D = 0 \]  

(57)

Solving this equation for parameter \( u \), we obtain

\[ u = \frac{-D - \mathbf{N} \cdot \mathbf{P}_1}{\mathbf{N} \cdot (\mathbf{P}_2 - \mathbf{P}_1)} \]  

(58)

We then substitute this value of \( u \) into the vector parametric line representation 55 to obtain values for the \( x \), \( y \), and \( z \) intersection coordinates. For the example in Figure 53, the line segment from \( \mathbf{P}_1 \) to \( \mathbf{P} \) is clipped and we save the section of the line from \( \mathbf{P}_1 \) to \( \mathbf{P}_2 \).

For polyhedra, such as the pyramid in Figure 54, we apply similar clipping procedures. We first test to see if the object is completely behind or completely in front of the clipping plane. If not, we process the vertex list for each polygon surface. Line-clipping methods are applied to each polygon edge in succession, just as in view-volume clipping, to produce the surface vertex lists. But in this case, we have to deal with only one clipping plane.

Clipping a curved object against a single clipping plane is easier than clipping the object against the six planes of a view volume. However, we still need to solve a set of nonlinear equations to locate intersections, unless we approximate the curve boundaries with straight-line sections.
**12 OpenGL Optional Clipping Planes**

In addition to the six clipping planes enclosing the view volume, OpenGL provides for the specification of additional clipping planes in a scene. Unlike the view-volume clipping planes, which are each perpendicular to one of the coordinate axes, these additional planes can have any orientation.

We designate an optional clipping plane and activate clipping against that plane with the statements

```c
glClipPlane (id, planeParameters);
glEnable (id);
```

Parameter `id` is used as an identifier for a clipping plane. This parameter is assigned one of the values `GL_CLIP_PLANE0`, `GL_CLIP_PLANE1`, and so forth, up to a facility-defined maximum. The plane is then defined using the four-element array `planeParameters`, whose elements are the double-precision, floating-point values for the four plane-equation parameters `A`, `B`, `C`, and `D`. An activated clipping plane that has been assigned the identifier `id` is turned off with

```c
glDisable (id);
```

The plane parameters `A`, `B`, `C`, and `D` are transformed to viewing coordinates and used to test viewing-coordinate positions in a scene. Subsequent changes in viewing or geometric-transformation parameters do not affect the stored plane parameters. Therefore, if we set up optional clipping planes before specifying any geometric or viewing transformations, the stored plane parameters are the same as the input parameters. Also, because the clipping routines for these planes are applied in viewing coordinates, and not in the normalized coordinate space, the performance of a program can be degraded when optional clipping planes are activated.
Any points that are “behind” an activated OpenGL clipping plane are eliminated. Thus, a viewing-coordinate position \((x, y, z)\) is clipped if it satisfies condition 52.

Six optional clipping planes are available in any OpenGL implementation, but more might be provided. We can find out how many optional clipping planes are possible for a particular OpenGL implementation with the inquiry

\[
glGetIntegerv (GL_MAX_CLIP_PLANES, numPlanes);
\]

Parameter \(numPlanes\) is the name of an integer array that is to be assigned an integer value equal to the number of optional clipping planes that we can use.

The default for the \(glClipPlane\) function is that the clipping-plane parameters \(A, B, C,\) and \(D\) are each assigned a value of 0 for all optional planes. And, initially, all optional clipping planes are disabled.

## 13 Summary

Viewing procedures for three-dimensional scenes follow the general approach used in two-dimensional viewing. We first create a world-coordinate scene, either from the definitions of objects in modeling coordinates or directly in world coordinates. Then we set up a viewing-coordinate reference frame and transfer object descriptions from world coordinates to viewing coordinates. Object descriptions are then processed through various routines to device coordinates.

Unlike two-dimensional viewing, however, three-dimensional viewing requires projection routines to transform object descriptions to a viewing plane before the transformation to device coordinates. Also, three-dimensional viewing operations involve more spatial parameters. We can use the camera analogy to describe three-dimensional viewing parameters. A viewing-coordinate reference frame is established with a view reference point (the camera position), a view-plane normal vector \(N\) (the camera lens direction), and a view-up vector \(V\) (the camera up direction). The view-plane position is then established along the viewing \(z\) axis, and object descriptions are projected to this plane. Either parallel-projection or perspective-projection methods can be used to transfer object descriptions to the view plane.

Parallel projections are either orthographic or oblique, and they can be specified with a projection vector. Orthographic parallel projections that display more than one face of an object are called axonometric projections. An isometric view of an object is obtained with an axonometric projection that foreshortens each principal axis by the same amount. Commonly used oblique projections are the cavalier projection and the cabinet projection. Perspective projections of objects are obtained with projection lines that meet at the projection reference point. Parallel projections maintain object proportions, but perspective projections decrease the size of distant objects. Perspective projections cause parallel lines to appear to converge to a vanishing point, provided the lines are not parallel to the view plane. Engineering and architectural displays can be generated with one-point, two-point, or three-point perspective projections, depending on the number of principal axes that intersect the view plane. An oblique perspective projection is obtained when the line from the projection reference point to the center of the clipping window is not perpendicular to the view plane.

Objects in a three-dimensional scene can be clipped against a view volume to eliminate unwanted sections of the scene. The top, bottom, and sides of the view volume are formed with planes that are parallel to the projection lines and that pass through the clipping-window edges. Near and far planes (also called front and back planes) are used to create a closed view volume. For a parallel
projection, the view volume is a parallelepiped. For a perspective projection, the view volume is a frustum. In either case, we can convert the view volume to a normalized cube with boundaries either at 0 and 1 for each coordinate or at −1 and 1 for each coordinate. Efficient clipping algorithms process objects in a scene against the bounding planes of the normalized view volume. Clipping is generally carried out in graphics packages in four-dimensional homogeneous coordinates following the projection and view-volume normalization transformations. Then, homogeneous coordinates are converted to three-dimensional, Cartesian projection coordinates. Additional clipping planes, with arbitrary orientations, can also be used to eliminate selected parts of a scene or to produce special effects.

A function is available in the OpenGL Utility library for specifying three-dimensional viewing parameters (see Table 1). This library also includes a function for setting up a symmetric perspective-projection transformation. Three other viewing functions are available in the OpenGL basic library for specifying an orthographic projection, a general perspective projection, and optional clipping planes. Table 1 summarizes the OpenGL viewing functions discussed in this chapter. In addition, the table lists some viewing-related functions.

REFERENCES

A complete listing of three-dimensional OpenGL viewing functions is given in Shreiner (2000). For OpenGL programming examples using three-dimensional viewing, see Woo, et al. (1999). Additional programming examples can be found at Nate Robins’s tutorial website: www.xmission.com/~nate/opengl.html.

EXERCISES
1. Write a procedure to set up the matrix that transforms world-coordinate positions to three-dimensional viewing coordinates, given $P_0$, $N$, and $V$. The view-up vector can be in any direction that is not parallel to $N$.
2. Write a procedure to transform the vertices of a polyhedron to projection coordinates using a parallel projection with any specified projection vector.
3. Write a program to obtain different parallel-projection views of a polyhedron by allowing the user to rotate the polyhedron via the keyboard.
4. Write a procedure to perform a one-point perspective projection of an object.
Three-Dimensional Viewing

5 Write a procedure to perform a two-point perspective projection of an object.
6 Develop a routine to perform a three-point perspective projection of an object.
7 Write a program that uses the routines in the previous three exercises to display a three-dimensional cube using a one-, two-, or three-point perspective projection according to input taken from the keyboard, which should be used to switch between projections. The program should also allow the user to rotate the cube in the \(xz\) plane around its center. Examine the visual differences of the three different types of projections.
8 Write a routine to convert a perspective projection frustum to a regular parallelepiped.
9 Modify the two-dimensional Cohen-Sutherland line-clipping algorithm to clip three-dimensional lines against the normalized symmetric view volume square.
10 Write a program to generate a set of 10 random lines, each of which has one endpoint within a normalized symmetric view volume and one without. Implement the three-dimensional Cohen-Sutherland line-clipping algorithm designed in the previous exercise to clip the set of lines against the viewing volume.
11 Modify the two-dimensional Liang-Barsky line-clipping algorithm to clip three-dimensional lines against a specified regular parallelepiped.
12 Write a program similar to that in Exercise 10 that generates a set of 10 random lines, each partially outside of a specified regular parallelepiped viewing volume. Use the three-dimensional Liang-Barsky line-clipping algorithm developed in the previous exercise to clip the lines against the viewing volume.
13 Modify the two-dimensional Liang-Barsky line-clipping algorithm to clip a given polyhedron against a specified regular parallelepiped.
14 Write a program to display a cube in a regular parallelepiped viewing volume and allow the user to translate the cube along each axis using keyboard input. Implement the algorithm in the previous exercise to clip the cube when it extends over any of the edges of the viewing volume.
15 Write a routine to perform line clipping in homogeneous coordinates.
16 Devise an algorithm to clip a polyhedron against a defined frustum. Compare the operations needed in this algorithm to those needed in an algorithm that clips against a regular parallelepiped.
17 Extend the Sutherland-Hodgman polygon-clipping algorithm to clip a convex polyhedron against a normalized symmetric view volume.
18 Write a routine to implement the preceding exercise.
19 Write a program similar to the one in Exercise 14 to display a cube in a normalized symmetric view volume that can be translated around the viewing volume via keyboard input. Use the implementation of the polygon-clipping algorithm developed in the previous exercise to clip the cube when it extends over the edge of the viewing volume.
20 Write a routine to perform polyhedron clipping in homogeneous coordinates.
21 Modify the program example in Section 10 to allow a user to specify a view for either the front or the back of the square.
22 Modify the program example in Section 10 to allow the perspective viewing parameters to be specified as user input.
23 Modify the program example in Section 10 to produce a view of any input polyhedron.
24 Modify the program in the preceding exercise to generate a view of the polyhedron using an orthographic projection.
25 Modify the program in the preceding exercise to generate a view of the polyhedron using an oblique parallel projection.

IN MORE DEPTH

1 In this exercise, you will give “depth” to the polygons that represent the objects in your scene and clip them against a normalized view volume. First, choose new \(z\)-coordinates and three-dimensional orientations for the polygons in your scene that are appropriate to the snapshot of your application. That is, they should be taken out of the \(xy\) plane in which they have been constrained so far and given appropriate depth. Once you have done this, implement an extension of the Sutherland-Hodgman polygon-clipping algorithm that allows clipping of convex polygons against a normalized symmetric view volume. You will use this algorithm in the next exercise to produce a view of some portion of your three-dimensional scene.
2 Choose a view of the scene from the previous exercise that produces a view volume in which all objects are not fully contained. Apply the algorithm for polygon clipping that you developed in the previous exercise against the view volume. Write routines to display the scene using a parallel projection and a perspective projection. Use the OpenGL three-dimensional viewing functions to do this, choosing appropriate parameters to specify the viewing volume in each case. Allow the user to switch between the two projections via keyboard input and note the differences in the visual appearance of the scene in the two cases.
Three-Dimensional Viewing

Color Plates

Color Plate 9
A realistic room display, achieved with a perspective projection, illumination effects, and selected surface properties. (Courtesy of John Snyder, Jed Lengyel, Devendra Kalra, and Al Barr, California Institute of Technology. © 1992 Caltech.)
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In setting up the definition of a complex object or system, it is usually easiest to specify the subparts first and then describe how the subparts fit together to form the overall object or system. For instance, a bicycle can be described in terms of a frame, wheels, fenders, handlebars, seat, chain, and pedals, along with the rules for positioning these components to form the bicycle. A hierarchical description of this type can be given as a tree structure, consisting of the subparts as the tree nodes and the construction rules as the tree branches.

Architectural and engineering systems, such as building layouts, automobile design, electronic circuits, and home appliances, are now routinely developed using computer-aided design (CAD) packages. And graphical design methods are used also for representing economic, financial, organizational, scientific, social, and environmental systems. Simulations are often constructed to study the behavior of a system under various conditions, and the outcome of a simulation can serve as an instructional tool or as a basis for making decisions.
about the system. Design packages generally provide routines for creating and managing
hierarchical models, and some packages also contain predefined shapes, such as wheels,
doors, gears, shafts, and electric-circuit components.

1 Basic Modeling Concepts

The creation and manipulation of a system representation is termed **modeling**. Any single representation is called a **model** of the system, which could be defined graphically or purely descriptively, such as a set of equations that describe the relationships among system parameters. Graphical models are often referred to as **geometric models**, because the component parts of a system are represented with geometric entities such as straight-line segments, polygons, polyhedra, cylinders, or spheres. Because we are concerned here only with graphics applications, we will use the term model to mean a computer-generated, geometric representation of a system.

System Representations

Figure 1 shows a graphical representation for a logic circuit, illustrating the features common to many system models. Component parts of the system are displayed as geometric structures, called **symbols**, and relationships among the symbols are represented in this example with a network of connecting lines. Three standard symbols are used to represent logic gates for the Boolean operations: **and**, **or**, and **not**. The connecting lines define relationships in terms of input and output flow (from left to right) through the system parts. One symbol, the **and** gate, is displayed at two different positions within the logic circuit. Repeated positioning of a few basic symbols is a common method for building complex models. Each such occurrence of a symbol within a model is called an **instance** of that symbol. We have one instance for the **or** and **not** symbols in Figure 1 and two instances of the **and** symbol.

In many cases, the particular graphical symbols chosen to represent the parts of a system are dictated by the system description. For circuit models, standard electrical or logic symbols are used. But with models representing abstract concepts, such as political, financial, or economic systems, symbols may be any convenient geometric pattern.

Information describing a model is usually provided as a combination of geometric and nongeometric data. Geometric information includes coordinate positions for locating the component parts, output primitives and attribute functions to define the structure of the parts, and data for constructing connections between the parts. Nongeometric information includes text labels, algorithms describing the operating characteristics of the model, and rules for determining the relationships or connections between component parts, if these are not specified as geometric data.

![Figure 1](image-url) Model of a logic circuit.
TABLE 1
Data table defining the structure and position of each gate in the circuit of Figure 1

<table>
<thead>
<tr>
<th>Symbol Code</th>
<th>Geometric Description</th>
<th>Identifying Label</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gate 1</td>
<td>(Coordinates and other parameters)</td>
<td>and</td>
</tr>
<tr>
<td>Gate 2</td>
<td>:</td>
<td>or</td>
</tr>
<tr>
<td>Gate 3</td>
<td>:</td>
<td>not</td>
</tr>
<tr>
<td>Gate 4</td>
<td>:</td>
<td>and</td>
</tr>
</tbody>
</table>

There are two methods for specifying the information needed to construct and manipulate a model. One method is to store the information in a data structure, such as a table or linked list. The other method is to specify the information in procedures. In general, a model specification will contain both data structures and procedures, although some models are defined completely with data structures and others use only procedural specifications. An application to perform solid modeling of objects might use mostly information taken from some data structure to define coordinate positions, with very few procedures. A weather model, on the other hand, may need mostly procedures to calculate plots of temperature and pressure variations.

As an example of how combinations of data structures and procedures can be used, we consider some alternative model specifications for the logic circuit of Figure 1. One method is to define the logic components in a data table (Table 1), with processing procedures used to specify how the network connections are to be made and how the circuit operates. Geometric data in this table include coordinates and parameters necessary for drawing and positioning the gates. These symbols could all be drawn as polygons, or they could be formed as combinations of straight-line segments and elliptical arcs. Labels for each of the component parts also have been included in the table, although the labels could be omitted if the symbols are displayed as commonly recognized shapes. Procedures would then be used to display the gates and construct the connecting lines, based on the coordinate positions of the gates and a specified order for connecting them. An additional procedure is used to produce the circuit output (binary values) for any given input. This procedure could be set up to display only the final output, or it could be designed to display intermediate output values to illustrate the internal functioning of the circuit.

Alternatively, we might specify graphical information for the circuit model in data structures. The connecting lines, as well as the gates, could then be defined in a data table that explicitly lists endpoints for each of the lines in the circuit. A single procedure might then display the circuit and calculate the output. At the other extreme, we could completely define the model in procedures, using no external data structures.

Symbol Hierarchies

Many models can be organized as a hierarchy of symbols. The basic elements for the model are defined as simple geometric shapes appropriate to the type of model under consideration. These basic symbols can be used to form composite objects, sometimes called modules, which themselves can be grouped to form
higher-level objects, and so on, for the various components of the model. In the simplest case, we can describe a model by a one-level hierarchy of component parts, as in Figure 2. For this circuit example, we assume that the gates are positioned and connected to each other with straight lines according to connection rules that are specified with each gate description. The basic symbols in this hierarchical description are the logic gates. Although the gates themselves could be described as hierarchies—formed from straight lines, elliptical arcs, and text—that description would not be a convenient one for constructing logic circuits, in which the simplest building blocks are gates. For an application in which we were interested in designing different geometric shapes, the basic symbols could be defined as straight-line segments and arcs.

An example of a two-level symbol hierarchy appears in Figure 3. Here, a facility layout is planned as an arrangement of work areas. Each work area is outfitted with a collection of furniture. The basic symbols are the furniture items: worktable, chair, shelves, file cabinet, and so forth. Higher-order objects are the work areas, which are put together with different furniture organizations. An instance of a basic symbol is defined by specifying its position, size, and orientation within each work area. Positions are given as coordinate locations in the work areas, and orientations are specified as rotations that determine which way the symbols are facing. At the first level below the root node for the facility tree, each work area is defined by specifying its position, size, and orientation within the facility layout. The boundary for each work area might be defined with a divider that encloses the work area and provides aisles within the facility.

More complex symbol hierarchies are formed with repeated groupings of symbol clusters at each higher level. The facility layout of Figure 3 could be extended to include symbol clusters that form different rooms, different floors
of a building, different buildings within a complex, and different complexes at widely separated geographical locations.

## 2 Modeling Packages

Although system models can be designed and manipulated using a general computer-graphics package, specialized modeling systems are available to facilitate modeling in particular applications. Modeling systems provide a means for defining and rearranging model representations in terms of symbol hierarchies, which are then processed by graphics routines for display. General-purpose graphics systems often do not provide routines to accommodate extensive modeling applications. But some graphics packages, such as GL and PHIGS, do include integrated sets of modeling and graphics functions.

If a graphics library contains no modeling functions, we can often use a modeling-package interface to the graphics routines. Alternatively, we could create our own modeling routines using the geometric transformations and other functions available in the graphics library.

Specialized modeling packages, such as some CAD systems, are defined and structured according to the type of application the package has been designed to handle. These packages provide menus of symbol shapes and functions for the intended application. And they can be designed for either two-dimensional or three-dimensional modeling.

## 3 General Hierarchical Modeling Methods

We create a hierarchical model of a system by nesting the descriptions of its subparts into one another to form a tree organization. As each node is placed into the hierarchy, it is assigned a set of transformations to position it appropriately into the overall model. For an office-facility design, work areas and offices are formed with arrangements of furniture items. The offices and work areas are then placed into departments, and so forth on up the hierarchy. An example of the use of multiple coordinate systems and hierarchical modeling with three-dimensional objects is given in Figure 4. This figure illustrates simulation of tractor movement. As the tractor moves, the tractor coordinate system and front-wheel coordinate system move in the world coordinate system. The front wheels rotate in the wheel system, and the wheel system rotates in the tractor system when the tractor turns.

**FIGURE 4**
Possible coordinate systems used in simulating tractor movement. A rotation of the front-wheel system causes the tractor to turn. Both the wheel and tractor reference frames move in the world coordinate system.
Local Coordinates

In general design applications, models are constructed with instances (transformed copies) of the geometric shapes that are defined in a basic symbol set. Each instance is positioned, with the proper orientation, in the world-coordinate reference of the overall structure of the model. The various graphical objects to be used in an application are each defined relative to the world-coordinate reference system, which is referred to as the local coordinate system for that object. Local coordinates are also called modeling coordinates, or sometimes master coordinates. Figure 5 illustrates local-coordinate definitions for two symbols that could be used in a two-dimensional facility-layout application.

Modeling Transformations

To construct a graphical model, we apply transformations to the local-coordinate definitions of symbols to produce instances of the symbols within the overall structure of the model. Transformations applied to the modeling-coordinate definitions of symbols to give them a particular position and orientation within a model are referred to as modeling transformations. The typical transformations available in a modeling package are translation, rotation, and scaling, but other transformations might also be used in some applications.

Creating Hierarchical Structures

A first step in a hierarchical modeling application is to construct modules that are compositions of basic symbols. The modules themselves may then be combined into higher-level modules, and so on. We define each initial module as a list of symbol instances, along with appropriate transformation parameters for each symbol. At the next level, we define each higher-level module as a list of symbol and lower-level module instances along with their transformation parameters. This process is continued up to the root of the tree, which represents the total model in world coordinates.
In a modeling package, a module is created with a sequence of commands of the form

```
createModule1
  setSymbolTransformation1
  insertSymbol1
  setSymbolTransformation2
  insertSymbol2
  .
  .
closeModule1
```

Each instance of a basic symbol is assigned a set of transformation parameters for that module. Similarly, modules are combined to form higher-level modules with functions such as

```
createModule6
  setModuleTransformation1
  insertModule1
  setModuleTransformation2
  insertModule2
  setSymbolTransformation5
  insertSymbol5
  .
  .
closeModule6
```

The transformation function for each module or symbol specifies how that object is to be fitted into the higher-level module. Often, options are provided so that a specified transformation matrix could premultiply, postmultiply, or replace the current transformation matrix.

Although a basic set of symbols could be available in a modeling package, the symbol set might not contain the shapes we need for a particular application. In that case, we can create additional shapes within a modeling program. As an example, the following pseudocode illustrates the specification of a simple model for a bicycle:

```
createWheelSymbol
createFrameSymbol
createBicycleModule
  setFrameTransformation
  insertFrameSymbol
```

Hierarchical Modeling
A number of other modeling routines are usually available in a system designed for hierarchical modeling. Modules often can be selectively displayed or temporarily taken out of a system representation. This allows a designer to experiment with different shapes and design structures. And selected modules could be highlighted or moved around in the display during the design process.

4 Hierarchical Modeling Using OpenGL
Display Lists

Complex objects can be described in OpenGL using nested display lists to form a hierarchical model. Each symbol and module for the model is created with a glNewList function. And we insert one display list into another display list using the glCallList function within the definition of the higher-order list. Geometric transformations can be associated with each inserted object to specify a position, orientation, and size within the higher-level module. As an example, the following code could be used to describe a bicycle that is simply composed of a frame and two identical wheels:

```
glNewList (bicycle, GL_COMPILE);
glCallList (frame);

  glPushMatrix ( );
  glTranslatef (tx1, ty1, tz1);
  glCallList (wheel);
  glPopMatrix ( );

  glPushMatrix ( );
  glTranslatef (tx2, ty2, tz2);
  glCallList (wheel);
  glPopMatrix ( );
glEndList ( );
```

This code creates a new display list which, when executed, will invoke glCallList to execute two additional display lists which will draw the frame and the wheels of the bicycle. Because we are positioning the two wheels relative to the location of the frame of the bicycle, when we draw each wheel we use a call to glPushMatrix before applying the translation which positions the wheel, followed by a call to glPopMatrix after drawing the wheel to restore the transformation matrix to its previous state. This isolates the per-wheel translations; without these calls to glPushMatrix and glPopMatrix, the translations would be cumulative rather than separate—in effect, we would position the
second wheel relative to the position of the first, rather than relative to the position of the frame.

Just as this display list is composed from other lists, so could the frame display list be composed from individual display lists describing the handlebars, chain, pedals, and other components, and the wheel display could be composed from other lists describing the wheel rim, its spokes, and the tire that surrounds the rim.

5 Summary

The term “model,” in computer-graphics applications, refers to a graphical representation for some system. Basic components of a system are represented as symbols, defined in local-coordinate reference frames, which are also referred to as modeling, or master, coordinates. We create a model, such as an electrical circuit, by placing instances of the symbols at selected locations with prescribed orientations.

Many models are constructed as symbol hierarchies. We can construct a hierarchical model by nesting modules, which are composed of instances of basic symbols and other modules. This nesting process may continue down to symbols that are defined with graphical output primitives and their attributes. As each symbol or module is nested within a higher-level module, an associated modeling transformation is specified for the nested structure.

A hierarchical model can be set up in OpenGL using display lists. The glNewList function can be used to define the overall structure of a system and its component modules. Individual symbol structures or other modules are inserted within a module using the glCallList function, preceded by an appropriate set of transformations to specify the position, orientation, and size of the inserted component.

REFERENCE

Examples of modeling applications using OpenGL are given in Woo, et al. (1999).

EXERCISES

1. Discuss model representations that would be appropriate for several distinctly different kinds of systems. Also discuss how graphical representations might be implemented for each system.

2. Devise a two-dimensional neighborhood layout package. A menu of various building structures (e.g., residential buildings, commercial buildings, roads, etc.) is to be provided to a designer, who can use a mouse to select and place an object in any location within a given tract of land that constitutes a neighborhood (a one-level hierarchy). Instance transformations can be limited to translations and rotations.

3. Extend the previous exercise so that building structures can also be scaled along each dimension.

4. Devise a two-dimensional city-planning package that presents a menu of building structure (e.g., residential buildings, commercial buildings, roads, etc.) to a designer. A two-level hierarchy is to be used so that building structures can be placed into various neighborhoods, and the neighborhoods can be arranged within a larger area comprising the land a given city can occupy. Building structures are to be placed into neighborhoods using only translation and rotation instance transformations.

5. Extend the previous exercise so that building structures can also be scaled along each dimension.

6. Write a set of routines for creating and displaying symbols for logic-circuit design. As a minimum, the symbol set should include the and, or, and not gates shown in Figure 1.

7. Develop a modeling package for designing logic circuits that will allow a designer to position
Hierarchical Modeling

electrical symbols within a circuit network. Use the symbol set from the previous exercise, and use only translations to place an instance of one of the menu shapes into the network. Once a component has been placed in the network, it is to be connected to other specified components with straight-line segments.

8 Suppose you are designing a “creature creator” for a video game. The player should be given access to a list of various forms from a fixed set of body parts: heads, bodies, arms, and legs. A creature must have exactly one head and one body, but may have any number of arms or legs. Write a set of routines for editing creatures (a single creature in this instance would be a module) in the proposed game. Your routines should provide for the ability to replace one body part instance for another, insert additional arms or legs, and delete existing arms or legs.

9 Given the coordinate extents of all displayed objects in a model, write a routine to detect any selected object.

10 Write procedures to display and to delete a specified module in a model.

11 Write a routine that will selectively take modules out of a model display or return them to the display.

12 Write a procedure to highlight a selected module in some way. For example, the selected module could be displayed in a different color or it could be enclosed within a rectangular outline.

13 Write a procedure to highlight a selected module in a model by causing the module’s scale to oscillate slightly.

IN MORE DEPTH

1 Recall that in previous exercises, you may have organized a subset of the objects in your scene into a group that behaves in a way that is easier mod-

eled in terms of relative positions and orientations within the group. Then, you used transformations from those local modeling coordinates to world coordinates to convert local transformations of the objects in the group into transformations in the world coordinate system. In this exercise, you will take this concept further, or implement it if you haven’t already. Consider the ways in which the objects in your application interact, and identify groups of objects that make sense to model as a single unit. Alternatively, if none of the objects in your scene exhibit this property, consider modeling single objects in terms of several polygons that change position, scale, or orientation relative to other polygons that make up the object model. This will require that you modify the representation of each object so that it is composed of more than a single polygon. You will learn more about three-dimensional object representations in later chapters, but this temporary solution will allow you to experiment with hierarchical modeling for now. If possible, try to build a hierarchy of two or more levels to get the full appreciation of the utility of organizing objects hierarchically. For each group of objects that comprise a group, identify the subcomponents and how they interact with each other.

2 Use the hierarchical organization of the objects that you developed in the previous exercise to replicate the simple animation of your scene. The transformations that you developed earlier may need to be modified because you altered the coordinates and orientations of the objects in your scene. Set up display lists to define each group and perform the appropriate transformations on each by using the matrix stack appropriately, as described in Section 4. Use the perspective projection viewing scheme that you developed to display the animation in the display window.
Computer graphics methods are now commonly used to produce animations for a variety of applications, including entertainment (motion pictures and cartoons), advertising, scientific and engineering studies, and training and education. Although we tend to think of animation as implying object motion, the term computer animation generally refers to any time sequence of visual changes in a picture. In addition to changing object positions using translations or rotations, a computer-generated animation could display time variations in object size, color, transparency, or surface texture. Advertising animations often transition one object shape into another: for example, transforming a can of motor oil into an automobile engine. We can also generate computer animations by varying camera parameters, such as position, orientation, or focal length, and variations in lighting effects or other parameters and procedures associated with illumination and rendering can be used to produce computer animations.

Another consideration in computer-generated animation is realism. Many applications require realistic displays. An accurate
representation of the shape of a thunderstorm or other natural phenomena described with a numerical model is important for evaluating the reliability of the model. Similarly, simulators for training aircraft pilots and heavy-equipment operators must produce reasonably accurate representations of the environment. Entertainment and advertising applications, on the other hand, are sometimes more interested in visual effects. Thus, scenes may be displayed with exaggerated shapes and unrealistic motions and transformations. However, there are many entertainment and advertising applications that do require accurate representations for computer-generated scenes. Also, in some scientific and engineering studies, realism is not a goal. For example, physical quantities are often displayed with pseudo-colors or abstract shapes that change over time to help the researcher understand the nature of the physical process.

Two basic methods for constructing a motion sequence are real-time animation and frame-by-frame animation. In real-time computer-animation, each stage of the sequence is viewed as it is created. Thus the animation must be generated at a rate that is compatible with the constraints of the refresh rate. For a frame-by-frame animation, each frame of the motion is separately generated and stored. Later, the frames can be recorded on film, or they can be displayed consecutively on a video monitor in “real-time playback” mode. Simple animation displays are generally produced in real time, while more complex animations are constructed more slowly, frame by frame. However, some applications require real-time animation, regardless of the complexity of the animation. A flight-simulator animation, for example, is produced in real time because the video displays must be generated in immediate response to changes in the control settings. In such cases, special hardware and software systems are often developed to allow the complex display sequences to be developed quickly.

1 Raster Methods for Computer Animation

Most of the time, we can create simple animation sequences in our programs using real-time methods. In general, though, we can produce an animation sequence on a raster-scan system one frame at a time, so that each completed frame could be saved in a file for later viewing. The animation can then be viewed by cycling through the completed frame sequence, or the frames could be transferred to film. If we want to generate an animation in real time, however, we need to produce the motion frames quickly enough so that a continuous motion sequence is displayed. For a complex scene, one frame of the animation could take most of the refresh cycle time to construct. In that case, objects generated first would be displayed for most of the frame refresh time, but objects generated toward the end of the refresh cycle would disappear almost as soon as they were displayed. For very complex animations, the frame construction time could be greater than the time to refresh the screen, which can lead to erratic motion and fractured frame displays. Because the screen display is generated from successively modified pixel values in the refresh buffer, we can take advantage of some of the characteristics of the raster screen-refresh process to produce motion sequences quickly.

Double Buffering

One method for producing a real-time animation with a raster system is to employ two refresh buffers. Initially, we create a frame for the animation in one
of the buffers. Then, while the screen is being refreshed from that buffer, we construct the next frame in the other buffer. When that frame is complete, we switch the roles of the two buffers so that the refresh routines use the second buffer during the process of creating the next frame in the first buffer. This alternating buffer process continues throughout the animation. Graphics libraries that permit such operations typically have one function for activating the double-buffering routines and another function for interchanging the roles of the two buffers.

When a call is made to switch two refresh buffers, the interchange could be performed at various times. The most straightforward implementation is to switch the two buffers at the end of the current refresh cycle, during the vertical retrace of the electron beam. If a program can complete the construction of a frame within the time of a refresh cycle, say \( \frac{1}{60} \) of a second, each motion sequence is displayed in synchronization with the screen refresh rate. However, if the time to construct a frame is longer than the refresh time, the current frame is displayed for two or more refresh cycles while the next animation frame is being generated. For example, if the screen refresh rate is 60 frames per second and it takes \( \frac{1}{50} \) of a second to construct an animation frame, each frame is displayed on the screen twice and the animation rate is only 30 frames each second. Similarly, if the frame construction time is \( \frac{1}{25} \) of a second, the animation frame rate is reduced to 20 frames per second because each frame is displayed three times.

Irregular animation frame rates can occur with double buffering when the frame construction time is very nearly equal to an integer multiple of the screen refresh time. As an example of this, if the screen refresh rate is 60 frames per second, then an erratic animation frame rate is possible when the frame construction time is very close to \( \frac{1}{60} \) of a second, or \( \frac{2}{60} \) of a second, or \( \frac{3}{60} \) of a second, and so forth. Because of slight variations in the implementation time for the routines that generate the primitives and their attributes, some frames could take a little more time to construct and some a little less time. Thus, the animation frame rate can change abruptly and erratically. One way to compensate for this effect is to add a small time delay to the program. Another possibility is to alter the motion or scene description to shorten the frame construction time.

### Generating Animations Using Raster Operations

We can also generate real-time raster animations for limited applications using block transfers of a rectangular array of pixel values. This animation technique is often used in game-playing programs. A simple method for translating an object from one location to another in the \( xy \) plane is to transfer the group of pixel values that define the shape of the object to the new location. Two-dimensional rotations in multiples of 90º are also simple to perform, although we can rotate rectangular blocks of pixels through other angles using antialiasing procedures. For a rotation that is not a multiple of 90º, we need to estimate the percentage of area coverage for those pixels that overlap the rotated block. Sequences of raster operations can be executed to produce realtime animation for either two-dimensional or three-dimensional objects, so long as we restrict the animation to motions in the projection plane. Then no viewing or visible-surface algorithms need be invoked.

We can also animate objects along two-dimensional motion paths using **color-table transformations**. Here we redefine the object at successive positions along the motion path and set the successive blocks of pixel values to color-table entries. The pixels at the first position of the object are set to a foreground color, and the pixels at the other object positions are set to the background color. The animation
is then accomplished by changing the color-table values so that the object color at successive positions along the animation path becomes the foreground color as the preceding position is set to the background color (Figure 1).

2 Design of Animation Sequences

Constructing an animation sequence can be a complicated task, particularly when it involves a story line and multiple objects, each of which can move in a different way. A basic approach is to design such animation sequences using the following development stages:

- Storyboard layout
- Object definitions
- Key-frame specifications
- Generation of in-between frames

The storyboard is an outline of the action. It defines the motion sequence as a set of basic events that are to take place. Depending on the type of animation to be produced, the storyboard could consist of a set of rough sketches, along with a brief description of the movements, or it could just be a list of the basic ideas for the action. Originally, the set of motion sketches was attached to a large board that was used to present an overall view of the animation project. Hence, the name “storyboard.”

An object definition is given for each participant in the action. Objects can be defined in terms of basic shapes, such as polygons or spline surfaces. In addition, a description is often given of the movements that are to be performed by each character or object in the story.

A key frame is a detailed drawing of the scene at a certain time in the animation sequence. Within each key frame, each object (or character) is positioned according to the time for that frame. Some key frames are chosen at extreme positions in the action; others are spaced so that the time interval between key frames is not too great. More key frames are specified for intricate motions than for simple, slowly varying motions. Development of the key frames is generally the responsibility of the senior animators, and often a separate animator is assigned to each character in the animation.

In-betweens are the intermediate frames between the key frames. The total number of frames, and hence the total number of in-betweens, needed for an animation is determined by the display media that is to be used. Film requires 24 frames per second, and graphics terminals are refreshed at the rate of 60 or more frames per second. Typically, time intervals for the motion are set up so that there are from three to five in-betweens for each pair of key frames. Depending on the speed specified for the motion, some key frames could be duplicated. As an example, a 1-minute film sequence with no duplication requires a total of 1,440 frames. If five in-betweens are required for each pair of key frames, then 288 key frames would need to be developed.

There are several other tasks that may be required, depending on the application. These additional tasks include motion verification, editing, and the production and synchronization of a soundtrack. Many of the functions needed to produce general animations are now computer-generated. Figures 2 and 3 show examples of computer-generated frames for animation sequences.
3 Traditional Animation Techniques

Film animators use a variety of methods for depicting and emphasizing motion sequences. These include object deformations, spacing between animation frames, motion anticipation and follow-through, and action focusing.

One of the most important techniques for simulating acceleration effects, particularly for nonrigid objects, is **squash and stretch**. Figure 4 shows how this technique is used to emphasize the acceleration and deceleration of a bouncing ball. As the ball accelerates, it begins to stretch. When the ball hits the floor and stops, it is first compressed (squashed) and then stretched again as it accelerates and bounces upwards.

Another technique used by film animators is **timing**, which refers to the spacing between motion frames. A slower moving object is represented with more closely spaced frames, and a faster moving object is displayed with fewer frames over the path of the motion. This effect is illustrated in Figure 5, where the position changes between frames increase as a bouncing ball moves faster.

Object movements can also be emphasized by creating preliminary actions that indicate an **anticipation** of a coming motion. For example, a cartoon character
might lean forward and rotate its body before starting to run; or a character might perform a “windup” before throwing a ball. Similarly, follow-through actions can be used to emphasize a previous motion. After throwing a ball, a character can continue the arm swing back to its body; or a hat can fly off a character that is stopped abruptly. An action also can be emphasized with staging, which refers to any method for focusing on an important part of a scene, such as a character hiding something.

4 General Computer-Animation Functions

Many software packages have been developed either for general animation design or for performing specialized animation tasks. Typical animation functions include managing object motions, generating views of objects, producing camera motions, and the generation of in-between frames. Some animation packages, such as Wavefront for example, provide special functions for both the overall animation design and the processing of individual objects. Others are special-purpose packages for particular features of an animation, such as a system for generating in-between frames or a system for figure animation.

A set of routines is often provided in a general animation package for storing and managing the object database. Object shapes and associated parameters are stored and updated in the database. Other object functions include those for generating the object motion and those for rendering the object surfaces. Movements can be generated according to specified constraints using two-dimensional or three-dimensional transformations. Standard functions can then be applied to identify visible surfaces and apply the rendering algorithms.

Another typical function set simulates camera movements. Standard camera motions are zooming, panning, and tilting. Finally, given the specification for the key frames, the in-betweens can be generated automatically.

5 Computer-Animation Languages

We can develop routines to design and control animation sequences within a general-purpose programming language, such as C, C++, Lisp, or Fortran, but several specialized animation languages have been developed. These languages typically include a graphics editor, a key-frame generator, an in-between generator, and standard graphics routines. The graphics editor allows an animator to design and modify object shapes, using spline surfaces, constructive solid-geometry methods, or other representation schemes.

An important task in an animation specification is scene description. This includes the positioning of objects and light sources, defining the photometric parameters (light-source intensities and surface illumination properties), and setting the camera parameters (position, orientation, and lens characteristics). Another standard function is action specification, which involves the layout of motion paths for the objects and camera. We need the usual graphics routines: viewing and perspective transformations, geometric transformations to generate object movements as a function of accelerations or kinematic path specifications, visible-surface identification, and the surface-rendering operations.

Key-frame systems were originally designed as a separate set of animation routines for generating the in-betweens from the user-specified key frames. Now, these routines are often a component in a more general animation package. In the simplest case, each object in a scene is defined as a set of rigid bodies connected at the joints and with a limited number of degrees of freedom. As an example, the
single-armed robot in Figure 6 has 6 degrees of freedom, which are referred to as arm sweep, shoulder swivel, elbow extension, pitch, yaw, and roll. We can extend the number of degrees of freedom for this robot arm to 9 by allowing three-dimensional translations for the base (Figure 7). If we also allow base rotations, the robot arm can have a total of 12 degrees of freedom. The human body, in comparison, has more than 200 degrees of freedom.

**Parameterized systems** allow object motion characteristics to be specified as part of the object definitions. The adjustable parameters control such object characteristics as degrees of freedom, motion limitations, and allowable shape changes. **Scripting systems** allow object specifications and animation sequences to be defined with a user-input *script*. From the script, a library of various objects and motions can be constructed.

## 6 Key-Frame Systems

A set of in-betweens can be generated from the specification of two (or more) key frames using a key-frame system. Motion paths can be given with a *kinematic description* as a set of spline curves, or the motions can be *physically based* by specifying the forces acting on the objects to be animated.

For complex scenes, we can separate the frames into individual components or objects called *cels* (celluloid transparencies). This term developed from cartoon-animation techniques where the background and each character in a scene were placed on a separate transparency. Then, with the transparencies stacked in the order from background to foreground, they were photographed to obtain the completed frame. The specified animation paths are then used to obtain the next cel for each character, where the positions are interpolated from the key-frame times.

With complex object transformations, the shapes of objects may change over time. Examples are clothes, facial features, magnified detail, evolving shapes, and exploding or disintegrating objects. For surfaces described with polygon meshes, these changes can result in significant changes in polygon shape such that the number of edges in a polygon could be different from one frame to the next. These changes are incorporated into the development of the in-between frames by adding or subtracting polygon edges according to the requirements of the defining key frames.

**Morphing**

Transformation of object shapes from one form to another is termed **morphing**, which is a shortened form of “metamorphosing.” An animator can model morphing by transitioning polygon shapes through the in-betweens from one key frame to the next.
Given two key frames, each with a different number of line segments specifying an object transformation, we can first adjust the object specification in one of the frames so that the number of polygon edges (or the number of polygon vertices) is the same for the two frames. This preprocessing step is illustrated in Figure 8. A straight-line segment in key frame \( k \) is transformed into two line segments in key frame \( k + 1 \). Because key frame \( k + 1 \) has an extra vertex, we add a vertex between vertices 1 and 2 in key frame \( k \) to balance the number of vertices (and edges) in the two key frames. Using linear interpolation to generate the in-betweens, we transition the added vertex in key frame \( k \) into vertex 3′ along the straight-line path shown in Figure 9. An example of a triangle linearly expanding into a quadrilateral is given in Figure 10.

We can state general preprocessing rules for equalizing key frames in terms of either the number of edges or the number of vertices to be added to a key frame. We first consider equalizing the edge count, where parameters \( L_k \) and \( L_{k+1} \) denote the number of line segments in two consecutive frames. The maximum and minimum number of lines to be equalized can be determined as

\[
L_{\text{max}} = \max(L_k, L_{k+1}), \quad L_{\text{min}} = \min(L_k, L_{k+1})
\]

(1)

Next we compute the following two quantities:

\[
N_e = L_{\text{max}} \mod L_{\text{min}}
\]

\[
N_s = \text{int}\left(\frac{L_{\text{max}}}{L_{\text{min}}}\right)
\]

(2)

**Figure 8**
An edge with vertex positions 1 and 2 in key frame \( k \) evolves into two connected edges in key frame \( k + 1 \).

**Figure 9**
Linear interpolation for transforming a line segment in key frame \( k \) into two connected line segments in key frame \( k + 1 \).

**Figure 10**
Linear interpolation for transforming a triangle into a quadrilateral.
The preprocessing steps for edge equalization are then accomplished with the following two procedures:

1. Divide $N_e$ edges of $\text{keyframe}_{\text{min}}$ into $N_e + 1$ sections.
2. Divide the remaining lines of $\text{keyframe}_{\text{min}}$ into $N_e$ sections.

As an example, if $L_k = 15$ and $L_{k+1} = 11$, we would divide four lines of $\text{keyframe}_{k+1}$ into two sections each. The remaining lines of $\text{keyframe}_{k+1}$ are left intact.

If we equalize the vertex count, we can use parameters $V_k$ and $V_{k+1}$ to denote the number of vertices in the two consecutive key frames. In this case, we determine the maximum and minimum number of vertices as

$$V_{\text{max}} = \max(V_k, V_{k+1}), \quad V_{\text{min}} = \min(V_k, V_{k+1})$$  \hspace{1cm} (3)

Then we compute the following two values:

$$N_{ls} = (V_{\text{max}} - 1) \mod (V_{\text{min}} - 1)$$

$$N_p = \text{int} \left( \frac{V_{\text{max}} - 1}{V_{\text{min}} - 1} \right)$$  \hspace{1cm} (4)

These two quantities are then used to perform vertex equalization with the following procedures:

1. Add $N_p$ points to $N_{ls}$ line sections of $\text{keyframe}_{\text{min}}$.
2. Add $N_p - 1$ points to the remaining edges of $\text{keyframe}_{\text{min}}$.

For the triangle-to-quadrilateral example, $V_k = 3$ and $V_{k+1} = 4$. Both $N_{ls}$ and $N_p$ are 1, so we would add one point to one edge of $\text{keyframe}_k$. No points would be added to the remaining lines of $\text{keyframe}_k$.

### Simulating Accelerations

Curve-fitting techniques are often used to specify the animation paths between key frames. Given the vertex positions at the key frames, we can fit the positions with linear or nonlinear paths. Figure 11 illustrates a nonlinear fit of keyframe positions. To simulate accelerations, we can adjust the time spacing for the in-betweens.

If the motion is to occur at constant speed (zero acceleration), we use equal-interval time spacing for the in-betweens. For instance, with $n$ in-betweens and
key-frame times of \( t_1 \) and \( t_2 \) (Figure 12), the time interval between the key frames is divided into \( n + 1 \) equal subintervals, yielding an in-between spacing of

\[
\Delta t = \frac{t_2 - t_1}{n + 1}
\]  

(5)

The time for the \( j \)th in-between is

\[
t_{B_j} = t_1 + j \Delta t, \quad j = 1, 2, \ldots, n
\]  

(6)

and this time value is used to calculate coordinate positions, color, and other physical parameters for that frame of the motion.

Speed changes (nonzero accelerations) are usually necessary at some point in an animation film or cartoon, particularly at the beginning and end of a motion sequence. The startup and slowdown portions of an animation path are often modeled with spline or trigonometric functions, but parabolic and cubic time functions have been applied to acceleration modeling. Animation packages commonly furnish trigonometric functions for simulating accelerations.

To model increasing speed (positive acceleration), we want the time spacing between frames to increase so that greater changes in position occur as the object moves faster. We can obtain an increasing size for the time interval with the function

\[
1 - \cos \theta, \quad 0 < \theta < \pi/2
\]

For \( n \) in-betweens, the time for the \( j \)th in-between would then be calculated as

\[
t_{B_j} = t_1 + \Delta t \left( 1 - \cos \frac{j \pi}{2(n + 1)} \right), \quad j = 1, 2, \ldots, n
\]  

(7)

where \( \Delta t \) is the time difference between the two key frames. Figure 13 gives a plot of the trigonometric acceleration function and the in-between spacing for \( n = 5 \).

We can model decreasing speed (deceleration) using the function \( \sin \theta \), with \( 0 < \theta < \pi/2 \). The time position of an in-between is then determined as

\[
t_{B_j} = t_1 + \Delta t \sin \frac{j \pi}{2(n + 1)}, \quad j = 1, 2, \ldots, n
\]  

(8)
A plot of this function and the decreasing size of the time intervals is shown in Figure 14 for five in-betweens.

Often, motions contain both speedups and slowdowns. We can model a combination of increasing–decreasing speed by first increasing the in-between time spacing and then decreasing this spacing. A function to accomplish these time changes is

\[ \frac{1}{2}(1 - \cos \theta), \quad 0 < \theta < \pi/2 \]

The time for the \( j \)th in-between is now calculated as

\[ t_{B_j} = t_1 + \Delta t \left\{ \frac{1 - \cos [j\pi/(n + 1)]}{2} \right\}, \quad j = 1, 2, \ldots, n \]  

(9)

with \( \Delta t \) denoting the time difference between the two key frames. Time intervals for a moving object first increase and then decrease, as shown in Figure 15.

---

**Figure 14**
A trigonometric deceleration function and the corresponding in-between spacing for \( n = 5 \) and \( \theta = j\pi/12 \) in Equation 8, producing decreased coordinate changes as the object moves through each time interval.

**Figure 15**
The trigonometric accelerate–decelerate function \((1 - \cos \theta)/2\) and the corresponding in-between spacing for \( n = 5 \) in Equation 9.
Processing the in-betweens is simplified by initially modeling “skeleton” (wire-frame) objects so that motion sequences can be interactively adjusted. After the animation sequence is completely defined, objects can be fully rendered.

7 Motion Specifications

General methods for describing an animation sequence range from an explicit specification of the motion paths to a description of the interactions that produce the motions. Thus, we could define how an animation is to take place by giving the transformation parameters, the motion path parameters, the forces that are to act on objects, or the details of how objects interact to produce motion.

Direct Motion Specification

The most straightforward method for defining an animation is direct motion specification of the geometric-transformation parameters. Here, we explicitly set the values for the rotation angles and translation vectors. Then the geometric transformation matrices are applied to transform coordinate positions. Alternatively, we could use an approximating equation involving these parameters to specify certain kinds of motions. We can approximate the path of a bouncing ball, for instance, with a damped, rectified, sine curve (Figure 16):

\[ y(x) = A|\sin(\omega x + \theta_0)|e^{-kx} \]  

where \( A \) is the initial amplitude (height of the ball above the ground), \( \omega \) is the angular frequency, \( \theta_0 \) is the phase angle, and \( k \) is the damping constant. This method for motion specification is particularly useful for simple user-programmed animation sequences.

Goal-Directed Systems

At the opposite extreme, we can specify the motions that are to take place in general terms that abstractly describe the actions in terms of the final results. In other words, an animation is specified in terms of the final state of the movements. These systems are referred to as goal-directed, since values for the motion parameters are determined from the goals of the animation. For example, we could specify that

![Approximating the motion of a bouncing ball with a damped sine function (Eq. 10).](image)
we want an object to “walk” or to “run” to a particular destination; or we could state that we want an object to “pick up” some other specified object. The input directives are then interpreted in terms of component motions that will accomplish the described task. Human motions, for instance, can be defined as a hierarchical structure of submotions for the torso, limbs, and so forth. Thus, when a goal, such as “walk to the door” is given, the movements required of the torso and limbs to accomplish this action are calculated.

**Kinematics and Dynamics**

We can also construct animation sequences using *kinematic* or *dynamic* descriptions. With a kinematic description, we specify the animation by giving motion parameters (position, velocity, and acceleration) without reference to causes or goals of the motion. For constant velocity (zero acceleration), we designate the motions of rigid bodies in a scene by giving an initial position and velocity vector for each object. For example, if a velocity is specified as $(3, 0, -4)$ km per sec, then this vector gives the direction for the straight-line motion path and the speed (magnitude of velocity) is calculated as 5 km per sec. If we also specify accelerations (rate of change of velocity), we can generate speedups, slowdowns, and curved motion paths. Kinematic specification of a motion can also be given by simply describing the motion path. This is often accomplished using spline curves.

An alternate approach is to use *inverse kinematics*. Here, we specify the initial and final positions of objects at specified times and the motion parameters are computed by the system. For example, assuming zero acceleration, we can determine the constant velocity that will accomplish the movement of an object from the initial position to the final position. This method is often used with complex objects by giving the positions and orientations of an end node of an object, such as a hand or a foot. The system then determines the motion parameters of other nodes to accomplish the desired motion.

Dynamic descriptions, on the other hand, require the specification of the forces that produce the velocities and accelerations. The description of object behavior in terms of the influence of forces is generally referred to as *physically based modeling*. Examples of forces affecting object motion include electromagnetic, gravitational, frictional, and other mechanical forces.

Object motions are obtained from the force equations describing physical laws, such as Newton’s laws of motion for gravitational and frictional processes, Euler or Navier-Stokes equations describing fluid flow, and Maxwell’s equations for electromagnetic forces. For example, the general form of Newton’s second law for a particle of mass $m$ is

$$F = \frac{d}{dt} (mv)$$  \hspace{1cm} (11)

where $F$ is the force vector and $v$ is the velocity vector. If mass is constant, we solve the equation $F = ma$, with $a$ representing the acceleration vector. Otherwise, mass is a function of time, as in relativistic motions or the motions of space vehicles that consume measurable amounts of fuel per unit time. We can also use *inverse dynamics* to obtain the forces, given the initial and final positions of objects and the type of motion required.

Applications of physically based modeling include complex rigid-body systems and such nonrigid systems as cloth and plastic materials. Typically, numerical methods are used to obtain the motion parameters incrementally from the dynamical equations using initial conditions or boundary values.
Character Animation

Animation of simple objects is relatively straightforward. When we consider the animation of more complex figures such as humans or animals, however, it becomes much more difficult to create realistic animation. Consider the animation of walking or running human (or humanoid) characters. Based upon observations in their own lives of walking or running people, viewers will expect to see animated characters move in particular ways. If an animated character’s movement doesn’t match this expectation, the believability of the character may suffer. Thus, much of the work involved in character animation is focused on creating believable movements.

Articulated Figure Animation

A basic technique for animating people, animals, insects, and other critters is to model them as articulated figures, which are hierarchical structures composed of a set of rigid links that are connected at rotary joints (Figure 17). In less formal terms, this just means that we model animate objects as moving stick figures, or simplified skeletons, that can later be wrapped with surfaces representing skin, hair, fur, feathers, clothes, or other outer coverings.

The connecting points, or hinges, for an articulated figure are placed at the shoulders, hips, knees, and other skeletal joints, which travel along specified motion paths as the body moves. For example, when a motion is specified for an object, the shoulder automatically moves in a certain way and, as the shoulder moves, the arms move. Different types of movement, such as walking, running, or jumping, are defined and associated with particular motions for the joints and connecting links.

A series of walking leg motions, for instance, might be defined as in Figure 18. The hip joint is translated forward along a horizontal line, while the connecting links perform a series of movements about the hip, knee, and angle joints. Starting with a straight leg [Figure 18(a)], the first motion is a knee bend as the hip moves forward [Figure 18(b)]. Then the leg swings forward, returns to the vertical position, and swings back, as shown in Figures 18(c), (d), and (e). The final motions are a wide swing back and a return to the straight position.

Figure 17
A simple articulated figure with nine joints and twelve connecting links, not counting the oval head.

Figure 18
Possible motions for a set of connected links representing a walking leg.
vertical position, as in Figures 18(f) and (g). This motion cycle is repeated for the duration of the animation as the figure moves over a specified distance or time interval.

As a figure moves, other movements are incorporated into the various joints. A sinusoidal motion, often with varying amplitude, can be applied to the hips so that they move about on the torso. Similarly, a rolling or rocking motion can be imparted to the shoulders, and the head can bob up and down.

Both kinematic-motion descriptions and inverse kinematics are used in figure animations. Specifying the joint motions is generally an easier task, but inverse kinematics can be useful for producing simple motion over arbitrary terrain. For a complicated figure, inverse kinematics may not produce a unique animation sequence: Many different rotational motions may be possible for a given set of initial and final conditions. In such cases, a unique solution may be possible by adding more constraints, such as conservation of momentum, to the system.

**Motion Capture**

An alternative to determining the motion of a character computationally is to digitally record the movement of a live actor and to base the movement of an animated character on that information. This technique, known as *motion capture* or *mo-cap*, can be used when the movement of the character is predetermined (as in a scripted scene). The animated character will perform the same series of movements as the live actor.

The classic motion capture technique involves placing a set of markers at strategic positions on the actor’s body, such as the arms, legs, hands, feet, and joints. It is possible to place the markers directly on the actor, but more commonly they are affixed to a special skintight body suit worn by the actor. The actor is then filmed performing the scene. Image processing techniques are then used to identify the positions of the markers in each frame of the film, and their positions are translated to coordinates. These coordinates are used to determine the positioning of the body of the animated character. The movement of each marker from frame to frame in the film is tracked and used to control the corresponding movement of the animated character.

To accurately determine the positions of the markers, the scene must be filmed by multiple cameras placed at fixed positions. The digitized marker data from each recording can then be used to triangulate the position of each marker in three dimensions. Typical motion capture systems will use up to two dozen cameras, but systems with several hundred cameras exist.

Optical motion capture systems rely on the reflection of light from a marker into the camera. These can be relatively simple passive systems using photo-reflective markers that reflect illumination from special lights placed near the cameras, or more advanced active systems in which the markers are powered and emit light. Active systems can be constructed so that the markers illuminate in a pattern or sequence, which allows each marker to be uniquely identified in each frame of the recording, simplifying the tracking process.

Non-optical systems rely on the direct transmission of position information from the markers to a recording device. Some non-optical systems use inertial sensors that provide gyroscope-based position and orientation information. Others use magnetic sensors that measure changes in magnetic flux. A series of transmitters placed around the stage generate magnetic fields that induce current in the magnetic sensors; that information is then transmitted to receivers.

Some motion capture systems record more than just the gross movements of the parts of the actor’s body. It is possible to record even the actor’s
Facial movements. Often called *performance capture* systems, these typically use a camera trained on the actor’s face and small light-emitting diode (LED) lights that illuminate the face. Small photoreflective markers attached to the face reflect the light from the LEDs and allow the camera to capture the small movements of the muscles of the face, which can then be used to create realistic facial animation on a computer-generated character.

**9 Periodic Motions**

When we construct an animation with repeated motion patterns, such as a rotating object, we need to be sure that the motion is sampled frequently enough to represent the movements correctly. In other words, the motion must be synchronized with the frame-generation rate so that we display enough frames per cycle to show the true motion. Otherwise, the animation may be displayed incorrectly.

A typical example of an undersampled periodic-motion display is the wagon wheel in a Western movie that appears to be turning in the wrong direction. Figure 19 illustrates one complete cycle in the rotation of a wagon wheel with one red spoke that makes 18 clockwise revolutions per second. If this motion is recorded on film at the standard motion-picture projection rate of 24 frames per second, then the first five frames depicting this motion would be as shown in Figure 20. Because the wheel completes \( \frac{2}{3} \) of a turn every \( \frac{1}{24} \) of a second, only one animation frame is generated per cycle, and the wheel thus appears to be rotating in the opposite (counterclockwise) direction.

In a computer-generated animation, we can control the sampling rate in a periodic motion by adjusting the motion parameters. For example, we can set the

![Figure 19](image1.png)

*Figure 19* Five positions for a red spoke during one cycle of a wheel motion that is turning at the rate of 18 revolutions per second.

![Figure 20](image2.png)

*Figure 20* The first five film frames of the rotating wheel in Figure 19 produced at the rate of 24 frames per second.
angular increment for the motion of a rotating object so that multiple frames are generated in each revolution. Thus, a $3^\circ$ increment for a rotation angle produces 120 motion steps during one revolution, and a $4^\circ$ increment generates 90 steps. For faster motion, larger rotational steps could be used, so long as the number of samples per cycle is not too small and the motion is clearly displayed. When complex objects are to be animated, we also must take into account the effect that the frame construction time might have on the refresh rate, as discussed in Section 1. The motion of a complex object can be much slower than we want it to be if it takes too long to construct each frame of the animation.

Another factor that we need to consider in the display of a repeated motion is the effect of round-off in the calculations for the motion parameters. We can reset parameter values periodically to prevent the accumulated error from producing erratic motions. For a continuous rotation, we could reset parameter values once every cycle ($360^\circ$).

### 10 OpenGL Animation Procedures

Raster operations and color-index assignment functions are available in the core library, and routines for changing color-table values are provided in GLUT. Other raster-animation operations are available only as GLUT routines because they depend on the window system in use. In addition, computer-animation features such as double buffering may not be included in some hardware systems.

Double-buffering operations, if available, are activated using the following GLUT command:

```c
glutInitDisplayMode (GLUT_DOUBLE);
```

This provides two buffers, called the front buffer and the back buffer, that we can use alternately to refresh the screen display. While one buffer is acting as the refresh buffer for the current display window, the next frame of an animation can be constructed in the other buffer. We specify when the roles of the two buffers are to be interchanged using

```c
glutSwapBuffers ( );
```

To determine whether double-buffer operations are available on a system, we can issue the following query:

```c
glGetBooleanv (GL_DOUBLEBUFFER, status);
```

A value of `GL_TRUE` is returned to array parameter `status` if both front and back buffers are available on a system. Otherwise, the returned value is `GL_FALSE`.

For a continuous animation, we can also use

```c
glutIdleFunc (animationFcn);
```

where parameter `animationFcn` can be assigned the name of a procedure that is to perform the operations for incrementing the animation parameters. This procedure is continuously executed whenever there are no display-window events that must be processed. To disable the `glutIdleFunc`, we set its argument to the value `NULL` or the value `0`.

An example animation program is given in the following code, which continuously rotates a regular hexagon in the $xy$ plane about the $z$ axis. The origin of
three-dimensional screen coordinates is placed at the center of the display window so that the z axis passes through this center position. In procedure init, we use a display list to set up the description of the regular hexagon, whose center position is originally at the screen-coordinate position (150, 150) with a radius (distance from the polygon center to any vertex) of 100 pixels. In the display function, displayHex, we specify an initial 0° rotation about the z axis and invoke the glutSwapBuffers routine. To activate the rotation, we use procedure mouseFcn, which continually increments the rotation angle by 3° once we press the middle mouse button. The calculation of the incremented rotation angle is performed in procedure rotateHex, which is called by the glutIdleFunc routine in procedure mouseFcn. We stop the rotation by pressing the right mouse button, which causes the glutIdleFunc to be invoked with a NULL argument.

```c
#include <GL/glut.h>
#include <math.h>
#include <stdlib.h>

const double TWO_PI = 6.2831853;

GLsizei winWidth = 500, winHeight = 500; // Initial display window size.
GLuint regHex; // Define name for display list.
static GLfloat rotTheta = 0.0;

class scrPt {
    public:
        GLint x, y;
};

static void init (void)
{
    scrPt hexVertex;
    GLdouble hexTheta;
    GLint k;

    glClearColor (1.0, 1.0, 1.0, 0.0);

    /* Set up a display list for a red regular hexagon. *
    * Vertices for the hexagon are six equally spaced  *
    * points around the circumference of a circle.     *
    */
    regHex = glGenLists (1);
    glNewList (regHex, GL_COMPILE);
    glColor3f (1.0, 0.0, 0.0);
    glBegin (GL_POLYGON);
    for (k = 0; k < 6; k++) {
        hexTheta = TWO_PI * k / 6;
        hexVertex.x = 150 + 100 * cos (hexTheta);
        hexVertex.y = 150 + 100 * sin (hexTheta);
        glVertex2i (hexVertex.x, hexVertex.y);
    }
    glEnd ( );
    glEndList ( );
}
```

void displayHex (void)
{
    glClear (GL_COLOR_BUFFER_BIT);
    glPushMatrix ( );
    glRotatef (rotTheta, 0.0, 0.0, 1.0);
    glCallList (regHex);
    glPopMatrix ( );
    glutSwapBuffers ( );
    glFlush ( );
}

void rotateHex (void)
{
    rotTheta += 3.0;
    if (rotTheta > 360.0)
        rotTheta -= 360.0;
    glutPostRedisplay ( );
}

void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    glViewport (0, 0, (GLsizei) newWidth, (GLsizei) newHeight);
    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ( );
    gluOrtho2D (-320.0, 320.0, -320.0, 320.0);
    glMatrixMode (GL_MODELVIEW);
    glLoadIdentity ( );
    glClear (GL_COLOR_BUFFER_BIT);
}

void mouseFcn (GLint button, GLint action, GLint x, GLint y)
{
    switch (button) {
    case GLUT_MIDDLE_BUTTON: // Start the rotation.
        if (action == GLUT_DOWN)
            glutIdleFunc (rotateHex);
        break;
    case GLUT_RIGHT_BUTTON:  // Stop the rotation.
        if (action == GLUT_DOWN)
            glutIdleFunc (NULL);
        break;
    default:
        break;
    }
void main (int argc, char** argv)
{
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_DOUBLE | GLUT_RGB);
    glutInitWindowPosition (150, 150);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Animation Example");

    init ( );
    glutDisplayFunc (displayHex);
    glutReshapeFunc (winReshapeFcn);
    glutMouseFunc (mouseFcn);

    glutMainLoop ( );
}

11 Summary

An animation sequence can be constructed frame by frame, or it can be generated in real time. When separate frames of an animation are constructed and stored, the frames can later be transferred to film or displayed in rapid succession on a video monitor. Animations involving complex scenes and motions are commonly produced one frame at a time, while simpler motion sequences are displayed in real time.

On a raster system, double-buffering methods can be used to facilitate motion displays. One buffer is used to refresh the screen, while a second buffer is being loaded with the screen values for the next frame of the motion. Then the roles of the two buffers are interchanged, usually at the end of a refresh cycle.

Another raster method for displaying an animation is to perform motion sequences using block transfers of pixel values. Translations are accomplished by a simple move of a rectangular block of pixel colors from one frame-buffer position to another. And rotations in $90^\circ$ increments can be performed with combinations of translations and row-column interchanges within the pixel array.

Color-table methods can be used for simple raster animations by storing an image of an object at multiple locations in the frame buffer, using different color-table values. One image is stored in the foreground color, and the copies of the image at the other locations are assigned a background color. By rapidly interchanging the foreground and background color values stored in the color table, we can display the object at various screen positions.

Several developmental stages can be used to produce an animation, starting with the storyboard, object definitions, and specification of key frames. The storyboard is an outline of the action, and the key frames define the details of the object motions for selected positions in the animation. Once the key frames have been established, in-between frames are generated to construct a smooth motion from one key frame to the next. A computer animation can involve motion specifications for the “camera,” as well as motion paths for the objects and characters involved in the animation.

Various techniques have been developed for simulating and emphasizing motion effects. Squash and stretch effects are standard methods for stressing
accelerations, and the timing between motion frames can be varied to produce speed variations. Other methods include a preliminary windup motion, a follow-through at the end of an action, and staging methods that focus on an important action in a scene. Trigonometric functions are typically used to generate the time spacing for in-between frames when the motions involve accelerations.

Animations can be generated with special-purpose software or with a general-purpose graphics package. Systems that are available for automated computer animation include key-frame systems, parameterized systems, and scripting systems.

Many animations include morphing effects, in which one object shape is transformed into another. These effects are accomplished by using the in-between frames to transition the defining points and lines in one object into the points and lines of the other object.

Motions in an animation can be described with direct motion specifications or they can be goal-directed. Thus, an animation can be defined in terms of translation and rotation parameters, or motions can be described with equations or with kinematic or dynamic parameters. Kinematic motion descriptions specify positions, velocities, and accelerations; dynamic motion descriptions are given in terms of the forces acting on the objects in a scene.

Articulated figures are often used to model the motions of people and animals. Rigid links, connected at rotary joints, are defined in a hierarchical structure. When a motion is imparted to an object, each subpart is programmed to move in a particular way in response to the overall motion.

Motion capture techniques provide an alternative to computed character motion. They can be used to produce more realistic movement for articulated characters.

The sampling rate for periodic motions should produce enough frames per cycle to display the animation correctly. Otherwise, erratic or misleading motions may result.

In addition to the raster ops and color-table methods, a few functions are available in the OpenGL Utility Toolkit (GLUT) for developing animation programs. These provide routines for double-buffering operations and for incrementing motion parameters during idle-processing intervals. In Table 1, we list the GLUT functions for producing animations with OpenGL programs.

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glutInitDisplayMode (GLUT_DOUBLE)</td>
<td>Activates double-buffering operations.</td>
</tr>
<tr>
<td>glutSwapBuffers</td>
<td>Interchanges front and back refresh buffers.</td>
</tr>
<tr>
<td>glGetBooleanv (GL_DOUBLEBUFFER, status)</td>
<td>Queries a system to determine whether double buffering is available.</td>
</tr>
<tr>
<td>glutIdleFunc</td>
<td>Specifies a function for incrementing animation parameters.</td>
</tr>
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REFERENCES


EXERCISES

1 Design a storyboard layout and accompanying key frames for an animation of a simple stick figure, as in Figure 17.
2 Write a program to generate the in-betweens for the key frames specified in Exercise 1 using linear interpolation.
3 Expand the animation sequence in Exercise 1 to include two or more moving objects.
4 Write a program to generate the in-betweens for the key frames in Exercise 3 using linear interpolation.
5 Write a morphing program to transform any given polygon into another specified polygon, using five in-betweens.
6 Write a morphing program to transform a sphere into a specified polyhedron, using five in-betweens.
7 Set up an animation specification involving accelerations and implementing Eq. 7.
8 Set up an animation specification involving both accelerations and decelerations, implementing the in-between spacing calculations given in Equations 7 and 8.
9 Set up an animation specification implementing the acceleration–deceleration calculations of Equation 9.
10 Write a program to simulate the linear, two-dimensional motion of a filled circle inside a given rectangular area. The circle is to be given an initial position and velocity, and the circle is to rebound from the walls with the angle of reflection equal to the angle of incidence.
11 Convert the program of the previous exercise into a two-player ball and paddle game by replacing two opposite sides of the rectangle with short line segments that can be moved back and forth along each of the rectangle edges. Interactive movement of each line segment simulates a paddle that can be positioned to prevent the bouncing ball from escaping that side of the rectangle. Each time the circle escapes from the interior of the rectangle the score of the player assigned to the opposite side is increased. Initial input parameters include circle position, direction, and speed. The game scores can be displayed in one corner of the display window.
12 Modify the ball and paddle game in the previous exercise to vary the speed of the bouncing ball. After each successful block by a player, the speed of the ball is incremented by a small amount.
13 Modify the game in the previous exercise to include two balls, each initialized to the same speed but different positions and opposite directions.
14 Modify the two-dimensional bouncing ball inside a rectangle to a three-dimensional motion of a sphere bouncing around inside a parallelepiped. Interactive viewing parameters can be set to view the motion from different directions.
15 Write a program to implement the simulation of a bouncing ball using Eq. 10.
16 Expand the program in the previous exercise to include squash and stretch effects.
17 Write a program to implement the motion of a bouncing ball using dynamics. The motion of the ball is to be governed by a downward gravitational force and a ground-plane friction force. Initially, the ball is to be projected into space with a given velocity vector.
18 Write a program to implement dynamic motion specifications. Specify a scene with two or more objects, initial motion parameters, and specified forces. Then generate the animation from the solution of the force equations. (For example, the objects could be the earth, moon, and sun with attractive gravitational forces that are proportional to mass and inversely proportional to distance squared.)
19 Modify the rotating hexagon program in Section 10 to allow a user to interactively choose a three-dimensional object from a list of menu options to be rotated about the y axis. Use a perspective projection to display the object.
20 Modify the program in the previous exercise so that the rotation about the y axis is an elliptical path in the xz plane.
21 Modify the program in the previous exercise to allow interactive variation of the rotation speed.
IN MORE DEPTH

1. The goal of this chapter’s exercises is to increase the sophistication of the animation of your developing application. You may do this by modifying or adding to the existing animation that you have developed, or by using the techniques that you have learned so far to design a different animation of some portion of your application. In either case, draw a storyboard of your planned animation and outline the methods that you will use to implement it. Identify key frames that represent critical points in the animation. Consider methods by which you can generate the in-between frames that will move the objects in the scene realistically from one key frame to the next. Sketch a timeline that will help you determine the rate at which each object or group of objects should move in between each key frame. Try to have some objects exhibit nonzero accelerations in your animation. If possible, include an instance of linearly interpolated morphing from one polygon or set of polygons to another. If your application contains objects that behave like a physical dynamical system, try to incorporate dynamics in the form of physical models to produce the animation. Finally, make an attempt to include periodic motion in the animation if appropriate.

2. Given the storyboard and specification that you designed in the previous exercise, implement the animation in an OpenGL program with just a single buffer as previously done, and then with a double buffer. Note any differences in the quality of the animation between the two cases.
Graphics scenes can contain many different kinds of objects and material surfaces: trees, flowers, clouds, rocks, water, bricks, wood paneling, rubber, paper, marble, steel, glass, plastic, and cloth, just to mention a few. So it may not be surprising that there is no single method that we can use to describe objects that will include all the characteristics of these different materials.

Polygon and quadric surfaces provide precise descriptions for simple Euclidean objects such as polyhedrons and ellipsoids. They are examples of boundary representations (B-reps), which describe a three-dimensional object as a set of surfaces that separate the object interior from the environment. In this chapter, we consider the features of these types of representation schemes and how they are used in computer-graphics applications.
1 Polyhedra

The most commonly used boundary representation for a three-dimensional graphics object is a set of surface polygons that enclose the object interior. Many graphics systems store all object descriptions as sets of surface polygons. This simplifies and speeds up the surface rendering and display of objects because all surfaces are described with linear equations. For this reason, polygon descriptions are often referred to as standard graphics objects. In some cases, a polygonal representation is the only one available, but many packages also allow object surfaces to be described with other schemes, such as spline surfaces, which are usually converted to polygonal representations for processing through the viewing pipeline.

To describe an object as a set of polygon facets, we give the list of vertex coordinates for each polygon section over the object surface. The vertex coordinates and edge information for the surface sections are then stored in tables along with other information, such as the surface normal vector for each polygon. Some graphics packages provide routines for generating a polygon-surface mesh as a set of triangles or quadrilaterals. This allows us to describe a large section of an object’s bounding surface, or even the entire surface, with a single command. And some packages also provide routines for displaying common shapes, such as a cube, sphere, or cylinder, represented with polygon surfaces. Sophisticated graphics systems use fast hardware-implemented polygon renderers that have the capability for displaying a million or more shaded polygons (usually triangles) per second, including the application of surface texture and special lighting effects.

2 OpenGL Polyhedron Functions

We have two methods for specifying polygon surfaces in an OpenGL program. Using the polygon primitives we can generate a variety of polyhedron shapes and surface meshes. In addition, we can use GLUT functions to display the five regular polyhedra.

OpenGL Polygon Fill-Area Functions

A set of polygon patches for a section of an object surface, or a complete description for a polyhedron, can be given using the OpenGL primitive constants GL_POLYGON, GL_TRIANGLES, GL_TRIANGLE_STRIP, GL_TRIANGLE_FAN, GL_QUADS, and GL_QUAD_STRIP. For example, we could tessellate the lateral (axial) surface of a cylinder using a quadrilateral strip. Similarly, all faces of a parallelogram can be described with a set of rectangles, and all faces of a triangular pyramid could be specified using a set of connected triangular surfaces.

GLUT Regular Polyhedron Functions

Some standard shapes—the five regular polyhedra—are predefined by routines in the GLUT library. These polyhedra, also called the Platonic solids, are distinguished by the fact that all the faces of any regular polyhedron are identical regular polygons. Thus, all edges in a regular polyhedron are equal, all edge angles are equal, and all angles between faces are equal. Polyhedra are named according to the number of faces in each of the solids, and the five regular polyhedra are the regular tetrahedron (or triangular pyramid, with 4 faces), the regular hexahedron (or cube, with 6 faces), the regular octahedron (8 faces), the regular dodecahedron (12 faces), and the regular icosahedron (20 faces).
Ten functions are provided in GLUT for generating these solids: five of the functions produce wire-frame objects, and five display the polyhedra facets as shaded fill areas. The displayed surface characteristics for the fill areas are determined by the material properties and the lighting conditions that we set for a scene. Each regular polyhedron is described in modeling coordinates, so that each is centered at the world-coordinate origin.

We obtain the four-sided, regular triangular pyramid using either of these two functions:

\[ \text{glutWireTetrahedron ( )}; \]

or

\[ \text{glutSolidTetrahedron ( )}; \]

This polyhedron is generated with its center at the world-coordinate origin and with a radius (distance from the center of the tetrahedron to any vertex) equal to \( \sqrt{3} \).

The six-sided regular hexahedron (cube) is displayed with

\[ \text{glutWireCube (edgeLength)}; \]

or

\[ \text{glutSolidCube (edgeLength)}; \]

Parameter \( \text{edgeLength} \) can be assigned any positive, double-precision floating-point value, and the cube is centered on the coordinate origin.

To display the eight-sided regular octahedron, we invoke either of the following commands:

\[ \text{glutWireOctahedron ( )}; \]

or

\[ \text{glutSolidOctahedron ( )}; \]

This polyhedron has equilateral triangular faces, and the radius (distance from the center of the octahedron at the coordinate origin to any vertex) is 1.0.

The twelve-sided regular dodecahedron, centered at the world-coordinate origin, is generated with

\[ \text{glutWireDodecahedron ( )}; \]

or

\[ \text{glutSolidDodecahedron ( )}; \]

Each face of this polyhedron is a pentagon.

The following two functions generate the twenty-sided regular icosahedron:

\[ \text{glutWireIcosahedron ( )}; \]

or

\[ \text{glutSolidIcosahedron ( )}; \]

Default radius (distance from the polyhedron center at the coordinate origin to any vertex) for the icosahedron is 1.0, and each face is an equilateral triangle.
**Example GLUT Polyhedron Program**

Using the GLUT functions for the Platonic solids, the following program generates a transformed, wire-frame perspective display of these polyhedrons. All five solids are positioned within one display window (shown in Figure 1).

```c
#include <GL/glut.h>
GLsizei winWidth = 500, winHeight = 500; // Initial display-window size.

void init (void)
{
    glClearColor (1.0, 1.0, 1.0, 0.0); // White display window.
}

void displayWirePolyhedra (void)
{
    glClear (GL_COLOR_BUFFER_BIT); // Clear display window.
    glColor3f (0.0, 0.0, 1.0); // Set line color to blue.
    /* Set viewing transformation. */
    gluLookAt (5.0, 5.0, 5.0, 0.0, 0.0, 0.0, 0.0, 1.0, 0.0);

    /* Scale cube and display as wire-frame parallelepiped. */
    glScalef (1.5, 2.0, 1.0);
    glutWireCube (1.0);
}
```

**FIGURE 1**
A perspective view of the five GLUT polyhedra, scaled and positioned within a display window by procedure `displayWirePolyhedra`. 
/* Scale, translate, and display wire-frame dodecahedron. */
glScalef (0.8, 0.5, 0.8);
glTranslatef (-6.0, -5.0, 0.0);
glutWireDodecahedron ( );

/* Translate and display wire-frame tetrahedron. */
glTranslatef (8.6, 8.6, 2.0);
glutWireTetrahedron ( );

/* Translate and display wire-frame octahedron. */
glTranslatef (-3.0, -1.0, 0.0);
glutWireOctahedron ( );

/* Scale, translate, and display wire-frame icosahedron. */
glScalef (0.8, 0.8, 1.0);
glTranslatef (4.3, -2.0, 0.5);
glutWireIcosahedron ( );
glFlush ( );
}

void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    glViewport (0, 0, newWidth, newHeight);
    glMatrixMode (GL_PROJECTION);
    glFrustum (-1.0, 1.0, -1.0, 1.0, 2.0, 20.0);
    glMatrixMode (GL_MODELVIEW);
    glClear (GL_COLOR_BUFFER_BIT);
}

void main (int argc, char** argv)
{
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Wire-Frame Polyhedra");

    init ( );
    glutDisplayFunc (displayWirePolyhedra);
    glutReshapeFunc (winReshapeFcn);
    glutMainLoop ( );
}

3 Curved Surfaces

Equations for objects with curved boundaries can be expressed in either a parametric or a nonparametric form. The various objects that are often useful in graphics
applications include quadric surfaces, superquadrics, polynomial and exponential functions, and spline surfaces. These input object descriptions typically are tessellated to produce polygon-mesh approximations for the surfaces.

4 Quadric Surfaces

A frequently used class of objects are the quadric surfaces, which are described with second-degree equations (quadratics). They include spheres, ellipsoids, tori, paraboloids, and hyperboloids. Quadric surfaces, particularly spheres and ellipsoids, are common elements of graphics scenes, and routines for generating these surfaces are often available in graphics packages. Also, quadric surfaces can be produced with rational spline representations.

Sphere

In Cartesian coordinates, a spherical surface with radius $r$ centered on the coordinate origin is defined as the set of points $(x, y, z)$ that satisfy the equation

$$x^2 + y^2 + z^2 = r^2$$

We can also describe the spherical surface in parametric form, using latitude and longitude angles (Figure 2):

$$x = r \cos \phi \cos \theta, \quad -\pi/2 \leq \phi \leq \pi/2$$
$$y = r \cos \phi \sin \theta, \quad -\pi \leq \theta \leq \pi$$
$$z = r \sin \phi$$

The parametric representation in Equations 2 provides a symmetric range for the angular parameters $\theta$ and $\phi$. Alternatively, we could write the parametric equations using standard spherical coordinates, where angle $\phi$ is specified as the colatitude (Figure 3). Then, $\phi$ is defined over the range $0 \leq \phi \leq \pi$, and $\theta$ is often taken in the range $0 \leq \theta \leq 2\pi$. We could also set up the representation using parameters $u$ and $v$ defined over the range from 0 to 1 by substituting $\phi = \pi u$ and $\theta = 2\pi v$.

Ellipsoid

An ellipsoidal surface can be described as an extension of a spherical surface where the radii in three mutually perpendicular directions can have different values (Figure 4). The Cartesian representation for points over the surface of an ellipsoid centered on the origin is

$$\left(\frac{x}{r_x}\right)^2 + \left(\frac{y}{r_y}\right)^2 + \left(\frac{z}{r_z}\right)^2 = 1$$

And a parametric representation for the ellipsoid in terms of the latitude angle $\phi$ and the longitude angle $\theta$ in Figure 2 is

$$x = r_x \cos \phi \cos \theta, \quad -\pi/2 \leq \phi \leq \pi/2$$
$$y = r_y \cos \phi \sin \theta, \quad -\pi \leq \theta \leq \pi$$
$$z = r_z \sin \phi$$
A doughnut-shaped object is called a torus or anchor ring. Most often it is described as the surface generated by rotating a circle or an ellipse about a coplanar axis line that is external to the conic. The defining parameters for a torus are then the distance of the conic center from the rotation axis and the dimensions of the conic. A torus generated by the rotation of a circle with radius \( r \) in the \( yz \) plane about the \( z \) axis is shown in Figure 5. With the circle center on the \( y \) axis, the axial radius, \( r_{\text{axial}} \), of the resulting torus is equal to the distance along the \( y \) axis to the circle center from the \( z \) axis (the rotation axis); and the cross-sectional radius of the torus is the radius of the generating circle.

The equation for the cross-sectional circle shown in the side view of Figure 5 is

\[
(y - r_{\text{axial}})^2 + z^2 = r^2
\]

Rotating this circle about the \( z \) axis produces the torus whose surface positions are described with the Cartesian equation

\[
\left(\sqrt{x^2 + y^2} - r_{\text{axial}}\right)^2 + z^2 = r^2
\]

The corresponding parametric equations for the torus with a circular cross-section are

\[
x = (r_{\text{axial}} + r \cos \phi) \cos \theta, \quad -\pi \leq \phi \leq \pi
\]
\[
y = (r_{\text{axial}} + r \cos \phi) \sin \theta, \quad -\pi \leq \theta \leq \pi
\]
\[
z = r \sin \phi
\]

We could also generate a torus by rotating an ellipse, instead of a circle, about the \( z \) axis. For an ellipse in the \( yz \) plane with semimajor and semiminor axes denoted as \( r_y \) and \( r_z \), we can write the ellipse equation as

\[
\left(\frac{y - r_{\text{axial}}}{r_y}\right)^2 + \left(\frac{z}{r_z}\right)^2 = 1
\]

where \( r_{\text{axial}} \) is the distance along the \( y \) axis from the rotation \( z \) axis to the ellipse center. This generates a torus that can be described with the Cartesian equation

\[
\left(\sqrt{x^2 + y^2} - r_{\text{axial}}\right)^2 + \left(\frac{z}{r_z}\right)^2 = 1
\]
The corresponding parametric representation for the torus with an elliptical cross-section is

\[
\begin{align*}
x &= (r_{\text{axial}} + r_y \cos \phi) \cos \theta, \quad -\pi \leq \phi \leq \pi \\
y &= (r_{\text{axial}} + r_y \cos \phi) \sin \theta, \quad -\pi \leq \theta \leq \pi \\
z &= r_z \sin \phi
\end{align*}
\] (8)

Other variations on the preceding torus equations are possible. For example, we could generate a torus surface by rotating either a circle or an ellipse along an elliptical path around the rotation axis.

5 Superquadrics

The class of objects called **Superquadrics** is a generalization of the quadric representations. Superquadrics are formed by incorporating additional parameters into the quadric equations to provide increased flexibility for adjusting object shapes. One additional parameter is added to curve equations, and two additional parameters are used in surface equations.

**Superellipse**

We obtain a Cartesian representation for a superellipse from the corresponding equation for an ellipse by allowing the exponent on the \( x \) and \( y \) terms to be variable. One way to do this is to write the Cartesian superellipse equation in the form

\[
\left( \frac{x}{r_x} \right)^{2/s} + \left( \frac{y}{r_y} \right)^{2/s} = 1
\] (9)

where parameter \( s \) can be assigned any real value. When \( s = 1 \), we have an ordinary ellipse.

Corresponding parametric equations for the superellipse of Equation 9 can be expressed as

\[
\begin{align*}
x &= r_x \cos^{s} \theta, \quad -\pi \leq \theta \leq \pi \\
y &= r_y \sin^{s} \theta
\end{align*}
\] (10)

Figure 6 illustrates superellipse shapes that can be generated using various values for parameter \( s \).

**F I G U R E 6**

Superellipses plotted with values for parameter \( s \) ranging from 0.5 to 3.0 and with \( r_x = r_y \).
Superellipsoid

A Cartesian representation for a superellipsoid is obtained from the equation for an ellipsoid by incorporating two exponent parameters as follows:

\[
\left(\frac{x}{r_x}\right)^{2/s_2} + \left(\frac{y}{r_y}\right)^{2/s_2} + \left(\frac{z}{r_z}\right)^{2/s_1} = 1 \tag{11}
\]

For \(s_1 = s_2 = 1\), we have an ordinary ellipsoid.

We can then write the corresponding parametric representation for the superellipsoid of Equation 11 as

\[
x = r_x \cos^{s_1} \phi \cos^{s_2} \theta, \quad -\pi/2 \leq \phi \leq \pi/2 \]
\[
y = r_y \cos^{s_1} \phi \sin^{s_2} \theta, \quad -\pi \leq \theta \leq \pi \]
\[
z = r_z \sin^{s_1} \phi \tag{12}
\]

Color Plate 10 illustrates superellipsoid shapes that can be generated using various values for parameters \(s_1\) and \(s_2\). These and other superquadric shapes can be combined to create more complex structures, such as depictions of furniture, threaded bolts, and other hardware.

6 OpenGL Quadric-Surface and Cubic-Surface Functions

A sphere and a number of other three-dimensional quadric-surface objects can be displayed using functions that are included in the OpenGL Utility Toolkit (GLUT) and in the OpenGL Utility (GLU). In addition, GLUT has one function for displaying a teapot shape that is defined with bicubic surface patches. The GLUT functions, which are easy to incorporate into an application program, have two versions each. One version of each function displays a wire-frame surface, and the other displays the surface as a rendered set of fill-area polygon patches. With the GLUT functions, we can display a sphere, cone, torus, or the teapot. Quadric-surface GLU functions are a little more involved to set up, but they provide a few more options. With the GLU functions, we can display a sphere, cylinder, tapered cylinder, cone, flat circular ring (or hollow disk), and a section of a circular ring (or disk).

GLUT Quadric-Surface Functions

We generate a GLUT sphere with either of these two functions:

```c
    glutWireSphere (r, nLongitudes, nLatitudes);
```

or

```c
    glutSolidSphere (r, nLongitudes, nLatitudes);
```

where the sphere radius is determined by the double-precision floating-point number assigned to parameter \(r\). Parameters nLongitudes and nLatitudes are used to select the integer number of longitude and latitude lines that will be used to approximate the spherical surface as a quadrilateral mesh. Edges of the quadrilateral surface patches are straight-line approximations of the longitude and latitude lines. The sphere is defined in modeling coordinates, centered at the world-coordinate origin with its polar axis along the \(z\) axis.
A GLUT cone is obtained with

\[ \text{glutWireCone (rBase, height, nLongitudes, nLatitudes);} \]

or

\[ \text{glutSolidCone (rBase, height, nLongitudes, nLatitudes);} \]

We set double-precision, floating-point values for the radius of the cone base and for the cone height using parameters \textit{rbase} and \textit{height}, respectively. As with a GLUT sphere, parameters \textit{nLongitudes} and \textit{nLatitudes} are assigned integer values that specify the number of orthogonal surface lines for the quadrilateral mesh approximation. A cone longitude line is a straight-line segment along the cone surface from the apex to the base that lies in a plane containing the cone axis. Each latitude line is displayed as a set of straight-line segments around the circumference of a circle on the cone surface that is parallel to the cone base and that lies in a plane perpendicular to the cone axis. The cone is described in modeling coordinates, with the center of the base at the world-coordinate origin and with the cone axis along the world \textit{z} axis.

Wire-frame or surface-shaded displays of a torus with a circular cross-section are produced with

\[ \text{glutWireTorus (rCrossSection, rAxial, nConcentrics, nRadialSlices);} \]

or

\[ \text{glutSolidTorus (rCrossSection, rAxial, nConcentrics, nRadialSlices);} \]

The torus obtained with these GLUT routines can be described as the surface generated by rotating a circle with radius \textit{rCrossSection} about the coplanar \textit{z} axis, where the distance of the circle center from the \textit{z} axis is \textit{rAxial} (see Section 4). We select a size for the torus using double-precision, floating-point values for these radii in the GLUT functions. And the size of the quadrilaterals in the approximating surface mesh for the torus is set with integer values for parameters \textit{nConcentrics} and \textit{nRadialSlices}. Parameter \textit{nConcentrics} specifies the number of concentric circles (with center on the \textit{z} axis) to be used on the torus surface, and parameter \textit{nRadialSlices} specifies the number of radial slices through the torus surface. These two parameters designate the number of orthogonal grid lines over the torus surface, with the grid lines displayed as straight-line segments (the boundaries of the quadrilaterals) between intersection positions. The displayed torus is centered on the world-coordinate origin, with its axis along the world \textit{z} axis.

**GLUT Cubic-Surface Teapot Function**

During the early development of computer-graphics methods, sets of polygon-mesh data tables were constructed for the description of several three-dimensional objects that could be used to test rendering techniques. These objects included the surfaces of a Volkswagen automobile and a teapot, developed at the University of Utah. The data set for the Utah teapot, as constructed by Martin Newell in 1975, contains 306 vertices, defining 32 bicubic Bézier surface patches. Since determining the surface coordinates for a complex object is time-consuming,
these data sets, particularly the teapot surface mesh, became widely used.

We can display the teapot, as a mesh of over 1,000 bicubic surface patches, using either of the following two GLUT functions:

    glutWireTeapot (size);

or

    glutSolidTeapot (size);

The teapot surface is generated using OpenGL Bézier curve functions. Parameter size sets the double-precision floating-point value for the maximum radius of the teapot bowl. The teapot is centered on the world-coordinate origin coordinate origin with its vertical axis along the y axis.

**GLU Quadric-Surface Functions**

To generate a quadric surface using GLU functions, we need to assign a name to the quadric, activate the GLU quadric renderer, and designate values for the surface parameters. In addition, we can set other parameter values to control the appearance of a GLU quadric surface.

The following statements illustrate the basic sequence of calls for displaying a wire-frame sphere centered on the world-coordinate origin:

    GLUquadricObj *sphere1;

    sphere1 = gluNewQuadric ( );
    gluQuadricDrawStyle (sphere1, GLU_LINE);

    gluSphere (sphere1, r, nLongitudes, nLatitudes);

A name for the quadric object is defined in the first statement, and, for this example, we have chosen the name sphere1. This name is then used in other GLU functions to reference this particular quadric surface. Next, the quadric renderer is activated with the gluNewQuadric function, and then the display mode GLU_LINE is selected for sphere1 with the gluQuadricDrawStyle command. Thus, the sphere is displayed in a wire-frame form with a straight-line segment between each pair of surface vertices. Parameter r is assigned a double-precision value for the sphere radius, and the sphere surface is divided into a set of polygon facets by the equally spaced longitude and latitude lines. We specify the integer number of longitude lines and latitude lines as values for parameters nLongitudes and nLatitudes.

Three other display modes are available for GLU quadric surfaces. Using the symbolic constant GLU_POINT in the gluQuadricDrawStyle, we display a quadric surface as a point plot. For the sphere, a point is displayed at each surface vertex formed by the intersection of a longitude line and a latitude line. Another option is the symbolic constant GLU_SILHOUETTE. This produces a wire-frame display without the shared edges between two coplanar polygon facets. And with the symbolic constant GLU_FILL, we display the polygon patches as shaded fill areas.
We generate displays of the other GLU quadric-surface primitives using the same basic sequence of commands. To produce a view of a cone, cylinder, or tapered cylinder, we replace the `gluSphere` function with

```c
    gluCylinder (quadricName, rBase, rTop, height, nLongitudes, nLatitudes);
```

The base of this object is in the \(xy\) plane \((z = 0)\), and the axis is the \(z\) axis. We assign a double-precision radius value to the base of this quadric surface using parameter \(rBase\), and we assign a radius to the top of the quadric surface using parameter \(rTop\). If \(rTop = 0.0\), we get a cone; if \(rTop = rBase\), we obtain a cylinder. Otherwise, a tapered cylinder is displayed. A double-precision height value is assigned to parameter \(height\), and the surface is divided into a number of equally spaced vertical and horizontal lines as determined by the integer values assigned to parameters \(nLongitudes\) and \(nLatitudes\).

A flat, circular ring or solid disk is displayed in the \(xy\) plane \((z = 0)\) and centered on the world-coordinate origin with

```c
    gluDisk (ringName, rInner, rOuter, nRadii, nRings);
```

We set double-precision values for an inner radius and an outer radius with parameters \(rInner\) and \(rOuter\). If \(rInner = 0\), the disk is solid. Otherwise, it is displayed with a concentric hole in the center of the disk. The disk surface is divided into a set of facets with integer parameters \(nRadii\) and \(nRings\), which specify the number of radial slices to be used in the tessellation and the number of concentric circular rings, respectively. Orientation for the ring is defined with respect to the \(z\) axis, with the front of the ring facing in the \(+z\) direction and the back of the ring facing in the \(-z\) direction.

We can specify a section of a circular ring with the following GLU function:

```c
    gluPartialDisk (ringName, rInner, rOuter, nRadii, nRings, startAngle, sweepAngle);
```

The double-precision parameter \(startAngle\) designates an angular position in degrees in the \(xy\) plane measured clockwise from the positive \(y\) axis. Similarly, parameter \(sweepAngle\) denotes an angular distance in degrees from the \(startAngle\) position. Thus, a section of a flat, circular disk is displayed from angular position \(startAngle\) to \(startAngle + sweepAngle\). For example, if \(startAngle = 0.0\) and \(sweepAngle = 90.0\), then the section of the disk lying in the first quadrant of the \(xy\) plane is displayed.

Allocated memory for any GLU quadric surface can be reclaimed and the surface eliminated with

```c
    gluDeleteQuadric (quadricName);
```

Also, we can define the front and back directions for any quadric surface with the following orientation function:

```c
    gluQuadricOrientation (quadricName, normalVectorDirection);
```

Parameter \(normalVectorDirection\) is assigned either \(GLU\_OUTSIDE\) or \(GLU\_INSIDE\) to indicate a direction for the surface normal vectors, where "outside" indicates the front-face direction and "inside" indicates the
**back-face direction.** The default value is GLU_OUTSIDE. For the flat, circular ring, the default front-face direction is in the direction of the positive z axis (“above” the disk). Another option is the generation of surface-normal vectors, as follows:

```c
    gluQuadricNormals (quadricName, generationMode);
```

A symbolic constant is assigned to parameter `generationMode` to indicate how surface-normal vectors should be generated. The default is GLU_NONE, which means that no surface normals are to be generated and no lighting conditions typically are applied to the quadric surface. For flat surface shading (a constant color value for each surface), we use the symbolic constant GLU_FLAT. This produces one surface normal for each polygon facet. When other lighting and shading conditions are to be applied, we use the constant GLU_SMOOTH, which generates a normal vector for each surface vertex position.

Other options for GLU quadric surfaces include setting surface-texture parameters. In addition, we can designate a function that is to be invoked if an error occurs during the generation of a quadric surface:

```c
    gluQuadricCallback (quadricName, GLU_ERROR, function);
```

**Example Program Using GLUT and GLU Quadric-Surface Functions**

Three quadric-surface objects (sphere, cone, and cylinder) are displayed in a wire-frame representation by the following example program. We set the view-up direction as the positive z axis so that the axis for all displayed objects is vertical. The three objects are positioned at different locations within a single display window, as shown in Figure 7.
program to display objects using GLUT wire-frame Routines

#include <GL/glut.h>

GLsizei winWidth = 500, winHeight = 500; // Initial display-window size.

void init (void)
{
    glClearColor (1.0, 1.0, 1.0, 0.0); // Set display-window color.
}

void wireQuadSurfs (void)
{
    glClear (GL_COLOR_BUFFER_BIT); // Clear display window.
    glColor3f (0.0, 0.0, 1.0); // Set line-color to blue.

    /* Set viewing parameters with world z axis as view-up direction. */
    gluLookAt (2.0, 2.0, 2.0, 0.0, 0.0, 0.0, 0.0, 0.0, 1.0);

    /* Position and display GLUT wire-frame sphere. */
    glPushMatrix ( );
    glTranslatef (1.0, 1.0, 0.0);
    glutWireSphere (0.75, 8, 6);
    glPopMatrix ( );

    /* Position and display GLUT wire-frame cone. */
    glPushMatrix ( );
    glTranslatef (1.0, -0.5, 0.5);
    glutWireCone (0.7, 2.0, 7, 6);
    glPopMatrix ( );

    /* Position and display GLU wire-frame cylinder. */
    GLUquadricObj *cylinder; // Set name for GLU quadric object.
    glPushMatrix ( );
    glTranslatef (0.0, 1.2, 0.8);
    cylinder = gluNewQuadric ( );
    gluQuadricDrawStyle (cylinder, GLU_LINE);
    gluCylinder (cylinder, 0.6, 0.6, 1.5, 6, 4);
    glPopMatrix ( );

    glFlush ( );
}

void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    glViewport (0, 0, newWidth, newHeight);
    glMatrixMode (GL_PROJECTION);
    glOrtho (-2.0, 2.0, -2.0, 2.0, 0.0, 5.0);
    glMatrixMode (GL_MODELVIEW);
    glClear (GL_COLOR_BUFFER_BIT);
}

void main (int argc, char** argv)
{
    glutInit (&argc, argv);
    Three-Dimensional Object Representations
}

#include <GL/glut.h>

GLsizei winWidth = 500, winHeight = 500; // Initial display-window size.
void init (void)
{
    glClearColor (1.0, 1.0, 1.0, 0.0); // Set display-window color.
}

void wireQuadSurfs (void)
{
    glClear (GL_COLOR_BUFFER_BIT); // Clear display window.
    glColor3f (0.0, 0.0, 1.0); // Set line-color to blue.

    /* Set viewing parameters with world z axis as view-up direction. */
    gluLookAt (2.0, 2.0, 2.0, 0.0, 0.0, 0.0, 0.0, 0.0, 1.0);

    /* Position and display GLUT wire-frame sphere. */
    glPushMatrix ( );
    glTranslatef (1.0, 1.0, 0.0);
    glutWireSphere (0.75, 8, 6);
    glPopMatrix ( );

    /* Position and display GLUT wire-frame cone. */
    glPushMatrix ( );
    glTranslatef (1.0, -0.5, 0.5);
    glutWireCone (0.7, 2.0, 7, 6);
    glPopMatrix ( );

    /* Position and display GLU wire-frame cylinder. */
    GLUquadricObj *cylinder; // Set name for GLU quadric object.
    glPushMatrix ( );
    glTranslatef (0.0, 1.2, 0.8);
    cylinder = gluNewQuadric ( );
    gluQuadricDrawStyle (cylinder, GLU_LINE);
    gluCylinder (cylinder, 0.6, 0.6, 1.5, 6, 4);
    glPopMatrix ( );

    glFlush ( );
}

void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    glViewport (0, 0, newWidth, newHeight);
    glMatrixMode (GL_PROJECTION);
    glOrtho (-2.0, 2.0, -2.0, 2.0, 0.0, 5.0);
    glMatrixMode (GL_MODELVIEW);
    glClear (GL_COLOR_BUFFER_BIT);
}

void main (int argc, char** argv)
{
    glutInit (&argc, argv);
    Three-Dimensional Object Representations
}
7 Summary

Many representations have been developed for modeling the wide variety of objects and materials that we might want to display in a computer-graphics scene. In most cases, a three-dimensional object representation is rendered by a software package as a standard graphics object, whose surfaces are displayed as a polygon mesh.

Functions for displaying some common quadric surfaces, such as spheres and ellipsoids, are often available in graphics packages. Extensions of the quadrics, called superquadrics, provide additional parameters for creating a wider variety of object shapes.

Polygon surface facets for a standard graphics object can be specified in OpenGL using the polygon, triangle, or quadrilateral primitive functions. Also, GLUT routines are available for displaying the five regular polyhedra. Spheres, cones, and other quadric-surface objects can be displayed with GLUT and GLU functions, and a GLUT routine is provided for the generation of the cubic-surface Utah teapot. Tables 1 and 2 summarize the OpenGL polyhedron and quadric functions discussed in this chapter.

**Table 1**

Summary of OpenGL Polyhedron Functions

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glutWireTetrahedron</td>
<td>Displays a wire-frame tetrahedron.</td>
</tr>
<tr>
<td>glutSolidTetrahedron</td>
<td>Displays a surface-shaded tetrahedron.</td>
</tr>
<tr>
<td>glutWireCube</td>
<td>Displays a wire-frame cube.</td>
</tr>
<tr>
<td>glutSolidCube</td>
<td>Displays a surface-shaded cube.</td>
</tr>
<tr>
<td>glutWireOctahedron</td>
<td>Displays a wire-frame octahedron.</td>
</tr>
<tr>
<td>glutSolidOctahedron</td>
<td>Displays a surface-shaded octahedron.</td>
</tr>
<tr>
<td>glutWireDodecahedron</td>
<td>Displays a wire-frame dodecahedron.</td>
</tr>
<tr>
<td>glutSolidDodecahedron</td>
<td>Displays a surface-shaded dodecahedron.</td>
</tr>
<tr>
<td>glutWireIcosahedron</td>
<td>Displays a wire-frame icosahedron.</td>
</tr>
<tr>
<td>glutSolidIcosahedron</td>
<td>Displays a surface-shaded icosahedron.</td>
</tr>
</tbody>
</table>

Three-Dimensional Object Representations
TABLE 2

Summary of OpenGL Quadric-Surface and Cubic-Surface Functions

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glutWireSphere</td>
<td>Displays a wire-frame GLUT sphere.</td>
</tr>
<tr>
<td>glutSolidSphere</td>
<td>Displays a surface-shaded GLUT sphere.</td>
</tr>
<tr>
<td>glutWireCone</td>
<td>Displays a wire-frame GLUT cone.</td>
</tr>
<tr>
<td>glutSolidCone</td>
<td>Displays a surface-shaded GLUT cone.</td>
</tr>
<tr>
<td>glutWireTorus</td>
<td>Displays a wire-frame GLUT torus with a circular cross-section.</td>
</tr>
<tr>
<td>glutSolidTorus</td>
<td>Displays a surface-shaded, circular cross-section GLUT torus.</td>
</tr>
<tr>
<td>glutWireTeapot</td>
<td>Displays a wire-frame GLUT teapot.</td>
</tr>
<tr>
<td>glutSolidTeapot</td>
<td>Displays a surface-shaded GLUT teapot.</td>
</tr>
<tr>
<td>gluNewQuadric</td>
<td>Activates the GLU quadric renderer for an object name that has been defined with the declaration: GLUquadricObj *nameOfObject;</td>
</tr>
<tr>
<td>gluQuadricDrawStyle</td>
<td>Selects a display mode for a predefined GLU object name.</td>
</tr>
<tr>
<td>gluSphere</td>
<td>Displays a GLU sphere.</td>
</tr>
<tr>
<td>gluCylinder</td>
<td>Displays a GLU cone, cylinder, or tapered cylinder.</td>
</tr>
<tr>
<td>gluDisk</td>
<td>Displays a GLU flat, circular ring or solid disk.</td>
</tr>
<tr>
<td>gluPartialDisk</td>
<td>Displays a section of a GLU flat, circular ring or solid disk.</td>
</tr>
<tr>
<td>gluDeleteQuadric</td>
<td>Eliminates a GLU quadric object.</td>
</tr>
<tr>
<td>gluQuadricOrientation</td>
<td>Defines inside and outside orientations for a GLU quadric object.</td>
</tr>
<tr>
<td>gluQuadricNormals</td>
<td>Specifies how surface-normal vectors should be generated for a GLU quadric object.</td>
</tr>
<tr>
<td>gluQuadricCallback</td>
<td>Specifies a callback error function for a GLU quadric object.</td>
</tr>
</tbody>
</table>

REFERENCES


EXERCISES

1. Set up an algorithm for converting a given sphere to a polygon-mesh representation.
2. Set up an algorithm for converting a given ellipsoid to a polygon-mesh representation.
3. Set up an algorithm for converting a given cylinder to a polygon-mesh representation.
4. Set up an algorithm for converting a given superellipsoid to a polygon-mesh representation.
5. Set up an algorithm for converting a given torus with a circular cross section to a polygon mesh representation.
6. Set up an algorithm for converting a given torus with an ellipsoidal cross section to a polygon mesh representation.
7 Write a program that displays a sphere in the display window and allows the user to switch between solid and wire-frame views of the sphere, translate the sphere along any dimension, rotate the sphere around its center in any direction, and change the size of the sphere (i.e., its radius).

8 Write a program that displays a torus in the display window and allows the user to switch between solid and wire-frame views of the torus, translate the torus along any dimension, rotate the torus around its center in any direction, and change the sizes of the torus’ defining properties (i.e., the radius of its cross section ellipse and its axial radius).

9 Write a program that displays a sphere of fixed radius at world coordinate origin and allows the user to adjust the number of longitude and latitude lines used to approximate the sphere’s surface as a quadrilateral mesh. The user should also be able to switch between solid and wire-frame views of the sphere. Vary the resolution of the mesh approximation and observe the visual appearance of the sphere in both solid and wire-frame mode.

10 Write a program that displays a cylinder of fixed height and radius at world coordinate origin and allows the user to adjust the number of longitude and latitude lines used to approximate the cylinder’s surface as a quadrilateral mesh. The user should also be able to switch between solid and wire-frame views of the cylinder. Vary the resolution of the mesh approximation and observe the visual appearance of the cylinder in both solid and wire-frame mode.

IN MORE DEPTH

1 The material presented in this chapter will allow you to increase the complexity of the representations of the objects in your application by constructing more complex three-dimensional shapes. Choose the most appropriate three-dimensional shapes introduced in this chapter to replace the polygonal approximations of the objects in your application with which you have been working so far. Be sure to include at least a few curved-surface objects, using the GLU and GLUT functions for generating spheres, ellipsoids, and other quadric and cubic surfaces. Use the shaded fill areas to render the objects, not wire-frame views. Choose a reasonable setting for the number of latitude and longitude lines used to generate the polygon mesh approximation to these curved-surface objects. Write routines to call the appropriate functions and display the shapes in the appropriate positions and orientations in the scene. Use techniques in hierarchical modeling to generate objects that are better approximated as a group of these more-primitive shapes if appropriate.

2 In this exercise, you will experiment with varying the resolution of the polygon meshes that serve as the approximations to the curved-surface objects specified in the previous exercise. Choose a minimum number of latitude and longitude lines at which the representation of the objects is minimally acceptable as far as visual appearance goes. Using this as a baseline, render the scene from the previous exercise several times, each time increasing the number of latitude and longitude lines that define the mesh approximations of the objects by some fixed amount. For each setting of resolution, record the amount of time that it takes to render the scene using shaded fill areas to render the objects. Continue doing this until the resolution produces little or no noticeable difference in approximation quality. Then, make a plot of rendering time as a function of resolution parameters (number of latitude and longitude lines) and discuss the properties of the plot. Is there an ideal setting for this scene that balances visual quality with performance?
Three-Dimensional Object Representations Color Plate

Color Plate 10
Superellipsoids plotted with values for parameters $s_1$ and $s_2$ ranging from 0.0 to 2.5 and with $r_x = r_y = r_z$. 

Splines are another example of boundary representation modeling techniques. In drafting terminology, a spline is a flexible strip used to produce a smooth curve through a designated set of points. Several small weights are distributed along the length of the strip to hold it in position on the drafting table as the curve is drawn. The term spline curve originally referred to a curve drawn in this manner. We can mathematically describe such a curve with a piecewise cubic polynomial function whose first and second derivatives are continuous across the various curve sections. In computer graphics, the term spline curve now refers to any composite curve formed with polynomial sections satisfying any specified continuity conditions at the boundary of the pieces. A spline surface can be described with two sets of spline curves. There are several different kinds of spline specifications that are used in computer-graphics applications. Each individual specification simply refers to a particular type of polynomial with certain prescribed boundary conditions.
Splines are used to design curve and surface shapes, to digitize drawings, and to specify animation paths for the objects or the camera position in a scene. Typical computer-aided design (CAD) applications for splines include the design of automobile bodies, aircraft and spacecraft surfaces, ship hulls, and home appliances.

1 Interpolation and Approximation Splines

We specify a spline curve by giving a set of coordinate positions, called control points, which indicate the general shape of the curve. These coordinate positions are then fitted with piecewise-continuous, parametric polynomial functions in one of two ways. When polynomial sections are fitted so that all the control points are connected, as in Figure 1, the resulting curve is said to interpolate the set of control points. On the other hand, when the generated polynomial curve is plotted so that some, or all, of the control points are not on the curve path, the resulting curve is said to approximate the set of control points (Figure 2). Similar methods are used to construct interpolation or approximation spline surfaces.

Interpolation methods are commonly used to digitize drawings or to specify animation paths. Approximation methods are used primarily as design tools to create object shapes. Figure 3 shows the screen display of an approximation spline surface for a design application. Straight lines connect the control-point positions above the surface.

A spline curve or surface is defined, modified, and manipulated with operations on the control points. By interactively selecting spatial positions for the control points, a designer can set up an initial shape. After the polynomial fit is displayed for a given set of control points, the designer can then reposition some of or all the control points to restructure the shape of the object. Geometric transformations (translation, rotation, and scaling) are applied to the object by transforming the control points. In addition, CAD packages sometimes insert extra control points to aid a designer in adjusting the object shapes.

A set of control points forms a boundary for a region of space that is called the convex hull. One way to envision the shape of a convex hull for a two-dimensional curve is to imagine a rubber band stretched around the positions of the control points.
points so that each control point is either on the perimeter of this boundary or inside it (Figure 4). Thus, the convex hull for a two-dimensional spline curve is a convex polygon. In three-dimensional space, the convex hull for a set of spline control points forms a convex polyhedron. Convex hulls provide a measure for the deviation of a curve or surface from the region of space near the control points. In most cases, a spline is bounded by its convex hull, which ensures that the object shape follows the control points without erratic oscillations. Also, the convex hull provides a measure of the coordinate extents of a designed curve or surface, so it is useful in clipping and viewing routines.

A polyline connecting the sequence of control points for an approximation spline curve is usually displayed to remind a designer of the control-point positions and ordering. This set of connected line segments is called the control graph for the curve. Often the control graph is alluded to as the “control polygon” or the “characteristic polygon,” even though the control graph is a polyline and not a polygon. Figure 5 shows the shape of the control graph for the control-point sequences in Figure 4. For a spline surface, two sets of polyline control-point connectors form the edges for the polygon facets in a quadrilateral mesh for the surface control graph, as in Figure 3.

2 Parametric Continuity Conditions

To ensure a smooth transition from one section of a piecewise parametric spline to the next, we can impose various continuity conditions at the connection points. If each section of a spline curve is described with a set of parametric coordinate
functions of the form
\[ x = x(u), \quad y = y(u), \quad z = z(u), \quad u_1 \leq u \leq u_2 \quad (1) \]
we set \textit{parametric continuity} by matching the parametric derivatives of adjoining curve sections at their common boundary.

\textbf{Zero-order parametric continuity}, represented as \( C^0 \) continuity, means simply that the curves meet. That is, the values of \( x, y, \) and \( z \) evaluated at \( u_2 \) for the first curve section are equal, respectively, to the values of \( x, y, \) and \( z \) evaluated at \( u_1 \) for the next curve section. \textbf{First-order parametric continuity}, referred to as \( C^1 \) continuity, means that the first parametric derivatives (tangent lines) of the coordinate functions in Equation 1 for two successive curve sections are equal at their joining point. \textbf{Second-order parametric continuity}, or \( C^2 \) continuity, means that both the first and second parametric derivatives of the two curve sections are the same at the intersection. Higher-order parametric continuity conditions are defined similarly. Figure 6 shows examples of \( C^0, C^1, \) and \( C^2 \) continuity.

With second-order parametric continuity, the rates of change of the tangent vectors of connecting sections are equal at their intersection. Thus, the tangent line transitions smoothly from one section of the curve to the next [Figure 6(c)]. With first-order parametric continuity, however, the rate of change of tangent vectors for the two sections can be quite different [Figure 6(b)], so that the general shapes of the two adjacent sections can change abruptly. First-order parametric continuity is often sufficient for digitizing drawings and some design applications, while second-order parametric continuity is useful for setting up animation paths for camera motion and for many precision CAD requirements. A camera traveling along the curve path in Figure 6(b) with equal steps in parameter \( u \) would experience an abrupt change in acceleration at the boundary of the two sections, producing a discontinuity in the motion sequence. But if the camera was traveling along the path in Figure 6(c), the frame sequence for the motion would smoothly transition across the boundary.

\section*{3 Geometric Continuity Conditions}

Another method for joining two successive curve sections is to specify conditions for \textit{geometric continuity}. In this case, we require only that the parametric derivatives of the two sections are proportional to each other at their common boundary, instead of requiring equality.

\textbf{Zero-order geometric continuity}, described as \( G^0 \) continuity, is the same as zero-order parametric continuity. That is, two successive curve sections must have the same coordinate position at the boundary point. \textbf{First-order geometric continuity}, or \( G^1 \) continuity, means that the parametric first derivatives are proportional at the intersection of two successive sections. If we denote the parametric position on the curve as \( \mathbf{P}(u) \), the direction of the tangent vector \( \mathbf{T}(u) \), but not necessarily its magnitude, will be the same for two successive curve sections at their common point under \( G^1 \) continuity. \textbf{Second-order geometric continuity}, or \( G^2 \) continuity, means that both the first and second parametric derivatives of the two curve sections are proportional at their boundary. Under \( G^2 \) continuity, curvatures of two curve sections will match at the joining position.

A curve generated with geometric continuity conditions is similar to one generated with parametric continuity, but with slight differences in curve shape. Figure 7 provides a comparison of geometric and parametric continuity. With geometric continuity, the curve is pulled toward the section with the greater magnitude for the tangent vector.
4 Spline Specifications

There are three equivalent methods for specifying a particular spline representation, given the degree of the polynomial and the control-point positions: (1) We can state the set of boundary conditions that are imposed on the spline; or (2) we can state the matrix that characterizes the spline; or (3) we can state the set of blending functions (or basis functions) that determine how specified constraints on the curve are combined to calculate positions along the curve path.

To illustrate these three equivalent specifications, suppose we have the following parametric cubic polynomial representation for the $x$ coordinate along the path of a spline-curve section:

$$x(u) = a_x u^3 + b_x u^2 + c_x u + d_x, \quad 0 \leq u \leq 1 \quad (2)$$

Boundary conditions for this curve can be set for the endpoint coordinate positions $x(0)$ and $x(1)$ and for the parametric first derivatives at the endpoints: $x'(0)$ and $x'(1)$. These four boundary conditions are sufficient to determine the values of the four coefficients $a_x, b_x, c_x,$ and $d_x$.

From the boundary conditions, we can obtain the matrix that characterizes this spline curve by first rewriting Equation 2 as the following matrix product:

$$x(u) = \begin{bmatrix} u^3 & u^2 & u & 1 \end{bmatrix} \begin{bmatrix} a_x \\ b_x \\ c_x \\ d_x \end{bmatrix} = U \cdot C \quad (3)$$

where $U$ is the row matrix of powers of parameter $u$ and $C$ is the coefficient column matrix. Using Equation 3, we can write the boundary conditions in matrix form and solve for the coefficient matrix $C$ as

$$C = M_{\text{spline}} \cdot M_{\text{geom}} \quad (4)$$

where $M_{\text{geom}}$ is a four-element column matrix containing the geometric constraint values (boundary conditions) on the spline, and $M_{\text{spline}}$ is the 4 by 4 matrix that transforms the geometric constraint values to the polynomial coefficients and provides a characterization for the spline curve. Matrix $M_{\text{geom}}$ contains control-point coordinate values and other geometric constraints that have been specified. Thus, we can substitute the matrix representation for $C$ into Equation 3 to obtain

$$x(u) = U \cdot M_{\text{spline}} \cdot M_{\text{geom}} \quad (5)$$

The matrix $M_{\text{spline}}$, characterizing a spline representation, sometimes called the basis matrix, is particularly useful for transforming from one spline representation to another.
Finally, we can expand Equation 5 to obtain a polynomial representation for coordinate $x$ in terms of the geometric constraint parameters $g_k$, such as the control-point coordinates and slope of the curve at the control points:

$$x(u) = \sum_{k=0}^{3} g_k \cdot BF_k(u) \quad (6)$$

The polynomials $BF_k(u)$, for $k = 0, 1, 2, 3$, are called blending functions or basis functions because they combine (blend) the geometric constraint values to obtain coordinate positions along the curve. In subsequent sections, we explore the features of the various spline curves and surfaces that are useful in computer-graphics applications, including the specification of their matrix and blending-function representations.

## 5 Spline Surfaces

The usual procedure for defining a spline surface is to specify two sets of spline curves using a mesh of control points over some region of space. If we denote the control-point positions as $p_{k_u,k_v}$, then any point position on the spline surface can be computed as the product of the spline-curve blending functions as follows:

$$P(u, v) = \sum_{k_u,k_v} p_{k_u,k_v} BF_{k_u}(u) BF_{k_v}(v) \quad (7)$$

Surface parameters $u$ and $v$ often vary over the range from 0 to 1, but this range depends on the type of spline curves we use. One method for designating the three-dimensional control-point positions is to select height values above a two-dimensional mesh of positions on a ground plane.

## 6 Trimming Spline Surfaces

In CAD applications, a surface design may require some features that are not implemented just by adjusting control-point positions. For instance, a section of a spline surface may need to be snipped off to fit two design pieces together, or a hole may be needed so that a conduit can pass through the surface. For these applications, graphics packages often provide functions to generate trimming curves that can be used to take out sections of a spline surface, as illustrated in Figure 8. Trimming curves are typically defined in parametric $uv$ surface coordinates, and often they must be specified as closed curves.
7 Cubic-Spline Interpolation Methods

This class of splines is most often used to set up paths for object motions or to provide a representation for an existing object or drawing, but interpolation splines are also used sometimes to design object shapes. Cubic polynomials offer a reasonable compromise between flexibility and speed of computation. Compared to higher-order polynomials, cubic splines require less calculations and storage space, and they are more stable. Compared to quadratic polynomials and straight-line segments, cubic splines are more flexible for modeling object shapes.

Given a set of control points, cubic interpolation splines are obtained by fitting the input points with a piecewise cubic polynomial curve that passes through every control point. Suppose that we have \( n + 1 \) control points specified with coordinates

\[
p_k = (x_k, y_k, z_k), \quad k = 0, 1, 2, \ldots, n
\]

A cubic interpolation fit of these points is illustrated in Figure 9. We can describe the parametric cubic polynomial that is to be fitted between each pair of control points with the following set of equations:

\[
\begin{align*}
x(u) &= a_x u^3 + b_x u^2 + c_x u + d_x \\
y(u) &= a_y u^3 + b_y u^2 + c_y u + d_y, \quad (0 \leq u \leq 1) \\
z(u) &= a_z u^3 + b_z u^2 + c_z u + d_z
\end{align*}
\]

For each of these three equations, we need to determine the values for the four coefficients \( a, b, c, \) and \( d \) in the polynomial representation for each of the \( n \) curve sections between the \( n + 1 \) control points. We do this by setting enough boundary conditions at the control-point positions between curve sections so that we can obtain numerical values for all the coefficients. In the following sections, we discuss common methods for setting the boundary conditions for cubic interpolation splines.

Natural Cubic Splines

One of the first spline curves to be developed for graphics applications is the natural cubic spline. This interpolation curve is a mathematical representation of the original drafting spline. We formulate a natural cubic spline by requiring that two adjacent curve sections have the same first and second parametric derivatives at their common boundary. Thus, natural cubic splines have \( C^2 \) continuity.

If we have \( n + 1 \) control points, as in Figure 9, then we have \( n \) curve sections with a total of \( 4n \) polynomial coefficients to be determined. At each of the \( n - 1 \) interior control points, we have four boundary conditions: The two curve sections on either side of a control point must have the same first and second parametric derivatives at that control point, and each curve must pass through that control point. This gives us \( 4n - 4 \) equations to be satisfied by the \( 4n \) polynomial coefficients. We obtain an additional equation from the first control point \( p_0 \), the position of the beginning of the curve, and another condition from control point \( p_n \), which must be the last point on the curve. However, we still need...
two more conditions to be able to determine values for all the coefficients. One method for obtaining the two additional conditions is to set the second derivatives at \( P_0 \) and \( P_n \) equal to 0. Another approach is to add two extra control points (called *dummy points*), one at each end of the original control-point sequence. That is, we add a control point labeled \( P_{-1} \) at the beginning of the curve and a control point labeled \( P_{n+1} \) at the end. Then all the original control points are interior points, and we have the necessary \( 4n \) boundary conditions.

Although natural cubic splines are a mathematical model for the drafting spline, they have a major disadvantage. If the position of any of the control points is altered, the entire curve is affected. Thus, natural cubic splines allow for no “local control,” so that we cannot restructure part of the curve without specifying an entirely new set of control points. For this reason, other representations for a cubic-spline interpolation have been developed.

**Hermite Interpolation**

A *Hermite spline* (named after the French mathematician Charles Hermite) is an interpolating piecewise cubic polynomial with a specified tangent at each control point. Unlike the natural cubic splines, Hermite splines can be adjusted locally because each curve section depends only on its endpoint constraints.

If \( P(u) \) represents a parametric cubic point function for the curve section between control points \( P_k \) and \( P_{k+1} \), as shown in Figure 10, then the boundary conditions that define this Hermite curve section are

\[
\begin{align*}
P(0) &= P_k \\
P(1) &= P_{k+1} \\
P'(0) &= Dp_k \\
P'(1) &= Dp_{k+1}
\end{align*}
\]

with \( Dp_k \) and \( Dp_{k+1} \) specifying the values for the parametric derivatives (slope of the curve) at control points \( P_k \) and \( P_{k+1} \), respectively.

We can write the vector equivalent of Equations 8 for this Hermite curve section as

\[
P(u) = a u^3 + b u^2 + c u + d, \quad 0 \leq u \leq 1
\]

where the \( x \) component of \( P(u) \) is \( x(u) = a_x u^3 + b_x u^2 + c_x u + d_x \), and similarly for the \( y \) and \( z \) components. The matrix equivalent of Equation 10 is

\[
P(u) = \begin{bmatrix} u^3 & u^2 & u & 1 \end{bmatrix} \cdot \begin{bmatrix} a \\ b \\ c \\ d \end{bmatrix}
\]

and the derivative of the point function can be expressed as

\[
P'(u) = \begin{bmatrix} 3u^2 & 2u & 1 \end{bmatrix} \cdot \begin{bmatrix} a \\ b \\ c \\ d \end{bmatrix}
\]

Substituting endpoint values 0 and 1 for parameter \( u \) into the preceding two equations, we can express the Hermite boundary conditions 9 in the matrix form

\[
\begin{bmatrix}
P_k \\
P_{k+1} \\
Dp_k \\
Dp_{k+1}
\end{bmatrix} = \begin{bmatrix}
0 & 0 & 0 & 1 \\
1 & 1 & 1 & 1 \\
0 & 0 & 1 & 0 \\
3 & 2 & 1 & 0
\end{bmatrix} \cdot \begin{bmatrix} a \\ b \\ c \\ d \end{bmatrix}
\]
Solving this equation for the polynomial coefficients, we get
\[
\begin{bmatrix}
  a \\
  b \\
  c \\
  d
\end{bmatrix} = \begin{bmatrix}
  0 & 0 & 0 & 1 \\
  1 & 1 & 1 & 1 \\
  0 & 0 & 1 & 0 \\
  3 & 2 & 1 & 0
\end{bmatrix}^{-1} \begin{bmatrix}
  p_k \\
  p_{k+1} \\
  Dp_k \\
  Dp_{k+1}
\end{bmatrix}
\]
\[
= \begin{bmatrix}
  2 & -2 & 1 & 1 \\
  -3 & 3 & -2 & -1 \\
  0 & 0 & 1 & 0 \\
  1 & 0 & 0 & 0
\end{bmatrix} \begin{bmatrix}
  p_k \\
  p_{k+1} \\
  Dp_k \\
  Dp_{k+1}
\end{bmatrix}
\]
\[
= M_H \cdot \begin{bmatrix}
  p_k \\
  p_{k+1} \\
  Dp_k \\
  Dp_{k+1}
\end{bmatrix}
\]  

where $M_H$, the Hermite matrix, is the inverse of the boundary constraint matrix. Equation 11 can thus be written in terms of the boundary conditions as
\[
\mathbf{P}(u) = \begin{bmatrix}
  u^3 & u^2 & u & 1
\end{bmatrix} \cdot M_H \cdot \begin{bmatrix}
  p_k \\
  p_{k+1} \\
  Dp_k \\
  Dp_{k+1}
\end{bmatrix}
\]  

Finally, we can determine expressions for the polynomial Hermite blending functions, $H_k(u)$ for $k = 0, 1, 2, 3$, by carrying out the matrix multiplications in Equation 15 and collecting coefficients for the boundary constraints to obtain the polynomial form
\[
\mathbf{P}(u) = p_k(2u^3 - 3u^2 + 1) + p_{k+1}(-2u^3 + 3u^2) + Dp_k(u^3 - 2u^2 + u) + Dp_{k+1}(u^3 - u^2)
\]
\[
= p_k H_0(u) + p_{k+1} H_1 + Dp_k H_2 + Dp_{k+1} H_3
\]

Figure 11 shows the shape of the four Hermite blending functions.

Hermite polynomials can be useful for some digitizing applications, where it may not be too difficult to specify or approximate the curve slopes. But for most problems in computer graphics, it is more useful to generate spline curves without requiring input values for curve slopes or other geometric information, in addition to control-point coordinates. Cardinal splines and Kochanek-Bartels splines, discussed in the following two sections, are variations on the Hermite splines that do not require input values for the curve derivatives at the control points. Procedures for these splines compute parametric derivatives from the coordinate positions of the control points.

**Cardinal Splines**

As with Hermite splines, the **cardinal splines** are interpolating piecewise cubic polynomials with specified endpoint tangents at the boundary of each curve section. The difference is that we do not input the values for the endpoint tangents. For a cardinal spline, the slope at a control point is calculated from the coordinates of the two adjacent control points.
A cardinal spline section is completely specified with four consecutive control-point positions. The middle two control points are the section endpoints, and the other two points are used in the calculation of the endpoint slopes. If we take $P(u)$ as the representation for the parametric cubic point function for the curve section between control points $p_k$ and $p_{k+1}$, as in Figure 12, then the four control points from $p_{k-1}$ to $p_{k+1}$ are used to set the boundary conditions for the cardinal-spline section as

$$
\begin{align*}
P(0) &= p_k \\
P(1) &= p_{k+1} \\
P'(0) &= \frac{1}{2} (1 - t)(p_{k+1} - p_{k-1}) \\
P'(1) &= \frac{1}{2} (1 - t)(p_{k+2} - p_k)
\end{align*}
$$

(17)

Thus, the slopes at control points $p_k$ and $p_{k+1}$ are taken to be proportional, respectively, to the chords $p_{k-1}p_{k+1}$ and $p_kp_{k+2}$ (Figure 13). Parameter $t$ is called the tension parameter because it controls how loosely or tightly the cardinal spline fits the input control points. Figure 14 illustrates the shape of a cardinal curve for very small and very large values of tension $t$. When $t = 0$, this class of curves is referred to as Catmull-Rom splines, or Overhauser splines.
Using methods similar to those for Hermite splines, we can convert the boundary conditions 17 into the matrix form

\[ P(u) = \begin{bmatrix} u^3 & u^2 & u & 1 \end{bmatrix} \cdot M_C \cdot \begin{bmatrix} p_{k-1} \\ p_k \\ p_{k+1} \\ p_{k+2} \end{bmatrix} \]  

(18)

where the cardinal matrix is

\[ M_C = \begin{bmatrix} -s & 2 - s & s - 2 & s \\ 2s & s - 3 & 3 - 2s & -s \\ -s & 0 & s & 0 \\ 0 & 1 & 0 & 0 \end{bmatrix} \]  

(19)

with \( s = (1 - t)/2 \).

Expanding Equation 18 into polynomial form, we have

\[ P(u) = p_{k-1}(-s u^3 + 2s u^2 - s u) + p_k((2 - s)u^3 + (s - 3)u^2 + 1) + p_{k+1}((s - 2)u^3 + (3 - 2s)u^2 + s u) + p_{k+2}(s u^3 - s u^2) \]

\[ = p_{k-1} \text{CAR}_0(u) + p_k \text{CAR}_1(u) + p_{k+1} \text{CAR}_2(u) + p_{k+2} \text{CAR}_3(u) \]  

(20)

where the polynomials \( \text{CAR}_k(u) \) for \( k = 0, 1, 2, 3 \) are the cardinal-spline blending (basis) functions. Figure 15 gives a plot of the basis functions for cardinal splines with \( t = 0 \).
Examples of curves produced with the cardinal-spline blending functions are given in Figures 16, 17, and 18. In Figure 16, four cardinal-spline sections are plotted to form a closed curve. The first curve section is generated using the control-point set \( \{p_0, p_1, p_2, p_3\} \), the second curve is produced with the control-point set \( \{p_1, p_2, p_3, p_0\} \), the third curve section has control points \( \{p_2, p_3, p_0, p_1\} \), and the final curve section has control points \( \{p_3, p_0, p_1, p_2\} \). In Figure 17, a closed curve is obtained with a single cardinal-spline section by setting the position of the third control point to the coordinate position of the
second control point. In Figure 18, a self-intersecting cardinal-spline section is produced by setting the position of the third control point very near the coordinate position of the second control point. The resulting self-intersection is due to the constraints on the curve slope at the endpoints \(p_1\) and \(p_2\).

**Kochanek-Bartels Splines**

These interpolating cubic polynomials are extensions of the cardinal splines. Two additional parameters are introduced into the constraint equations defining Kochanek-Bartels splines to provide further flexibility in adjusting the shapes of curve sections.

Given four consecutive control points, labeled \(p_{k-1}\), \(p_k\), \(p_{k+1}\), and \(p_{k+2}\), we define the boundary conditions for a Kochanek-Bartels curve section between \(p_k\) and \(p_{k+1}\) as

\[
\begin{align*}
P(0) &= p_k \\
P(1) &= p_{k+1} \\
P'(0)_\text{in} &= \frac{1}{2}(1-t)[(1+b)(1-c)(p_k - p_{k-1}) + (1-b)(1+c)(p_{k+1} - p_k)] \\
P'(1)_\text{out} &= \frac{1}{2}(1-t)[(1+b)(1+c)(p_{k+1} - p_k) + (1-b)(1-c)(p_{k+2} - p_{k+1})]
\end{align*}
\]

where \(t\) is the tension parameter, \(b\) is the bias parameter, and \(c\) is the continuity parameter. In the Kochanek-Bartels formulation, parametric derivatives might not be continuous across section boundaries.

Tension parameter \(t\) has the same interpretation as in the cardinal spline formulation; that is, it controls the looseness or tightness of the curve sections. Bias, \(b\), is used to adjust the curvature at each end of a section so that curve sections can be skewed toward one end or the other (Figure 19). Parameter \(c\) controls the continuity of the tangent vector across the boundaries of sections. If \(c\) is assigned a nonzero value, there is a discontinuity in the slope of the curve across section boundaries.

Kochanek-Bartels splines were designed to model animation paths. In particular, abrupt changes in the motion of an object can be simulated with nonzero values for parameter \(c\). These motion changes are used in cartoon animations, for example, when a cartoon character stops quickly, changes direction, or collides with some other object.

**Bézier Spline Curves**

This spline approximation method was developed by the French engineer Pierre Bézier for use in the design of Renault automobile bodies. Bézier splines have a number of properties that make them highly useful and convenient for curve and

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8 Bézier Spline Curves

This spline approximation method was developed by the French engineer Pierre Bézier for use in the design of Renault automobile bodies. Bézier splines have a number of properties that make them highly useful and convenient for curve and
Spline Representations

surface design. They are also easy to implement. For these reasons, Bézier splines
are widely available in various CAD systems, in general graphics packages, and
in assorted drawing and painting packages.

In general, a Bézier curve section can be fitted to any number of control points,
although some graphic packages limit the number of control points to four. The
degree of the Bézier polynomial is determined by the number of control points
to be approximated and their relative position. As with the interpolation splines,
we can specify the Bézier curve path in the vicinity of the control points using
blending functions, a characterizing matrix, or boundary conditions. For general
Bézier curves, with no restrictions on the number of control points, the blending-
function specification is the most convenient representation.

Bézier Curve Equations

We first consider the general case of \( n + 1 \) control-point positions, denoted as
\( p_k = (x_k, y_k, z_k) \), with \( k \) varying from 0 to \( n \). These coordinate points are blended
to produce the following position vector \( P(u) \), which describes the path of an
approximating Bézier polynomial function between \( p_0 \) and \( p_n \):

\[
P(u) = \sum_{k=0}^{n} p_k B_{k,n}(u) , \quad 0 \leq u \leq 1
\]  

Equation 22 represents a set of three parametric equations for the individual
curve coordinates:

\[
x(u) = \sum_{k=0}^{n} x_k B_{k,n}(u) \\
y(u) = \sum_{k=0}^{n} y_k B_{k,n}(u) \\
z(u) = \sum_{k=0}^{n} z_k B_{k,n}(u)
\]

In most cases, a Bézier curve is a polynomial of a degree that is one less than
the designated number of control points: Three points generate a parabola, four
points a cubic curve, and so forth. Figure 20 demonstrates the appearance
of some Bézier curves for various selections of control points in the xy plane
\((z = 0)\). With certain control-point placements, however, we obtain degenerate
Bézier polynomials. For example, a Bézier curve generated with three collinear
control points is a straight-line segment; and a set of control points that are all at
the same coordinate position produce a Bézier “curve” that is a single point.

Recursive calculations can be used to obtain successive binomial-coefficient
values as

\[
C(n, k) = \frac{n-k+1}{k} C(n, k-1)
\]

for \( n \geq k \). Also, the Bézier blending functions satisfy the recursive relationship

\[
B_{k,n}(u) = (1-u)B_{k,n-1}(u) + u B_{k-1,n-1}(u), \quad n > k \geq 1
\]

with \( B_{k,k} = u^k \) and \( B_{0,k} = (1-u)^k \).
Example Bézier Curve-Generating Program

An implementation for calculating the Bézier blending functions and generating a two-dimensional, cubic Bézier-spline curve is given in the following program. Four control points are defined in the xy plane, and 1000 pixel positions are plotted along the curve path using a pixel width of 4. Values for the binomial coefficients are calculated in procedure `binomialCoeffs`, and coordinate positions along the curve path are calculated in procedure `computeBezPt`. These values are passed to procedure `bezier`, and pixel positions are plotted using the OpenGL point-plotting routines. Alternatively, we could have approximated the curve path with straight-line sections, using fewer points. More efficient methods for generating coordinate positions along the path of a spline curve are explored in Section 15. For this example, the world-coordinate limits are set so that only the curve points are displayed within the viewport (Figure 21). If we also wanted to plot the control-point positions, the control graph, or the convex hull, we would need to extend the limits of the world-coordinate clipping window.

```
#include <GL/glut.h>
#include <stdlib.h>
#include <math.h>

/* Set initial size of the display window. */
GLsizei winWidth = 600, winHeight = 600;

/* Set size of world-coordinate clipping window. */
GLfloat xwcMin = -50.0, xwcMax = 50.0;
GLfloat ywcMin = -50.0, ywcMax = 50.0;
```
class wcPt3D {
    public:
        GLfloat x, y, z;
};

void init (void) {
    /* Set color of display window to white. */
    glClearColor (1.0, 1.0, 1.0, 0.0);
}

void plotPoint (wcPt3D bezCurvePt) {
    glBegin (GL_POINTS);
        glVertex2f (bezCurvePt.x, bezCurvePt.y);
    glEnd ();
}

/* Compute binomial coefficients C for given value of n. */
void binomialCoeffs (GLint n, GLint * C) {
    GLint k, j;

    for (k = 0; k <= n; k++) {
        /* Compute n!/(k!(n - k)!). */
    }
C[k] = 1;
for (j = n; j >= k + 1; j--)
  C[k] *= j;
for (j = n - k; j >= 2; j--)
  C[k] /= j;
}
}
void computeBezPt (GLfloat u, wcPt3D * bezPt, GLint nCtrlPts, wcPt3D * ctrlPts, GLint * C)
{
  GLint k, n = nCtrlPts - 1;
  GLfloat bezBlendFcn;
  bezPt->x = bezPt->y = bezPt->z = 0.0;
  /* Compute blending functions and blend control points. */
  for (k = 0; k < nCtrlPts; k++) {
    bezBlendFcn = C[k] * pow(u, k) * pow(1 - u, n - k);
    bezPt->x += ctrlPts[k].x * bezBlendFcn;
    bezPt->y += ctrlPts[k].y * bezBlendFcn;
    bezPt->z += ctrlPts[k].z * bezBlendFcn;
  }
}
void bezier (wcPt3D * ctrlPts, GLint nCtrlPts, GLint nBezCurvePts)
{
  wcPt3D bezCurvePt;
  GLfloat u;
  GLint *C, k;
  /* Allocate space for binomial coefficients */
  C = new GLint[nCtrlPts];
  binomialCoeffs(nCtrlPts - 1, C);
  for (k = 0; k <= nBezCurvePts; k++) {
    u = GLfloat(k) / GLfloat(nBezCurvePts);
    computeBezPt(u, &bezCurvePt, nCtrlPts, ctrlPts, C);
    plotPoint(bezCurvePt);
  }
  delete[] C;
}
void displayFcn (void)
{
  /* Set example number of control points and number of curve positions to be plotted along the Bezier curve. */
  GLint nCtrlPts = 4, nBezCurvePts = 1000;
  wcPt3D ctrlPts[4] = { {-40.0, -40.0, 0.0}, {-10.0, 200.0, 0.0}, {10.0, -200.0, 0.0}, {40.0, 40.0, 0.0} };
  glClear(GL_COLOR_BUFFER_BIT); // Clear display window.
Properties of Bézier Curves

A very useful property of a Bézier curve is that the curve connects the first and last control points. Thus, a basic characteristic of any Bézier curve is that

\[
P(0) = p_0 \\
P(1) = p_n
\]

(28)

Values for the parametric first derivatives of a Bézier curve at the endpoints can be calculated from control-point coordinates as

\[
P'(0) = -np_0 + np_1 \\
P'(1) = np_{n-1} - np_n
\]

(29)

From these expressions, we see that the slope at the beginning of the curve is along the line joining the first two control points, and the slope at the end of the curve is along the line joining the last two endpoints. Similarly, the parametric second derivatives of a Bézier curve at the endpoints are calculated as

\[
P''(0) = n(n-1)[(p_2 - p_1) - (p_1 - p_0)] \\
P''(1) = n(n-1)[(p_{n-2} - p_{n-1}) - (p_{n-1} - p_n)]
\]

(30)
Another important property of any Bézier curve is that it lies within the convex hull (convex polygon boundary) of the control points. This follows from the fact that the Bézier blending functions are all positive and their sum is always 1:

$$\sum_{k=0}^{n} \text{BEZ}_{k,n}(u) = 1$$

(31)

so that any curve position is simply the weighted sum of the control-point positions. The convex-hull property for a Bézier curve ensures that the polynomial smoothly follows the control points without erratic oscillations.

**Design Techniques Using Bézier Curves**

A closed Bézier curve is generated when we set the last control-point position to the coordinate position of the first control point, as in the example shown in Figure 22. Also, specifying multiple control points at a single coordinate position gives more weight to that position. In Figure 23, a single coordinate position is input as two control points, and the resulting curve is pulled nearer to this position.

We can fit a Bézier curve to any number of control points, but this requires the calculation of polynomial functions of higher degree. When complicated curves are to be generated, they can be formed by piecing together several Bézier sections of lower degree. Generating smaller Bézier-curve sections also gives us better local control over the shape of the curve. Because Bézier curves connect the first and last control points, it is easy to match curve sections (zero-order continuity). Also, Bézier curves have the important property that the tangent to the curve at an endpoint is along the line joining that endpoint to the adjacent control point. Therefore, to obtain first-order continuity between curve sections, we can pick control points \(p_0\) and \(p_1\) for the next curve section to be along the same straight line as control points \(p_n\) and \(p_0\) of the preceding section (Figure 24). If the first curve section has \(n\) control points and the next curve section has \(n'\) control points, then we match curve tangents by placing control point \(p_1'\) at the position

$$p_1' = p_n + \frac{n}{n'} (p_n - p_{n-1})$$

(32)

To simplify the placement of \(p_1'\), we can require only geometric continuity and place \(p_1\) anywhere along the line of \(p_{n-1}\) and \(p_n\).

We obtain \(C^2\) continuity by using the expressions in Equations 30 to match parametric second derivatives for two adjacent Bézier sections. This establishes a coordinate position for control point \(p_2\), in addition to the fixed positions for

**Figure 22**
A closed Bézier curve generated by specifying the first and last control points at the same location.

**Figure 23**
A Bézier curve can be made to pass closer to a given coordinate position by assigning multiple control points to that position.

**Figure 24**
Piecewise approximation curve formed with two Bézier sections. Zero-order and first-order continuity is attained between the two curve sections by setting \(p_0 = p_2\) and by setting \(p_1\) along the line formed with points \(p_1\) and \(p_2\).
p₀′ and p₁′ that we need for $C^0$ and $C^1$ continuity. However, requiring second-order continuity for Bézier curve sections can be unnecessarily restrictive. This is particularly true with cubic curves, which have only four control points per section. In this case, second-order continuity fixes the position of the first three control points and leaves us only one point that we can use to adjust the shape of the curve segment.

**Cubic Bézier Curves**

Many graphics packages provide functions for displaying only cubic splines. This allows reasonable design flexibility while avoiding the increased calculations needed with higher-order polynomials. Cubic Bézier curves are generated with four control points. The four blending functions for cubic Bézier curves, obtained by substituting $n = 3$ into Equation 23, are

\[
\begin{align*}
BEZ_{0,3} & = (1 - u)^3 \\
BEZ_{1,3} & = 3u(1 - u)^2 \\
BEZ_{2,3} & = 3u^2(1 - u) \\
BEZ_{3,3} & = u^3
\end{align*}
\]

Plots of the four cubic Bézier blending functions are given in Figure 25. The form of the blending functions determine how the control points influence the shape of the curve for values of parameter $u$ over the range from 0 to 1. At $u = 0$, $u = 1$, $u = 0.5$.
Spline Representations

The only nonzero blending function is \( \text{BEZ}_{0,3} \), which has the value 1. At \( u = 1 \), the only nonzero function is \( \text{BEZ}_{3,3}(1) = 1 \). Thus, a cubic Bézier curve always begins at control point \( p_0 \) and ends at the position of control point \( p_3 \). The other functions, \( \text{BEZ}_{1,3} \) and \( \text{BEZ}_{2,3} \), influence the shape of the curve at intermediate values of the parameter \( u \) so that the resulting curve tends toward the points \( p_1 \) and \( p_2 \). Blending function \( \text{BEZ}_{1,3} \) is maximized at \( u = 1/3 \), and \( \text{BEZ}_{2,3} \) is maximized at \( u = 2/3 \).

We note in Figure 25 that each of the four blending functions is nonzero over the entire range of parameter \( u \) between the endpoint positions. Thus, Bézier curves do not allow for local control of the curve shape. If we reposition any one of the control points, the entire curve is affected.

At the end positions of the cubic Bézier curve, the parametric first derivatives (slopes) are

\[
P'(0) = 3(p_1 - p_0), \quad P'(1) = 3(p_3 - p_2)
\]

and the parametric second derivatives are

\[
P''(0) = 6(p_0 - 2p_1 + p_2), \quad P''(1) = 6(p_1 - 2p_2 + p_3)
\]

We can construct complex spline curves using a series of cubic-Bézier sections. Using expressions for the parametric derivatives, we can equate curve tangents to attain \( C^1 \) continuity between the curve sections. In addition, we could use the expressions for the second derivatives to obtain \( C^2 \) continuity, although this leaves us with no options for the placement of the first three control points.

A matrix formulation for the cubic-Bézier curve function is obtained by expanding the polynomial expressions for the blending functions and restructuring the equations as

\[
P(u) = [u^3 \quad u^2 \quad u \quad 1] \cdot \begin{bmatrix} p_0 \\ p_1 \\ p_2 \\ p_3 \end{bmatrix}
\]

where the Bézier matrix is

\[
M_{\text{bez}} = \begin{bmatrix}
-1 & 3 & -3 & 1 \\
3 & -6 & 3 & 0 \\
-3 & 3 & 0 & 0 \\
1 & 0 & 0 & 0
\end{bmatrix}
\]

(35)

We could also introduce additional parameters to allow adjustment of curve “tension” and “bias,” as we did with the interpolating splines. But more versatile types of splines (such as B-splines and beta-splines, discussed later in this chapter) are often provided with this capability.

9 Bézier Surfaces

Two sets of orthogonal Bézier curves can be used to design an object surface. The parametric vector function for the Bézier surface is formed as the tensor product of Bézier blending functions:

\[
P(u, v) = \sum_{j=0}^{m} \sum_{k=0}^{n} p_{j,k} \cdot \text{BEZ}_{j,m}(v) \cdot \text{BEZ}_{k,n}(u)
\]

(36)

with \( p_{j,k} \) specifying the location of the \((m+1)\) by \((n+1)\) control points.

Figure 26 illustrates two Bézier surface plots. The control points are connected by dashed lines, and the solid lines show curves of constant \( u \) and
constant \( v \). Each curve of constant \( u \) is plotted by varying \( v \) over the interval from 0 to 1, with \( u \) fixed at one of the values in this unit interval. Curves of constant \( v \) are plotted similarly.

Bézier surfaces have the same properties as Bézier curves, and they provide a convenient method for interactive design applications. To specify the three-dimensional coordinate positions for the control points, we could first construct a rectangular grid in the \( xy \) “ground” plane. We then choose elevations above the ground plane at the grid intersections as the \( z \)-coordinate values for the control points.

Figure 27 illustrates a surface formed with two Bézier sections. As with curves, a smooth transition from one section to the other is assured by establishing both zero-order and first-order continuity at the boundary line. Zero-order continuity is obtained by matching control points at the boundary. First-order continuity is obtained by choosing control points along a straight line across the boundary and by maintaining a constant ratio of collinear line segments for each set of specified control points across section boundaries.
10 B-Spline Curves

This spline category is the most widely used, and B-spline functions are commonly available in CAD systems and many graphics-programming packages. Like Bézier splines, B-splines are generated by approximating a set of control points. But B-splines have two advantages over Bézier splines: (1) the degree of a B-spline polynomial can be set independently of the number of control points (with certain limitations), and (2) B-splines allow local control over the shape of a spline. The tradeoff is that B-splines are more complex than Bézier splines.

B-Spline Curve Equations

We can write a general expression for the calculation of coordinate positions along a B-spline curve using a blending-function formulation as

\[ P(u) = \sum_{k=0}^{n} p_k B_{k,d}(u), \quad u_{\text{min}} \leq u \leq u_{\text{max}}, \quad 2 \leq d \leq n + 1 \]  

(37)

where \( p_k \) is an input set of \( n + 1 \) control points. There are several differences between this B-spline formulation and the expression for a Bézier spline curve. The range of parameter \( u \) now depends on how we choose the other B-spline parameters. And the B-spline blending functions \( B_{k,d} \) are polynomials of degree \( d - 1 \), where \( d \) is the degree parameter. (Sometimes parameter \( d \) is alluded to as the “order” of the polynomial, but this can be misleading because the term order is also often used to mean simply the degree of the polynomial.) The degree parameter \( d \) can be assigned any integer value in the range from 2 up to the number of control points \( (n + 1) \). Actually, we could also set the value of the degree parameter at 1, but then our “curve” is just a point plot of the control points. Local control for B-splines is achieved by defining the blending functions over subintervals of the total range of \( u \).

Blending functions for B-spline curves are defined by the Cox-deBoor recursion formulas:

\[ B_{k,1}(u) = \begin{cases} 1 & \text{if } u_k \leq u \leq u_{k+1} \\ 0 & \text{otherwise} \end{cases} \]

\[ B_{k,d}(u) = \frac{u - u_k}{u_{k+d} - u_k} B_{k,d-1}(u) + \frac{u_{k+d} - u}{u_{k+d} - u_{k+1}} B_{k+1,d-1}(u) \]  

(38)

where each blending function is defined over \( d \) subintervals of the total range of \( u \). Each subinterval endpoint \( u_j \) is referred to as a knot, and the entire set of selected subinterval endpoints is called a knot vector. We can choose any values for the subinterval endpoints, subject to the condition \( u_j \leq u_{j+1} \). Values for \( u_{\text{min}} \) and \( u_{\text{max}} \) then depend on the number of control points we select, the value we choose for the degree parameter \( d \), and how we set up the subintervals (knot vector). Because it is possible to choose the elements of the knot vector so that some denominators in the Cox-deBoor calculations evaluate to 0, this formulation assumes that any terms evaluated as 0/0 are to be assigned the value 0.

Figure 28 demonstrates the local-control characteristics of B-splines. In addition to local control, B-splines allow us to vary the number of control points used to design a curve without changing the degree of the polynomial. Also, we can increase the number of values in the knot vector to aid in curve design. When we do this, however, we must add control points because the size of the knot vector depends on parameter \( n \).
B-spline curves have the following properties:

- The polynomial curve has degree \( d - 1 \) and \( C^{d-2} \) continuity over the range of \( u \).
- For \( n+1 \) control points, the curve is described with \( n+1 \) blending functions.
- Each blending function \( B_{k,d} \) is defined over \( d \) subintervals of the total range of \( u \), starting at knot value \( u_k \).
- The range of parameter \( u \) is divided into \( n+d \) subintervals by the \( n+d+1 \) values specified in the knot vector.
- With knot values labeled as \( \{u_0, u_1, \ldots, u_{n+d}\} \), the resulting B-spline curve is defined only in the interval from knot value \( u_{d-1} \) up to knot value \( u_{n+1} \). (Some blending functions are undefined outside this interval.)
- Each section of the spline curve (between two successive knot values) is influenced by \( d \) control points.
- Any one control point can affect the shape of at most \( d \) curve sections.

In addition, a B-spline curve lies within the convex hull of at most \( d+1 \) control points, so that B-splines are tightly bound to the input positions. For any value of \( u \) in the interval from knot value \( u_{d-1} \) to \( u_{n+1} \), the sum over all basis functions is 1, as follows:

\[
\sum_{k=0}^{n} B_{k,d}(u) = 1
\]  

(39)

Given the control-point positions and the value of the degree parameter \( d \), we then need to specify the knot values to obtain the blending functions using the recurrence relations 38. There are three general classifications for knot vectors: uniform, open uniform, and nonuniform. B-splines are commonly described according to the selected knot vector class.

### Uniform Periodic B-Spline Curves

When the spacing between knot values is constant, the resulting curve is called a **uniform** B-spline. For example, we can set up a uniform knot vector as

\[
\{-1.5, -1.0, -0.5, 0.0, 0.5, 1.0, 1.5, 2.0\}
\]

Often knot values are normalized to the range between 0 and 1, as in

\[
\{0.0, 0.2, 0.4, 0.6, 0.8, 1.0\}
\]
It is convenient in many applications to set up uniform knot values with a separation of 1 and a starting value of 0. The following knot vector is an example of this specification scheme:

\[ \{0, 1, 2, 3, 4, 5, 6, 7\} \]

Uniform B-splines have **periodic** blending functions. That is, for given values of \( n \) and \( d \), all blending functions have the same shape. Each successive blending function is simply a shifted version of the previous function:

\[ B_{k,d}(u) = B_{k+1,d}(u + \Delta u) = B_{k+2,d}(u + 2\Delta u) \quad (40) \]

where \( \Delta u \) is the interval between adjacent knot values. Figure 29 shows the quadratic, uniform B-spline blending functions generated in the following example for a curve with four control points.

**Example 1 Uniform, Quadratic B-Splines**

To illustrate the formulation of B-spline blending functions for a uniform, integer knot vector, we select parameter values \( d = n = 3 \). The knot vector must then contain \( n + d + 1 = 7 \) knot values:

\[ \{0, 1, 2, 3, 4, 5, 6\} \]

and the range for parameter \( u \) is from 0 to 6, with \( n + d = 6 \) subintervals.
Each of the four blending functions spans $d = 3$ subintervals of the total range for $u$. Using the recurrence relations 38, we obtain the first blending function as

$$B_{0,3}(u) = \begin{cases} \frac{1}{2}u^2, & \text{for } 0 \leq u < 1 \\ \frac{1}{2}u(2-u) + \frac{1}{2}(u-1)(3-u), & \text{for } 1 \leq u < 2 \\ \frac{1}{2}(3-u)^2, & \text{for } 2 \leq u < 3 \end{cases}$$

We obtain the next periodic blending function using Equation 40, substituting $u-1$ for $u$ in $B_{0,3}$, and shifting the starting positions up by 1:

$$B_{1,3}(u) = \begin{cases} \frac{1}{2}(u-1)^2, & \text{for } 1 \leq u < 2 \\ \frac{1}{2}(u-1)(3-u) + \frac{1}{2}(u-2)(4-u), & \text{for } 2 \leq u < 3 \\ \frac{1}{2}(4-u)^2, & \text{for } 3 \leq u < 4 \end{cases}$$

Similarly, the remaining two periodic functions are obtained by successively shifting $B_{1,3}$ to the right:

$$B_{2,3}(u) = \begin{cases} \frac{1}{2}(u-2)^2, & \text{for } 2 \leq u < 3 \\ \frac{1}{2}(u-2)(4-u) + \frac{1}{2}(u-3)(5-u), & \text{for } 3 \leq u < 4 \\ \frac{1}{2}(5-u)^2, & \text{for } 4 \leq u < 5 \end{cases}$$

$$B_{3,3}(u) = \begin{cases} \frac{1}{2}(u-3)^2, & \text{for } 3 \leq u < 4 \\ \frac{1}{2}(u-3)(5-u) + \frac{1}{2}(u-4)(6-u), & \text{for } 4 \leq u < 5 \\ \frac{1}{2}(6-u)^2, & \text{for } 5 \leq u < 6 \end{cases}$$

A plot of the four periodic, quadratic blending functions is given in Figure 29, which demonstrates the local feature of B-splines. The first control point is multiplied by blending function $B_{0,3}(u)$. Therefore, changing the position of the first control point affects the shape of the curve only up to $u = 3$. Similarly, the last control point influences the shape of the spline curve in the interval where $B_{3,3}$ is defined.

Figure 29 also illustrates the limits of the B-spline curve for this example. All blending functions are present in the interval from $u_{d-1} = 2$ to $u_{n+1} = 4$. Below 2 and above 4, not all blending functions are present. This interval, from 2 to 4, is the range of the polynomial curve, and the interval in which Equation 39 is valid. Thus, the sum of all blending functions is 1 within this interval. Outside this interval, we cannot sum all blending functions, since they are not all defined below 2 and above 4.
Because the range of the resulting polynomial curve is from 2 to 4, we can determine the starting and ending position of the curve by evaluating the blending functions at these points to obtain

\[ P_{\text{start}} = \frac{1}{2} (p_0 + p_1), \quad P_{\text{end}} = \frac{1}{2} (p_2 + p_3) \]

Thus, the curve starts at the midposition between the first two control points and ends at the midposition between the last two control points.

We can also determine the parametric derivatives at the starting and ending positions of the curve. Taking the derivatives of the blending functions and substituting the endpoint values for parameter \( u \), we find that

\[ P'_{\text{start}} = p_1 - p_0, \quad P'_{\text{end}} = p_3 - p_2 \]

The parametric slope of the curve at the start position is parallel to the line joining the first two control points, and the parametric slope at the end of the curve is parallel to the line joining the last two control points.

An example plot of the quadratic periodic B-spline curve is given in Figure 30 for four control points selected in the xy plane.

In the preceding example, we noted that the quadratic curve starts between the first two control points and ends at a position between the last two control points. This result is valid for a quadratic periodic B-spline fitted to any number of distinct control points. In general, for higher-order polynomials, the start and end positions are each weighted averages of \( d - 1 \) control points. We can pull a spline curve closer to any control-point position by specifying that position multiple times.

General expressions for the boundary conditions for periodic B-splines can be obtained by reparameterizing the blending functions so that parameter \( u \) is mapped onto the unit interval from 0 to 1. Beginning and ending conditions are then obtained at \( u = 0 \) and \( u = 1 \).

**Cubic Periodic B-Spline Curves**

Because cubic periodic B-splines are commonly used in graphics packages, we consider the formulation for this class of splines. Periodic splines are particularly useful for generating certain closed curves. For example, the closed curve in Figure 31 can be generated in sections by cyclically specifying four of the six control points for each section. Also, if the coordinate positions for three consecutive control points are identical, the curve passes through that point.

For cubic B-spline curves, \( d = 4 \) and each blending function spans four sub-intervals of the total range of \( u \). If we are to fit the cubic to four control points, then we could use the integer knot vector

\[ \{0, 1, 2, 3, 4, 5, 6, 7\} \]

and recurrence relations 38 to obtain the periodic blending functions, as we did in the last section for quadratic periodic B-splines.

To derive the curve equations for a periodic, cubic B-spline, we consider an alternate formulation by starting with the boundary conditions and obtaining the blending functions normalized to the interval \( 0 \leq u \leq 1 \). Using this formulation, we can also obtain the characteristic matrix easily. The boundary conditions for
periodic cubic B-splines with four control points, labeled $p_0, p_1, p_2,$ and $p_3,$ are

$$P(0) = \frac{1}{6}(p_0 + 4p_1 + p_2)$$
$$P(1) = \frac{1}{6}(p_1 + 4p_2 + p_3)$$
$$P'(0) = \frac{1}{2}(p_2 - p_0)$$
$$P'(1) = \frac{1}{2}(p_3 - p_1)$$

These boundary conditions are similar to those for cardinal splines: Curve sections are defined with four control points, and parametric derivatives (slopes) at the beginning and end of each curve section are parallel to the chords joining adjacent control points. The B-spline curve section starts at a position near $p_1$ and ends at a position near $p_2.$

A matrix formulation for a cubic periodic B-spline with four control points can then be written as

$$P(u) = [u^3 \quad u^2 \quad u \quad 1] \cdot M_B \cdot \begin{bmatrix} p_0 \\ p_1 \\ p_2 \\ p_3 \end{bmatrix}$$

where the B-spline matrix for periodic cubic polynomials is

$$M_B = \frac{1}{6} \begin{bmatrix} -1 & 3 & -3 & 1 \\ 3 & -6 & 3 & 0 \\ -3 & 0 & 3 & 0 \\ 1 & 4 & 1 & 0 \end{bmatrix}$$

This matrix can be obtained by solving for the coefficients in a general cubic polynomial expression, using the specified four boundary conditions.

We can also modify the B-spline equations to include a tension parameter $t$ (as in cardinal splines). The matrix for the periodic cubic B-spline, with tension parameter $t,$ is

$$M_{B_t} = \frac{1}{6} \begin{bmatrix} -t & 12 - 9t & 9t - 12 & t \\ 3t & 12t - 18 & 18 - 15t & 0 \\ -3t & 0 & 3t & 0 \\ t & 6 - 2t & t & 0 \end{bmatrix}$$

which reduces to $M_B$ when $t = 1.$

We obtain the periodic cubic B-spline blending functions over the parameter range from 0 to 1 by expanding the matrix representation into polynomial form. For example, using the tension value $t = 1,$ we have

$$B_{0,3}(u) = \frac{1}{6}(1 - u)^3, \quad 0 \leq u \leq 1$$
$$B_{1,3}(u) = \frac{1}{6}(3u^3 - 6u^2 + 4)$$
$$B_{2,3}(u) = \frac{1}{6}(-3u^3 + 3u^2 + 3u + 1)$$
$$B_{3,3}(u) = \frac{1}{6}u^3$$
Open Uniform B-Spline Curves

This class of B-splines is a cross between uniform B-splines and nonuniform B-splines. Sometimes it is treated as a special type of uniform B-spline, and sometimes it is considered to be in the nonuniform B-spline classification. For the **open uniform** B-splines, or simply **open** B-splines, the knot spacing is uniform except at the ends, where knot values are repeated \( d \) times.

Here are two examples of open, uniform, integer knot vectors, each with a starting value of 0:

\[
\{0, 0, 1, 2, 3, 3\} \quad \text{for } d = 2 \text{ and } n = 3 \\
\{0, 0, 0, 1, 2, 2, 2\} \quad \text{for } d = 4 \text{ and } n = 4
\]

We can normalize these knot vectors to the unit interval from 0 to 1 as

\[
\{0, 0, 0.33, 0.67, 1, 1\} \quad \text{for } d = 2 \text{ and } n = 3 \\
\{0, 0, 0, 0.5, 1, 1, 1, 1\} \quad \text{for } d = 4 \text{ and } n = 4
\]

For any values of parameters \( d \) and \( n \), we can generate an open uniform knot vector with integer values using the calculations

\[
\begin{align*}
  u_j &= \begin{cases} 
  0 & \text{for } 0 \le j < d \\
  j - d + 1 & \text{for } d \le j \le n \\
  n - d + 2 & \text{for } j > n
  \end{cases}
\end{align*}
\]

for values of \( j \) ranging from 0 to \( n + d \). With this assignment, the first \( d \) knots are assigned the value 0, and the last \( d \) knots have the value \( n - d + 2 \).

Open uniform B-splines have characteristics that are very similar to Bézier splines. In fact, when \( d = n + 1 \) (degree of the polynomial is \( n \)), open B-splines reduce to Bézier splines, and all knot values are either 0 or 1. For example, with a cubic open B-spline \((d = 4)\) and four control points, the knot vector is

\[
\{0, 0, 0, 1, 1, 1\}
\]

The polynomial curve for an open B-spline connects the first and last control points. Also, the parametric slope of the curve at the first control point is parallel to the straight line formed by the first two control points, and the parametric slope at the last control point is parallel to the line defined by the last two control points. Thus, the geometric constraints for matching curve sections are the same as for Bézier curves.

As with Bézier curves, specifying multiple control points at the same coordinate position pulls any B-spline curve closer to that position. Since open B-splines start at the first control point and end at the last control point, a closed curve can be generated by setting the first and last control points at the same coordinate position.

**Example 2** Open Uniform, Quadratic B-Splines

From conditions 48 with \( d = 3 \) and \( n = 4 \) (five control points), we obtain the following eight values for the knot vector:

\[
\{u_0, u_1, u_2, u_3, u_4, u_5, u_6, u_7\} = \{0, 0, 1, 2, 3, 3, 3, 3\}
\]

The total range of \( u \) is divided into seven subintervals, and each of the five blending functions \( B_{k,3} \) is defined over three subintervals, starting at knot position \( u_k \). Thus \( B_{0,3} \) is defined from \( u_0 = 0 \) to \( u_3 = 1 \), \( B_{1,3} \) is defined
from \( u_1 = 0 \) to \( u_4 = 2 \), and \( B_{4,3} \) is defined from \( u_4 = 2 \) to \( u_7 = 3 \). Explicit polynomial expressions are obtained for the blending functions from recurrence relations 38 as

\[
B_{0,3}(u) = (1 - u)^2 \quad 0 \leq u < 1
\]

\[
B_{1,3}(u) = \begin{cases} 
\frac{1}{2}u(4 - 3u) & 0 \leq u < 1 \\
\frac{1}{2}(2 - u)^2 & 1 \leq u < 2 
\end{cases}
\]

\[
B_{2,3}(u) = \begin{cases} 
\frac{1}{2}u^2 & 0 \leq u < 1 \\
\frac{1}{2}u(2 - u) + \frac{1}{2}(u - 1)(3 - u) & 1 \leq u < 2 \\
\frac{1}{2}(3 - u)^2 & 2 \leq u < 3 
\end{cases}
\]

\[
B_{3,3}(u) = \begin{cases} 
\frac{1}{2}(u - 1)^2 & 1 \leq u < 2 \\
\frac{1}{2}(3 - u)(3u - 5) & 2 \leq u < 3 
\end{cases}
\]

\[
B_{4,3}(u) = (u - 2)^2 \quad 2 \leq u < 3
\]

Figure 32 shows the shape of the five blending functions. The local features of B-splines are again demonstrated. Blending function \( B_{0,3} \) is nonzero only in the subinterval from 0 to 1, so the first control point influences the curve only in this interval. Similarly, function \( B_{4,3} \) is 0 outside the interval from 2 to 3, and the position of the last control point does not affect the shape of the beginning and middle parts of the curve.

Matrix formulations for open B-splines are not generated as conveniently as they are for periodic uniform B-splines. This is due to the multiplicity of knot values at the beginning and end of the knot vector.

**Nonuniform B-Spline Curves**

For this class of splines, we can specify any values and intervals for the knot vector. With **nonuniform** B-splines, we can choose multiple internal knot values and unequal spacing between the knot values. Some examples are

\[
\{0, 1, 2, 3, 3, 4\} \\
\{0, 2, 2, 3, 3, 6\} \\
\{0, 0, 0, 1, 1, 3, 3, 3\} \\
\{0, 0.2, 0.6, 0.9, 1.0\}
\]

Nonuniform B-splines provide increased flexibility in controlling a curve shape. With unequally spaced intervals in the knot vector, we obtain different shapes for the blending functions in different intervals, which can be used in designing the spline features. By increasing knot multiplicity, we can produce subtle variations in the curve path and introduce discontinuities. Multiple knot values also reduce the continuity by 1 for each repeat of a particular value.
We obtain the blending functions for a nonuniform B-spline using methods similar to those discussed for uniform and open B-splines. Given a set of \( n + 1 \) control points, we set the degree of the polynomial and select the knot values. Then, using the recurrence relations, we could either obtain the set of blending functions or evaluate curve positions directly for the display of the curve. Graphics packages often restrict the knot intervals to be either 0 or 1 to reduce computations. A set of characteristic matrices can then be stored and used to compute values along the spline curve without evaluating the recurrence relations for each curve point to be plotted.
11 B-Spline Surfaces

Formulation of a B-spline surface is similar to that for Bézier splines. We can obtain a vector point function over a B-spline surface using the tensor product of B-spline blending functions in the form

\[ P(u, v) = \sum_{k_u=0}^{n_u} \sum_{k_v=0}^{n_v} p_{k_u,k_v} B_{k_u,d_u}(u) B_{k_v,d_v}(v) \]  

(49)

where the vector values for \( p_{k_u,k_v} \) specify the positions of the \((n_u + 1)\) by \((n_v + 1)\) control points.

B-spline surfaces exhibit the same properties as those of their component B-spline curves. A surface can be constructed from selected values for degree parameters \( d_u \) and \( d_v \), which set the degrees for the orthogonal surface polynomials at \( d_u - 1 \) and \( d_v - 1 \). For each surface parameter \( u \) and \( v \), we also select values for the knot vectors, which determines the parameter range for the blending functions.

12 Beta-Splines

A generalization of B-splines are the beta-splines, also referred to as \( \beta \)-splines, that are formulated by imposing geometric continuity conditions on the first and second parametric derivatives. The continuity parameters for beta-splines are called \( \beta \) parameters.

Beta-Spline Continuity Conditions

For a specified knot vector, we designate the spline sections to the left and right of a particular knot \( u_j \) with the position vectors \( P_{j-1}(u) \) and \( P_j(u) \) (Figure 33). Zero-order continuity (positional continuity), \( G^0 \), at \( u_j \) is obtained by requiring that

\[ P_{j-1}(u_j) = P_j(u_j) \]  

(50)

First-order continuity (unit tangent continuity), \( G^1 \), is obtained by requiring tangent vectors to be proportional:

\[ \beta_1 P'_{j-1}(u_j) = P'_j(u_j), \quad \beta_1 > 0 \]  

(51)

Here, parametric first derivatives are proportional, and the unit tangent vectors are continuous across the knot.

Second-order continuity (curvature vector continuity), \( G^2 \), is imposed with the condition

\[ \beta_1^2 P''_{j-1}(u_j) + \beta_2 P''_j(u_j) = P''_j(u_j) \]  

(52)

where \( \beta_2 \) can be assigned any real number and \( \beta_1 > 0 \). The curvature vector provides a measure of the amount of bending for the curve at position \( u_j \). When \( \beta_1 = 1 \) and \( \beta_2 = 0 \), beta-splines reduce to B-splines.

Parameter \( \beta_1 \) is called the bias parameter since it controls the skewness of the curve. For \( \beta_1 > 1 \), the curve tends to flatten to the right in the direction of the unit tangent vector at the knots. For \( 0 < \beta_1 < 1 \), the curve tends to flatten to the left. The effect of \( \beta_1 \) on the shape of the spline curve is shown in Figure 34.

Parameter \( \beta_2 \) is called the tension parameter since it controls how tightly or loosely the spline fits the control graph. As \( \beta_2 \) increases, the curve approaches the shape of the control graph, as shown in Figure 35.
Cubic Periodic Beta-Spline Matrix Representation

Applying the beta-spline boundary conditions to a cubic polynomial with a uniform knot vector, we obtain the matrix representation for a periodic beta-spline as

$$M_\beta = \frac{1}{\delta} \begin{bmatrix} -2\beta_1^3 & 2(\beta_2 + \beta_1^3 + \beta_1^2 + \beta_1) & -2(\beta_2 + \beta_1^2 + \beta_1 + 1) & 2 \\ 6\beta_1^3 & -3(\beta_2 + 2\beta_1^3 + 2\beta_1^2) & 3(\beta_2 + 2\beta_1^2) & 0 \\ -6\beta_1^3 & 6(\beta_1^3 - \beta_1) & 6\beta_1 & 0 \\ 2\beta_1^3 & \beta_2 + 4(\beta_1^2 + \beta_1) & 2 & 0 \end{bmatrix}$$ (53)

where $\delta = \beta_2 + 2\beta_1^3 + 4\beta_1^2 + 4\beta_1 + 2$.

We obtain the B-spline matrix $M_B$ when $\beta_1 = 1$ and $\beta_2 = 0$. And we have the B-spline tension matrix $M_B$, (Eq. 44) when

$$\beta_1 = 1, \quad \beta_2 = \frac{12}{t}(1 - t)$$

13 Rational Splines

A rational function is simply the ratio of two polynomials. Thus, a rational spline is the ratio of two spline functions. For example, a rational B-spline curve can be described with the position vector

$$P(u) = \frac{\sum_{k=0}^{n} \omega_k p_k B_{k,d}(u)}{\sum_{k=0}^{n} \omega_k B_{k,d}(u)}$$ (54)

where the $p_k$ are the $n+1$ control-point positions. Parameters $\omega_k$ are weight factors for the control points. The greater the value of a particular $\omega_k$, the closer the curve is pulled toward the control point $p_k$ weighted by that parameter. When all weight factors are set to the value 1, we have the standard B-spline curve, because the denominator in Equation 54 is then just the sum of the blending functions, which has the value 1 (Equation 39).

Rational splines have two important advantages, compared to nonrational splines. First, they provide an exact representation for quadric curves (conics), such as circles and ellipses. Nonrational splines, which are polynomials, can only approximate conics. This allows graphics packages to model all curve shapes with one representation—rational splines—without needing a library of curve...
functions to handle different design shapes. The second advantage of rational splines is that they are invariant with respect to a perspective viewing transformation. This means that we can apply a perspective viewing transformation to the control points of the rational curve, and we will obtain the correct view of the curve. Nonrational splines, on the other hand, are not invariant with respect to a perspective viewing transformation. Typically, graphics design packages use nonuniform knot vector representations for constructing rational B-splines. These splines are referred to as nonuniform rational B-splines (NURBs).

Homogeneous coordinate representations are used for rational splines because the denominator can be treated as the homogeneous factor in a four-dimensional representation of the control points. Thus, a rational spline can be thought of as the projection of a four-dimensional nonrational spline into three-dimensional space.

In general, constructing a rational B-spline representation is carried out using the same procedures that we employed to obtain a nonrational representation. Given the set of control points, the degree of the polynomial, the weighting factors, and the knot vector, we apply the recurrence relations to obtain the blending functions. With some CAD systems, we construct a conic section by specifying three points on an arc. A rational homogeneous-coordinate spline representation is then determined by computing control-point positions that would generate the selected conic type.

As an example of describing conic sections with rational splines, we can use a quadratic B-spline function \((d = 3)\), three control points, and the open knot vector \(\{0, 0, 0, 1, 1, 1\}\) which is the same as a quadratic Bézier spline. We then set the weighting functions to the values

\[
\omega_0 = \omega_2 = 1 \\
\omega_1 = \frac{r}{1-r}, \quad 0 \leq r < 1
\]

and the rational B-spline representation is

\[
P(u) = \frac{p_0 B_{0,3}(u) + \frac{r}{1-r} p_1 B_{1,3}(u) + p_2 B_{2,3}(u)}{B_{0,3}(u) + \frac{r}{1-r} B_{1,3}(u) + B_{2,3}(u)}
\]

We then obtain the various conics (Figure 36) with the following values for parameter \(r\):

- \(r > 1/2, \quad \omega_1 > 1\) Hyperbola section
- \(r = 1/2, \quad \omega_1 = 1\) Parabola section
- \(r < 1/2, \quad \omega_1 < 1\) Ellipse section
- \(r = 0, \quad \omega_1 = 0\) Straight-line segment

We can generate a one-quarter arc of a unit circle in the first quadrant of the \(xy\) plane (Figure 37) by setting \(\omega_1 = \cos \phi\) and by choosing the control points as

\[
p_0 = (0, 1), \quad p_1 = (1, 1), \quad p_2 = (1, 0)
\]

A complete circle can be obtained by generating sections in the other three quadrants using similar control-point placements. Or we could produce a complete circle from the first-quadrant section using geometric transformations in the \(xy\) plane. For example, we can reflect the one-quarter circular arc about the \(x\) and \(y\) axes to produce the circular arcs in the other three quadrants.
A homogeneous representation for a unit circular arc in the first quadrant of the $xy$ plane is

$$
\begin{bmatrix}
  x_h(u) \\
  y_h(u) \\
  z_h(u) \\
  h(u)
\end{bmatrix} =
\begin{bmatrix}
  1 - u^2 \\
  2u \\
  0 \\
  1 + u^2
\end{bmatrix}
$$

(57)

This homogeneous representation yields the parametric circle equations for the first quadrant as

$$
x = \frac{x_h(u)}{h(u)} = \frac{1 - u^2}{1 + u^2}
$$

$$
y = \frac{y_h(u)}{h(u)} = \frac{2u}{1 + u^2}
$$

(58)

### 14 Conversion Between Spline Representations

Sometimes it is desirable to be able to switch from one spline representation to another. For instance, a Bézier representation is most convenient for subdividing a spline curve, while a B-spline representation offers greater design flexibility. Therefore, we might design a curve using B-spline sections, then convert to an equivalent Bézier representation to display the object using a recursive subdivision procedure to locate coordinate positions along the curve.

Suppose that we have a spline description of an object that can be expressed with the following matrix product:

$$
P(u) = U \cdot M_{\text{spline}} \cdot M_{\text{geom}}
$$

(59)

where $M_{\text{spline}}$ is the matrix characterizing the spline representation and $M_{\text{geom}}$ is the column matrix of geometric constraints (for example, control-point coordinates). To transform to a second representation with spline matrix $M_{\text{spline2}}$, we must determine the geometric constraint matrix $M_{\text{geom2}}$ that produces the same
vector point function for the object. That is,
\[ P(u) = U \cdot M_{\text{spline2}} \cdot M_{\text{geom2}} \] (60)
or
\[ U \cdot M_{\text{spline2}} \cdot M_{\text{geom2}} = U \cdot M_{\text{spline1}} \cdot M_{\text{geom1}} \] (61)
Solving for \( M_{\text{geom2}} \), we have
\[ M_{\text{geom2}} = M_{\text{spline2}}^{-1} \cdot M_{\text{spline1}} \cdot M_{\text{geom1}} = M_{s1,s2} \cdot M_{\text{geom1}} \] (62)
Thus, the required transformation matrix that converts from the first spline representation to the second is
\[ M_{s1,s2} = M_{\text{spline2}}^{-1} \cdot M_{\text{spline1}} \] (63)
A nonuniform B-spline cannot be characterized with a general spline matrix. But we can rearrange the knot sequence to change the nonuniform B-spline to a Bézier representation. Then the Bézier matrix could be converted to any other form.

The following example calculates the transformation matrix for conversion from a periodic, cubic B-spline representation to a cubic Bézier spline representation:

\[
M_{B,\text{Bez}} = \begin{bmatrix}
-1 & 3 & -3 & 1 \\
3 & -6 & 3 & 0 \\
-3 & 3 & 0 & 0 \\
1 & 0 & 0 & 0
\end{bmatrix}^{-1} \cdot \frac{1}{6} \begin{bmatrix}
-1 & 3 & -3 & 1 \\
3 & -6 & 3 & 0 \\
-3 & 0 & 3 & 0 \\
1 & 4 & 1 & 0
\end{bmatrix}
\]
\[
= \begin{bmatrix}
1 & 4 & 1 & 0 \\
0 & 4 & 2 & 0 \\
0 & 2 & 4 & 0 \\
0 & 1 & 4 & 1
\end{bmatrix}
\] (64)
The transformation matrix for converting from a cubic Bézier representation to a periodic, cubic B-spline representation is
\[
M_{\text{Bez},B} = \begin{bmatrix}
1 & 1 & 1 & 1 \\
-6 & 2 & -2 & 6 \\
1 & -1 & 1 & 0 \\
-2 & 0 & 2 & 2 \\
1 & 2 & 1 & 0 \\
\frac{7}{6} & 3 & \frac{7}{6} & 0
\end{bmatrix}^{-1} \cdot \begin{bmatrix}
-1 & 3 & -3 & 1 \\
3 & -6 & 3 & 0 \\
-3 & 0 & 3 & 0 \\
1 & 4 & 1 & 0
\end{bmatrix}
\]
\[
= \begin{bmatrix}
6 & -7 & 2 & 0 \\
0 & 2 & -1 & 0 \\
0 & -1 & 2 & 0 \\
0 & 2 & -7 & 6
\end{bmatrix}
\] (65)

15 Displaying Spline Curves and Surfaces
To display a spline curve or surface, we must determine coordinate positions on the curve or surface that project to pixel positions on the display device. This means that we must evaluate the parametric polynomial spline functions in certain increments over the range of the functions, and several methods have been developed for accomplishing this evaluation efficiently.
Horner’s Rule
The simplest method for evaluating a polynomial, other than direct calculation of each term in succession, is Horner’s rule, which performs the calculations by successive factoring. This requires one multiplication and one addition at each step. For a polynomial of degree \( n \), there are \( n \) steps.

For example, suppose that we have a cubic-spline representation where the \( x \) coordinate is expressed as
\[
x(u) = a x u^3 + b x u^2 + c x u + d x
\]
with similar expressions for the \( y \) and \( z \) coordinates. For a particular value of parameter \( u \), we evaluate this polynomial in the following factored order:
\[
x(u) = [(a x + b x) u + c x] u + d x
\]
The calculation of each \( x \) value requires three multiplications and three additions, so that the determination of each coordinate position \((x, y, z)\) along a cubic-spline curve requires nine multiplications and nine additions.

Additional factoring manipulations could be applied to reduce the number of computations required by Horner’s method, especially for higher-order polynomials (degree greater than 3). But repeated determination of coordinate positions over the range of a spline function can be computed much faster using forward-difference calculations or spline-subdivision methods.

Forward-Difference Calculations
A fast method for evaluating polynomial functions is to generate successive values recursively by incrementing previously calculated values as, for example,
\[
x_{k+1} = x_k + \Delta x_k
\]
Thus, once we know the increment and the value of \( x_k \) at any step, we get the next value simply by adding the increment to \( x_k \). The increment \( \Delta x_k \) at each step is called the forward difference. For the parametric curve representation, we obtain the forward differences from the intervals we select for parameter \( u \). If we divide the total range of \( u \) into subintervals of fixed size \( \delta \), then two successive \( x \) positions occur at \( x_k = x(u_k) \) and \( x_{k+1} = x(u_{k+1}) \), where
\[
u_{k+1} = u_k + \delta,
\]
and \( u_0 = 0 \).

As an illustration of this method, we first consider the polynomial representation \( x(u) = a x u + b x \) for the \( x \)-coordinate position along a linear-spline curve. Two successive \( x \)-coordinate positions are represented as
\[
x_k = a x u_k + b x
\]
\[
x_{k+1} = a x (u_k + \delta) + b x
\]
Subtracting the two equations, we obtain the forward difference:
\[
\Delta x_k = x_{k+1} - x_k = a x \delta
\]
In this case, the forward difference is a constant. With higher-order polynomials, the forward difference is itself a polynomial function of parameter \( u \). This forward-difference polynomial has degree one less than the original polynomial.

For the cubic-spline representation in Equation 66, two successive \( x \)-coordinate positions have the polynomial representations
\[
x_k = a x u_k^3 + b x u_k^2 + c x u_k + d x
\]
\[
x_{k+1} = a x (u_k + \delta)^3 + b x (u_k + \delta)^2 + c x (u_k + \delta) + d x
\]
The forward difference now evaluates to
\[
\Delta x_k = 3a x \delta u_k^2 + (3a x \delta^2 + 2b x \delta) u_k + (a x \delta^3 + b x \delta^2 + c x \delta)
\]
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which is a quadratic function of parameter \( u_k \). Because \( \Delta x_k \) is a polynomial function of \( u \), we can use the same incremental procedure to obtain successive values of \( \Delta x_k \). That is,

\[
\Delta x_{k+1} = \Delta x_k + \Delta_2 x_k
\]

(74)

where the second forward difference is the linear function

\[
\Delta_2 x_k = 6a_x \delta^2 u_k + 6a_x \delta^3 + 2b_x \delta^2
\]

(75)

Repeating this process once more, we can write

\[
\Delta_2 x_{k+1} = \Delta_2 x_k + \Delta_3 x_k
\]

(76)

with the third forward difference as the constant expression

\[
\Delta_3 x_k = 6a_x \delta^3
\]

(77)

Equations 68, 74, 76, and 77 provide an incremental forward-difference calculation of points along the cubic curve. Starting at \( u_0 = 0 \) with a constant step size \( \delta \), the initial values for the \( x \) coordinate and its first two forward differences are

\[
x_0 = d_x
\]

\[
\Delta x_0 = a_x \delta^3 + b_x \delta^2 + c_x \delta
\]

(78)

Once these initial values have been computed, the calculation for each successive \( x \)-coordinate position requires only three additions.

We can apply forward-difference methods to determine positions along spline curves of any degree \( n \). Each successive coordinate position \((x, y, z)\) is evaluated with a series of \( 3n \) additions. For surfaces, the incremental calculations are applied to both parameter \( u \) and parameter \( v \).

Subdivision Methods

Recursive spline-subdivision procedures are used to repeatedly divide a given curve section in half, increasing the number of control points at each step. Subdivision methods are useful for displaying approximation spline curves since we can continue the subdivision process until the control graph approximates the curve path. Control-point coordinates can then be plotted as curve positions. Another application of subdivision is to generate more control points for shaping a curve. Thus, we could design a general curve shape with a few control points, then apply a subdivision procedure to obtain additional control points. With the added control points, we can then make fine adjustments to small sections of the curve.

Spline subdivision is applied to a Bézier curve section most easily because the curve begins at the first control point and ends at the last control point, the range of parameter \( u \) is always between 0 and 1, and it is easy to determine when the control points are “near enough” to the curve path. Bézier subdivision can be applied to other spline representations with the following sequence of actions:

1. Convert the current spline representation to a Bézier representation.
2. Apply the Bézier subdivision algorithm.
3. Convert the Bézier representation to the original spline representation.

Figure 38 shows the first step in a recursive subdivision of a cubic Bézier curve section. Positions along the Bézier curve are described with the parametric point function \( P(u) \) for \( 0 \leq u \leq 1 \). At the first subdivision step, we use the halfway point \( P(0.5) \) to divide the original curve into two segments. The first segment is then described with the point function \( P_1(s) \), and the second segment is described
with $P_2(t)$, where

$$
\begin{align*}
  s &= 2u, & \text{for } 0.0 \leq u \leq 0.5 \\
  t &= 2u - 1, & \text{for } 0.5 \leq u \leq 1.0
\end{align*}
$$

(79)

Each of the two curve segments has the same number of control points as the original curve. Also, the boundary conditions (position and parametric slope) at the ends of each of the two curve segments must match the position and slope values for the original curve function $P(u)$. This gives us four conditions for each curve segment that we can use to determine the control-point positions. For the first segment, the four control points are

$$
\begin{align*}
  p_{1,0} &= p_0 \\
  p_{1,1} &= \frac{1}{2}(p_0 + p_1) \\
  p_{1,2} &= \frac{1}{4}(p_0 + 2p_1 + p_2) \\
  p_{1,3} &= \frac{1}{8}(p_0 + 3p_1 + 3p_2 + p_3)
\end{align*}
$$

(80)

For the second segment, we obtain the four control points

$$
\begin{align*}
  p_{2,0} &= \frac{1}{8}(p_0 + 3p_1 + 3p_2 + p_3) \\
  p_{2,1} &= \frac{1}{4}(p_1 + 2p_2 + p_3) \\
  p_{2,2} &= \frac{1}{2}(p_2 + p_3) \\
  p_{2,3} &= p_3
\end{align*}
$$

(81)

An efficient order for computing the new set of control points can be set up using only add and shift (division by 2) operations as

$$
\begin{align*}
  p_{1,0} &= p_0 \\
  p_{1,1} &= \frac{1}{2}(p_0 + p_1) \\
  T &= \frac{1}{2}(p_1 + p_2) \\
  p_{1,2} &= \frac{1}{2}(p_{1,1} + T) \\
  p_{2,3} &= p_3 \\
  p_{2,2} &= \frac{1}{2}(p_2 + p_3) \\
  p_{2,1} &= \frac{1}{2}(T + p_{2,2}) \\
  p_{2,0} &= \frac{1}{2}(p_{1,2} + p_{2,1}) \\
  p_{1,3} &= p_{2,0}
\end{align*}
$$

(82)
The preceding steps can be repeated any number of times, depending on whether we are subdividing the curve to gain more control points or trying to locate approximate curve positions. When we are subdividing to obtain a set of display points, we can terminate the subdivision procedure when the curve segments are small enough. One way to determine this is to check the distance from the first control point to the last control point for each segment. If this distance is “sufficiently” small, we can stop subdividing. Another test is to check the distances between adjacent pairs of control points. Alternatively, we could stop subdividing when the set of control points for each segment is nearly along a straight-line path.

Subdivision methods can be applied to Bézier curves of any degree. For a Bézier polynomial of degree $n - 1$, the $2^n$ control points for each of the initial two curve segments are

$$p_{1,k} = \frac{1}{2^k} \sum_{j=0}^{k} C(k, j)p_j, \quad k = 0, 1, 2, \ldots, n$$

$$p_{2,k} = \frac{1}{2^{n-k}} \sum_{j=k}^{n} C(n-k, n-j)p_j$$

where $C(k, j)$ and $C(n-k, n-j)$ are the binomial coefficients.

Subdivision methods can be applied directly to nonuniform B-splines by adding values to the knot vector. In general, however, these methods are not as efficient as Bézier subdivision.

16 OpenGL Approximation-Spline Functions

Both Bézier splines and B-splines can be displayed using OpenGL functions, as well as trimming curves for spline surfaces. The core library contains the Bézier functions, and the OpenGL Utility (GLU) has the B-spline and trimming-curve functions. Bézier functions are often hardware implemented, and the GLU functions provide a B-spline interface that accesses OpenGL point-plotting and line-drawing routines.

OpenGL Bézier-Spline Curve Functions

We specify parameters and activate the routines for Bézier-curve display with the OpenGL functions

```c
glMap1* (GL_MAP1_VERTEX_3, uMin, uMax, stride, nPts, *ctrlPts);
glEnable (GL_MAP1_VERTEX_3);
```

We deactivate the routines with

```c
glDisable (GL_MAP1_VERTEX_3);
```

A suffix code of f or d is used with glMap1 to indicate either floating-point or double precision for the data values. Minimum and maximum values for the curve parameter $u$ are specified in uMin and uMax, although these values for a Bézier curve are typically set to 0 and 1.0, respectively. The three-dimensional, floating-point, Cartesian-coordinate values for the Bézier control points are listed in array ctrlPts, and the number of elements in this array is given as a positive integer using parameter nPts. Parameter stride is assigned an integer offset that indicates the number of data values between the beginning of one coordinate position in array ctrlPts and the beginning of the next coordinate position. For a list of three-dimensional control-point positions, we set stride = 3. A
higher value for \textit{stride} would be used if we specified the control points using four-dimensional homogeneous coordinates or intertwined the coordinate values with other data, such as color values. To express control-point positions in four-dimensional homogeneous coordinates \((x, y, z, h)\), we need only change the value of \textit{stride} and change the symbolic constant in \texttt{glMap1} and \texttt{glEnable} to \texttt{GL\_MAP1\_VERTEX\_4}.

After we have set up the Bézier parameters and activated the curve-generation routines, we need to evaluate positions along the spline path and display the resulting curve. A coordinate position along the curve path is calculated with

\[
glEvalCoord1^* (uValue);\]

where parameter \(uValue\) is assigned some value in the interval from \(uMin\) to \(uMax\). The suffix code for this function can be either \(f\) or \(d\), and we can also use the suffix code \(v\) to indicate that the value for the argument is given in an array. Function \texttt{glEvalCoord1} calculates a coordinate position using Equation 22 with the parameter value

\[
u = \frac{u_{\text{value}} - u_{\text{min}}}{u_{\text{max}} - u_{\text{min}}}\]  
(84)

which maps the \(uValue\) to the interval from 0 to 1.0.

When \texttt{glEvalCoord1} processes a value for the curve parameter \(u\), it generates a \texttt{glVertex3} function. To obtain a Bézier curve, we thus repeatedly invoke the \texttt{glEvalCoord1} function to produce a set of points along the curve path, using selected values in the range from \(uMin\) to \(uMax\). Joining these points with straight-line segments, we can approximate the spline curve as a polyline.

As an example of the OpenGL Bézier-curve routines, the following code uses the four control-point positions from the program in Section 8 to generate a two-dimensional cubic Bézier curve. In this instance, 50 points are plotted along the curve path, and the curve points are connected with straight-line segments. The curve path is then displayed as a blue polyline, and the control points are plotted as red points of size 5 (Figure 39).
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```c
GLfloat ctrlPts [4][3] = { {-40.0, 40.0, 0.0}, {-10.0, 200.0, 0.0},
{10.0, -200.0, 0.0}, {40.0, 40.0, 0.0} };

glMap1f (GL_MAP1_VERTEX_3, 0.0, 1.0, 3, 4, *ctrlPts);
glEnable (GL_MAP1_VERTEX_3);

GLint k;

glColor3f (0.0, 0.0, 1.0); // Set line color to blue.
glBegin (GL_LINE_STRIP); // Generate Bezier "curve".
    for (k = 0; k <= 50; k++)
        glEvalCoord1f (GLfloat (k) / 50.0);
    glEnd ( );

glColor (1.0, 0.0, 0.0); // Set point color to red.
glPointSize (5.0); // Set point size to 5.0.
glBegin (GL_POINTS); // Plot control points.
    for (k = 0; k < 4; k++)
        glVertex3fv (&ctrlPts [k][0]);
    glEnd ( );
```

Although the previous example generated a spline curve with evenly spaced parameter values, we can use the glEvalCoord1f function to obtain any spacing for parameter u. Usually, however, a spline curve is generated with evenly spaced parameter values, and OpenGL provides the following functions, which we can use to produce a set of uniformly spaced parameter values:

```c
glMapGrid1f (50, 0.0, 1.0);
glEvalMesh1 (GL_LINE, 0, 50);
```

The suffix code for glMapGrid1 can be either f or d. Parameter n specifies the integer number of equal subdivisions over the range from u1 to u2, and parameters n1 and n2 specify an integer range corresponding to u1 and u2. Parameter mode is assigned either GL_POINT or GL_LINE, depending on whether we want to display the curve using discrete points (a dotted curve) or using straight-line segments. For a curve that is to be displayed as a polyline, the output of these two functions is the same as the output from the following code, except that the argument of glEvalCoord1 is set either to u1 or to u2 if \( k = 0 \) or \( k = n \), respectively, to avoid round-off error. In other words, with mode = GL_LINE, the preceding OpenGL commands are equivalent to

```c
glBegin (GL_LINE_STRIP);
    for (k = n1; k <= n2; k++)
        glEvalCoord1f (u1 + k * (u2 - u1) / n);
    glEnd ( );
```

Thus, in the previous programming example, we could replace the block of code containing the loop for generating the Bézier curve with the following statements.

```c
glColor3f (0.0, 0.0, 1.0);
glMapGrid1f (50.0, 0.0, 1.0);
glEvalMesh1 (GL_LINE, 0, 50);
```
Using the `glMapGrid1` and `glEvalMesh1` functions, we can divide a curve into a number of segments and select the parameter spacing for each segment according to its curvature. Therefore, a segment with more oscillations could be assigned more intervals, and a flatter section of the curve could be assigned fewer intervals.

Instead of displaying Bézier curves, we can use the `glMap1` function to designate values for other kinds of data, and seven other OpenGL symbolic constants are available for this purpose. With the symbolic constant `GL_MAP1_COLOR_4`, we use the array `ctrlPts` to specify a list of four-element (red, green, blue, alpha) colors. Then a linearly interpolated set of colors can be generated for use in an application, and these generated color values do not change the current setting for the color state. Similarly, we can designate a list of values from a color-index table with `GL_MAP1_INDEX`, and a list of three-dimensional, surface-normal vectors is specified in array `ctrlPts` when we use the symbolic constant `GL_MAP1_NORMAL`. The remaining four symbolic constants are used with lists of surface-texture information.

Multiple `glMap1` functions can be activated simultaneously, and calls to `glEvalCoord1` or to `glMapGrid1` and `glEvalMesh1` then produce data points for each data type that is enabled. This allows us to generate combinations of coordinate positions, color values, surface-normal vectors, and surface-texture data. Note, however, we cannot activate `GL_MAP1_VERTEX_3` and `GL_MAP1_VERTEX_4` simultaneously, and we can activate only one of the surface-texture generators at any one time.

**OpenGL Bézier-Spline Surface Functions**

Activation and parameter specification for the OpenGL Bézier-surface routines are accomplished with

```c
    glMap2* (GL_MAP2_VERTEX_3, uMin, uMax, uStride, nuPts,
             vMin, vMax, vStride, nvPts, *ctrlPts);
    glEnable (GL_MAP2_VERTEX_3);
```

A suffix code of `f` or `d` is used with `glMap2` to indicate either floating-point or double precision for the data values. For a surface, we specify minimum and maximum values for both parameter $u$ and parameter $v$. The three-dimensional Cartesian coordinates for the Bézier control points are listed in the double-subscripted array `ctrlPts`, and the integer size of the array is given with parameters `nuPts` and `nvPts`. If control points are to be specified using four-dimensional homogeneous coordinates, we use the symbolic constant `GL_MAP2_VERTEX_4` instead of `GL_MAP2_VERTEX_3`. The integer offset between the beginning of coordinate values for control point $p_{j,k}$ and the beginning of coordinate values for $p_{j+1,k}$ is given in `uStride`; and the integer offset between the beginning of coordinate values for control point $p_{j,k}$ and the beginning of coordinate values for $p_{j,k+1}$ is given in `vStride`. This allows the coordinate data to be intertwined with other data, so that we need to specify only the offsets to locate coordinate values. We deactivate the Bézier-surface routines with

```c
    glDisable (GL_MAP2_VERTEX_3)
```

Coordinate positions on the Bézier surface can be calculated with

```c
    glEvalCoord2* (uValue, vValue);
```

or

```c
    glEvalCoord2*v (uvArray);
```
Parameter uValue is assigned some value in the interval from uMin to uMax, and parameter vValue is assigned some value in the interval from vMin to vMax. With the vector version, uvArray = (uValue, vValue). The suffix code for either function can be f or d. Function glEvalCoord2 calculates a coordinate position using Equation 36 with the parameter values

\[
\begin{align*}
\quad & u = \frac{uValue - uMin}{uMax - uMin}, \\
\quad & v = \frac{vValue - vMin}{vMax - vMin}
\end{align*}
\] (85) which maps each of uValue and vValue to the interval from 0 to 1.0.

To display a Bezier surface, we repeatedly invoke glEvalCoord2, which generates a series of glVertex3 functions. This is similar to generating a spline curve, except that we now have two parameters, u and v. For example, a surface defined with 16 control points, arranged in a 4 × 4 grid, can be displayed as a set of surface lines with the following code. The offset for the coordinate values in the u direction is 3, and the offset in the v direction is 12. Each coordinate position is specified with three values, and the y coordinate for each group of four positions is constant.

```c
GLfloat ctrlPts[4][4][3] = {
    { {-1.5, -1.5, 4.0}, {-0.5, -1.5, 2.0},
      { -1.5, -0.5, 1.0}, {-0.5, -0.5, 3.0},
      { 0.5, -0.5, 0.0}, { 1.5, -0.5, -1.0} },
    { {-1.5, 0.5, 4.0}, {-0.5, 0.5, 0.0},
      { -1.5, 1.5, -2.0}, { 0.5, 1.5, 3.0},
      { 0.5, 1.5, 0.0}, { 1.5, 1.5, -1.0} }
};

glMap2f (GL_MAP2_VERTEX_3, 0.0, 1.0, 3, 4,
        0.0, 1.0, 12, 4, &ctrlPts[0][0][0]);
glEnable (GL_MAP2_VERTEX_3);
GLint k, j;

for (k = 0; k <= 8; k++)
    {
    glBegin (GL_LINE_STRIP); // Generate Bezier surface lines.
    for (j = 0; j <= 40; j++)
        glEvalCoord2f (GLfloat (j) / 40.0, GLfloat (k) / 8.0);
    glEnd ( );
    glEnd ( );
}
```

Instead of using the glEvalCoord2 function, we can generate evenly spaced parameter values over the surface with

```c
glMapGrid2* (nu, u1, u2, nv, v1, v2);
glEvalMesh2 (mode, nu1, nu2, nv1, nv2);
```
The suffix code for glMapGrid2 is again either f or d, and parameter mode can be assigned the value GL_POINT, GL_LINE, or GL_FILL. A two-dimensional grid of points is produced, with nu equally spaced intervals between u1 and u2, and with nv equally spaced intervals between v1 and v2. The corresponding integer range for parameter u is nu1 to nu2, and the corresponding integer range for parameter v is nv1 to nv2.

For a surface that is to be displayed as a grid of polylines, the output of glMapGrid2 and glEvalMesh2 is the same as the following program sequence except for the conditions that avoid round-off error at the beginning and ending values of the loop variables. At the beginning of the loops, the argument of glEvalCoord1 is set to (u1, v1), and at the end of the loops, the argument of glEvalCoord1 is set to (u2, v2).

```c
for (k = nu1; k <= nu2; k++) {
    glBegin (GL_LINES);
    for (j = nv1; j <= nv2; j++)
        glEvalCoord2f (u1 + k * (u2 - u1) / nu,
                        v1 + j * (v2 - v1) / nv);
    glEnd ( );
}
for (j = nv1; j <= nv2; j++) {
    glBegin (GL_LINES);
    for (k = nu1; k <= nu2; k++)
        glEvalCoord2f (u1 + k * (u2 - u1) / nu,
                        v1 + j * (v2 - v1) / nv);
    glEnd ( );
}
```

Similarly, for a surface displayed as a set of filled-polygon facets (mode = GL_FILL), the output of glMapGrid2 and glEvalMesh2 is the same as the following program sequence, except for the round-off avoiding conditions for the beginning and ending values of the loop variables:

```c
for (k = nu1; k < nu2; k++) {
    glBegin (GL_QUAD_STRIP);
    for (j = nv1; j <= nv2; j++) {
        glEvalCoord2f (u1 + k * (u2 - u1) / nu,
                        v1 + j * (v2 - v1) / nv);
        glEvalCoord2f (u1 + (k + 1) * (u2 - u1) / nu,
                        v1 + j * (v2 - v1) / nv);
    }
    glEnd ( );
}
```

We can use the glMap2 function to designate values for other kinds of data, just as we did with glMap1. Similar symbolic constants, such as GL_MAP2_COLOR_4 and GL_MAP2_NORMAL, are available for this purpose. And we can activate multiple glMap2 functions to generate various data combinations.

**GLU B-Spline Curve Functions**

Although the GLU B-spline routines are referred to as NURBs functions, they can be used to generate B-splines that are neither nonuniform nor rational. Thus, we
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can use these GLU routines to display a polynomial B-spline that has uniform knot spacing. And the GLU routines can also be used to produce Bézier splines, rational or nonrational. To generate a B-spline (or Bézier spline), we need to define a name for the spline, activate the GLU B-spline renderer, and then define the spline parameters.

The following statements illustrate the basic sequence of calls for displaying a B-spline curve:

```c
GLUnurbsObj *curveName;
curveName = gluNewNurbsRenderer ( );
gluBeginCurve (curveName);
gluNurbsCurve (curveName, nknots, *knotVector, stride, *ctrlPts, degParam, GL_MAP1_VERTEX_3);
gluEndCurve (curveName);
```

In the first statement, we assign a name to the curve, then we invoke the GLU B-spline rendering routines for that curve using the `gluNewNurbsRenderer` command. A value of 0 is assigned to `curveName` when there is not enough memory available to create a B-spline curve. Inside a `gluBeginCurve`/`gluEndCurve` pair, we next state the attributes for the curve using a `gluNurbsCurve` function. This allows us to set up multiple curve sections, and each section is referenced with a distinct curve name. Parameter `knotVector` designates the set of floating-point knot values, and integer parameter `nknots` specifies the number of elements in the knot vector. The degree of the polynomial is `degParam` − 1. We list the values for the three-dimensional, control-point coordinates in array parameter `ctrlPts`, which contains `nknots` − `degParam` elements. And the integer offset between the start of successive coordinate positions in array `ctrlPts` is specified by integer parameter `stride`. If the control-point positions are contiguous (not interspersed between other data types), the value of `stride` is set to 3. We eliminate a defined B-spline with

```c
gluDeleteNurbsRenderer (curveName);
```

As an example of the use of GLU routines to display a spline curve, the following code generates a cubic, Bézier polynomial. To obtain this cubic curve, we set the degree parameter to the value 4. We use four control points, and we select an eight-element, open-uniform knot sequence with four repeated values at each end.

```c
GLfloat knotVector [8] = {0.0, 0.0, 0.0, 0.0, 1.0, 1.0, 1.0, 1.0};
GLfloat ctrlPts [4][3] = { {-4.0, 0.0, 0.0}, {-2.0, 8.0, 0.0},
                          {2.0, -8.0, 0.0}, {4.0, 0.0, 0.0} };
GLUnurbsObj *cubicBezCurve;
cubicBezCurve = gluNewNurbsRenderer ( );
gluBeginCurve (cubicBezCurve);
gluNurbsCurve (cubicBezCurve, 8, knotVector, 3, &ctrlPts [0][0],
             4, GL_MAP1_VERTEX_3);
gluEndCurve(cubicBezCurve);
```
To create a rational B-spline curve, we replace the symbolic constant `GL_MAP1_VERTEX_3` with `GL_MAP1_VERTEX_4`. Four-dimensional, homogeneous coordinates \((x_h, y_h, z_h, h)\) are then used to specify the control points, and the resulting homogeneous division produces the desired rational polynomial form.

We can also use the `gluNurbsCurve` function to specify lists of color values, normal vectors, or surface-texture properties, just as we did with the `glMap1` and `glMap2` functions. Any of the symbolic constants, such as `GL_MAP1_COLOR_4` or `GL_MAP1_NORMAL`, can be used as the last argument in the `gluNurbsCurve` function. Each call is then listed inside the `gluBeginCurve/gluEndCurve` pair, with two restrictions: We cannot list more than one function for each data type, and we must include exactly one function to generate the B-spline curve.

A B-spline curve is divided automatically into a number of sections and displayed as a polyline by the GLU routines. However, a variety of B-spline rendering options can also be selected with repeated calls to the following GLU function:

```
    gluNurbsProperty (splineName, property, value);
```

Parameter `splineName` is assigned the name of a B-spline, parameter `property` is assigned a GLU symbolic constant that identifies the rendering property that we want to set, and parameter `value` is assigned either a floating-point numerical value or a GLU symbolic constant that sets the value for the selected property. Several `gluNurbsProperty` functions can be specified following the `gluNewNurbsRenderer` statement. Many of the properties that can be set using the `gluNurbsProperty` function are surface parameters, as described in the next section.

### GLU B-Spline Surface Functions

The following statements illustrate a basic sequence of calls for generating a B-spline surface:

```
GLUnurbsObj *surfName

    surfName = gluNewNurbsRenderer ( );
    gluNurbsProperty (surfName, property1, value1);
    gluNurbsProperty (surfName, property2, value2);
    gluNurbsProperty (surfName, property3, value3);
    
    gluBeginSurface (surfName);
    gluNurbsSurface (surfName, nuKnots, uKnotVector, nvKnots,
                            vKnotVector, uStride, vStride, &ctrlPts [0][0][0],
                            uDegParam, vDegParam, GL_MAP2_VERTEX_3);
    gluEndSurface (surfName);
```

In general, the GLU statements and parameters for defining a B-spline surface are similar to those for a B-spline curve. After invoking the B-spline rendering routines with `gluNewNurbsRenderer`, we could specify a number of optional surface-property values. Attributes for the surface are then set with a `gluNurbsSurface`
call. Multiple surfaces, each with a distinct identifying name, can be defined in this way. A value of 0 is returned to variable `surfName` by the system when there is not enough memory available to store a B-spline object. Parameters `uKnotVector` and `vKnotVector` designate the arrays of floating-point knot values in the parametric $u$ and $v$ directions. We specify the number of elements in each knot vector with parameters `nuKnots` and `nvKnots`. The degree of the polynomial in parameter $u$ is given by the value of $uDegParam - 1$, and the degree of the polynomial in parameter $v$ is the value of $vDegParam - 1$. We list the floating-point values for the three-dimensional, control-point coordinates in array parameter `ctrlPts`, which contains $(nuKnots - uDegParam) \times (nvKnots - vDegParam)$ elements. The integer offset between the start of successive control points in the parametric $u$ direction is specified with integer parameter `uStride`, and the offset in the parametric $v$ direction is specified with integer parameter `vStride`. We erase a spline surface to free its allocated memory with the same function (gluDeleteNurbsRenderer) we used for a B-spline curve.

A B-spline surface, by default, is displayed automatically as a set of polygon fill areas by the GLU routines, but we can choose other display options and parameters. Nine properties, with two or more possible values for each property, can be set for a B-spline surface. As an example of property setting, the following statements specify a wire-frame, triangularly tessellated display for a surface:

```c
  gluNurbsProperty (surfName, GLU_NURBS_MODE, GLU_NURBS_TESSELLATOR);
  gluNurbsProperty (surfName, GLU_DISPLAY_MODE, GLU_OUTLINE_POLYGON);
```

The GLU tessellating routines divide the surface into a set of triangles and display each triangle as a polygon outline. In addition, these triangle primitives can be retrieved using the `gluNurbsCallback` function. Other values for property `GLU_DISPLAY_MODE` are `GLU_OUTLINE_PATCH` and `GLU_FILL` (the default value). With the value `GLU_OUTLINE_PATCH`, we also obtain a wire-frame display, but the surface is not divided into triangular sections. Instead, the original surface is outlined, along with any trimming curves that have been specified. The only other value that can be set for the property `GLU_NURBS_MODE` is `GLU_NURBS_RENDERER` (the default value), which renders objects without making tessellated data available for callback.

We set the number of sampling points per unit length with the properties `GLU_U_STEP` and `GLU_V_STEP`. The default value for each is 100. To set the $u$ or $v$ sampling values, we also must set the property `GLU_SAMPLING_METHOD` to the value `GLU_DOMAIN_DISTANCE`. Several other values can be used with the property `GLU_SAMPLING_METHOD` to specify how surface tessellation is to be carried out. Properties `GLU_SAMPLING_TOLERANCE` and `GLU_PARAMETRIC_TOLERANCE` are used to set maximum sampling lengths. By setting property `GLU_CULLING` to the value `GL_TRUE`, we can improve rendering performance by not tessellating objects that are outside the viewing volume. The default value for GLU culling is `GL_FALSE`, and the property `GLU_AUTO_LOAD_MATRIX` allows the matrices for the viewing, projection, and viewport transformations to be downloaded from the OpenGL server when its value is `GL_TRUE` (the default value). Otherwise, if we set the value to `GL_FALSE`, an application must supply these matrices using the `gluLoadSamplingMatrices` function.
To determine the current value of a B-spline property, we use the following query function:

```c
gluGetNurbsProperty (splineName, property, value);
```

For a specified `splineName` and `property`, the corresponding value is returned to parameter `value`.

When the property `GLU_AUTO_LOAD_MATRIX` is set to the value `GL_FALSE`, we invoke

```c
    gluLoadSamplingMatrices (splineName, modelviewMat, projMat, 
                           viewport);
```

This function specifies the modelview matrix, projection matrix, and viewport that are to be used in the sampling and culling routines for a spline object. The current modelview and projection matrices can be obtained with calls to the `glGetFloatv` function, and the current viewport can be obtained with a call to `glGetIntegerv`.

Various events associated with spline objects are processed using

```c
    gluNurbsCallback (splineName, event, fcn);
```

Parameter `event` is assigned a GLU symbolic constant, and parameter `fcn` specifies a function that is to be invoked when the event corresponding to the GLU constant is encountered. For example, if we set parameter `event` to `GLU_NURBS_ERROR`, then `fcn` is called when an error occurs. Other events are used by the GLU spline routines to return the OpenGL polygons generated by the tessellation process. The symbolic constant `GLU_NURBS_BEGIN` indicates the start of a primitive such as line segments, triangles, or quadrilaterals, and `GLU_NURBS_END` indicates the end of the primitive. The function argument for the beginning of a primitive is then a symbolic constant such as `GL_LINE_STRIP`, `GL_TRIANGLES`, or `GL_QUAD_STRIP`. Symbolic constant `GLU_NURBS_VERTEX` indicates that three-dimensional coordinate data are to be supplied, and a vertex function is called. Additional constants are available for indicating other data, such as color values.

Data values for the `gluNurbsCallback` function are supplied by

```c
    gluNurbsCallbackData (splineName, dataValues);
```

Parameter `splineName` is assigned the name of the spline object that is to be tessellated, and parameter `dataValues` is assigned a list of data values.

**GLU Surface-Trimming Functions**

A set of one or more two-dimensional trimming curves is specified for a B-spline surface with the following statements:

```c
    gluBeginTrim (surfName);
    gluPwlCurve (surfName, nPts, *curvePts, stride, GLU_MAP1_TRIM_2);
    ...
    ...
    gluEndTrim (surfName);
```
Parameter `surfName` is the name of the B-spline surface to be trimmed. A set of floating-point coordinates for the trimming curve is specified in array parameter `curvePts`, which contains `nPts` coordinate positions. An integer offset between successive coordinate positions is given in parameter `stride`. The specified curve coordinates are used to generate a piecewise linear trimming function for the B-spline surface. In other words, the generated trimming “curve” is a polyline. If the curve points are to be given in three-dimensional, homogeneous \((u, v, h)\) parameter space, then the final argument in `gluPwlCurve` is set to the GLU symbolic constant `GLU_MAP1_TRIM_3`.

We can also use one or more `gluNurbsCurve` functions as a trimming curve. In addition, we can construct trimming curves that are combinations of `gluPwlCurve` functions and `gluNurbsCurve` functions. Any specified GLU trimming “curve” must be nonintersecting, and it must be a closed curve.

The following code illustrates the GLU trimming functions for a cubic Bézier surface. We first set the coordinate points for an outermost trimming curve. These positions are specified in a counterclockwise direction completely around the unit square. Next, we set the coordinate points for an innermost trimming curve in two sections, and these positions are specified in a clockwise direction. And the knot vectors for both the surface and the first inner trim-curve section are set up to produce cubic Bézier curves. A plot of the inner and outer trimming curves on the unit square is shown in Figure 40.

```c
GLUnurbsObj *bezSurface;
GLfloat outerTrimPts[5][2] = { {0.0, 0.0}, {1.0, 0.0}, {1.0, 1.0},
                              {0.0, 1.0}, {0.0, 0.0} };
GLfloat innerTrimPts1[3][2] = { {0.25, 0.5}, {0.5, 0.75},
                                 {0.75, 0.5} };
GLfloat innerTrimPts2[4][2] = { {0.75, 0.5}, {0.75, 0.25},
                                 {0.25, 0.25}, {0.25, 0.5} };
GLfloat surfKnots[8] = (0.0, 0.0, 0.0, 0.0, 1.0, 1.0, 1.0, 1.0);
GLfloat trimCurveKnots[8] = (0.0, 0.0, 0.0, 0.0, 1.0, 1.0, 1.0, 1.0);
```
bezSurface = gluNewNurbsRenderer ( );

gluBeginSurface (bezSurface);
    gluNurbsSurface (bezSurface, 8, surfKnots, 8, surfKnots, 4 * 3, 3,
        &ctrlPts [0][0][0], 4, 4, GL_MAP2_VERTEX_3);
    gluBeginTrim (bezSurface);
        /* Counterclockwise outer trim curve. */
        gluPwlCurve (bezSurface, 5, &outerTrimPts [0][0], 2,
            GLU_MAP1_TRIM_2);
    gluEndTrim (bezSurface);
    gluBeginTrim (bezSurface);
        /* Clockwise inner trim-curve sections. */
        gluPwlCurve (bezSurface, 3, &innerTrimPts1 [0][0], 2,
            GLU_MAP1_TRIM_2);
        gluNurbsCurve (bezSurface, 8, trimCurveKnots, 2,
            &innerTrimPts2 [0][0], 4, GLU_MAP1_TRIM_2);
    gluEndTrim (bezSurface);
    gluEndSurface (bezSurface);

### 17 Summary

The most widely used methods for representing objects in CAD applications are the spline representations, which are piecewise continuous polynomial functions. A spline curve or surface is defined with a set of control points and the boundary conditions on the spline sections. Lines connecting the sequence of control points form the control graph, and all control points are within the convex hull of a spline object. The boundary conditions can be specified using parametric or geometric derivatives, and most spline representations use parametric boundary conditions. Interpolation splines connect all control points; approximation splines do not connect all control points. A spline surface can be described with the tensor product of two polynomials. Cubic polynomials are commonly used for the interpolation representations, which include the Hermite, cardinal, and Kochanek-Bartels splines. Bézier splines provide a simple and powerful approximation method for describing curved lines and surfaces, however the polynomial degree is determined by the number of control points and local control over curve shapes is difficult to attain. B-splines, which include Bézier splines as a special case, are a more versatile approximation representation, but they require the specification of a knot vector. Beta splines are generalizations of B-splines that are specified with geometric boundary conditions. Rational splines are formulated as the ratio of two spline representations. Rational splines can be used to describe quadrics, and they are invariant with respect to a perspective viewing transformation. A rational B-spline with a nonuniform knot vector is commonly referred to as a NURB. To determine the coordinate positions along a spline curve or surface, we can use forward-difference calculations or subdivision methods.

The core library of OpenGL contains functions for producing Bézier splines, and GLU functions are furnished for specifying B-splines and spline-surface trimming curves. Tables 1 and 2 summarize the OpenGL spline functions discussed in this chapter.
### Table 1

**Summary of OpenGL Bezier Functions**

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>glMap1</code></td>
<td>Specifies parameters for Bézier-curve display, color values, etc., and activate these routines using <code>glEnable</code>.</td>
</tr>
<tr>
<td><code>glEvalCoord1</code></td>
<td>Calculates a coordinate position for a Bézier curve.</td>
</tr>
<tr>
<td><code>glMapGrid1</code></td>
<td>Specifies the number of equally spaced subdivisions between two Bézier-curve parameters.</td>
</tr>
<tr>
<td><code>glEvalMesh1</code></td>
<td>Specifies the display mode and integer range for a Bézier-curve display.</td>
</tr>
<tr>
<td><code>glMap2</code></td>
<td>Specifies parameters for a Bézier-surface display, color values, etc., and activate these routines using <code>glEnable</code>.</td>
</tr>
<tr>
<td><code>glEvalCoord2</code></td>
<td>Calculates a coordinate position for a Bézier surface.</td>
</tr>
<tr>
<td><code>glMapGrid2</code></td>
<td>Specifies a two-dimensional grid of equally spaced subdivisions over a Bézier surface.</td>
</tr>
<tr>
<td><code>glEvalMesh2</code></td>
<td>Specifies the display mode and integer range for the two-dimensional Bézier-surface grid.</td>
</tr>
</tbody>
</table>

### Table 2

**Summary of OpenGL B-Spline Functions**

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>gluNewNurbsRenderer</code></td>
<td>Activates the GLU B-spline renderer for an object name that has been defined with the declaration <code>GLUnurbsObj *bsplineName</code>.</td>
</tr>
<tr>
<td><code>gluBeginCurve</code></td>
<td>Begins the assignment of parameter values for a specified B-spline curve with one or more sections.</td>
</tr>
<tr>
<td><code>gluEndCurve</code></td>
<td>Signals the end of the B-spline curve parameter specifications.</td>
</tr>
<tr>
<td><code>gluNurbsCurve</code></td>
<td>Specifies the parameter values for a named B-spline curve section.</td>
</tr>
<tr>
<td><code>gluDeleteNurbsRenderer</code></td>
<td>Eliminates a specified B-spline.</td>
</tr>
<tr>
<td><code>gluNurbsProperty</code></td>
<td>Specifies rendering options for a designated B-spline.</td>
</tr>
<tr>
<td><code>gluGetNurbsProperty</code></td>
<td>Determines the current value of a designated property for a particular B-spline.</td>
</tr>
</tbody>
</table>
Spline Representations

**TABLE 2**

Summary of OpenGL B-Spline Functions (Continued)

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>gluBeginSurface</td>
<td>Begins the assignment of parameter values for a specified B-spline surface with one or more sections.</td>
</tr>
<tr>
<td>gluEndSurface</td>
<td>Signals the end of the B-spline surface parameter specifications.</td>
</tr>
<tr>
<td>gluNurbsSurface</td>
<td>Specifies the parameter values for a named B-spline surface.</td>
</tr>
<tr>
<td>gluLoadSamplingMatrices</td>
<td>Specifies viewing and geometric transformation matrices to be used in sampling and culling routines for a B-spline.</td>
</tr>
<tr>
<td>gluNurbsCallback</td>
<td>Specifies a callback function for a designated B-spline and associated event.</td>
</tr>
<tr>
<td>gluNurbsCallbackData</td>
<td>Specifies data values that are to be passed to the event callback function.</td>
</tr>
<tr>
<td>gluBeginTrim</td>
<td>Begins the assignment of trimming-curve parameter values for a B-spline surface.</td>
</tr>
<tr>
<td>gluEndTrim</td>
<td>Signals the end of the trimming curve parameter specifications.</td>
</tr>
<tr>
<td>gluPwlCurve</td>
<td>Specifies trimming-curve parameter values for a B-spline surface.</td>
</tr>
</tbody>
</table>

**REFERENCES**


**EXERCISES**

1. Write a routine to display a two-dimensional cardinal spline curve in the $xy$ plane along with the control points used to generate the curve. The curve should be drawn in black (on a white background) and the control points should be drawn in blue. Additionally, allow the user to modify the control points via keyboard input. The user should be able to cycle through the control points and move each one around in the $xy$ plane. The currently selected control point should be drawn in red. The curve should be redrawn each time a control point is moved.

2. Write a routine to display a two-dimensional Bézier-Bartels curve, given an input set of control points in the $xy$ plane.

3. Write a program using the routine developed in the previous exercise similar to the program in Exercise 2. Control points should be drawn in addition to the curve on a white background and the user should be able to edit the control points in the same manner. The curve should be redrawn each time a control point is moved.
5. What are the Bézier-curve blending functions for three control points specified in the $xy$ plane? Plot each function and identify the minimum and maximum blending-function values.

6. What are the Bézier-curve blending functions for five control points specified in the $xy$ plane? Plot each function and identify the minimum and maximum blending-function values.

7. Modify the program example in Section 8 to display any cubic Bézier curve, given a set of four input control points in the $xy$ plane.

8. Modify the program example in Section 8 to display a Bézier curve of degree $n – 1$, given a set of $n$ input control points in the $xy$ plane.

9. Complete the OpenGL programming example in Section 8 to display any cubic Bézier curve, given a set of four input control points in the $xy$ plane.

10. Modify the program in the previous exercise to allow the user to edit the control points using keyboard input as in Exercise 2. The currently selected control point should be drawn in red, and the others in blue. The curve should be drawn in black and redrawn each time a control point is moved.

11. Modify the OpenGL program example in Section 8 to display any spatial cubic Bézier curve, given a set of four input control points in $xyz$ space. Use an orthogonal projection to display the curve, with the viewing parameters specified as input.

12. Write a routine that can be used to design two-dimensional Bézier curve shapes that have first-order piecewise continuity. The number and position of the control points for each section of the curve are to be specified as input.

13. Use the routine developed in the previous exercise to allow the user to edit the control points using keyboard input as in Exercise 2. Controls points should be displayed in the same manner.

14. Write a routine that can be used to design two-dimensional Bézier curve shapes that have second-order piecewise continuity. The number and position of the control points for each section of the curve are to be specified as input.

15. Use the routine developed in the previous exercise to allow the user to edit the control points using keyboard input as in Exercise 2. Controls points should be displayed in the same manner.

16. Modify the program example in Section 8 to display any cubic Bézier curve, given a set of four input control points in the $xy$ plane, using the subdivision method to calculate curve points.

17. Modify the program example in Section 8 to display any cubic Bézier curve, given a set of four input control points in the $xy$ plane, using forward differences to calculate curve points.

18. What are the blending functions for a two-dimensional, uniform, periodic B-spline curve with $d = 5$?

19. What are the blending functions for a two-dimensional, uniform, periodic B-spline curve with $d = 6$?

20. Modify the programming example in Section 10 to display a two-dimensional, uniform, periodic B-spline curve, given an input set of control points, using forward differences to calculate positions along the curve path.

21. Modify the program in the previous example to display the B-spline curve using OpenGL functions.

22. Modify the program in the previous exercise to allow the user to edit the control points using keyboard input as in Exercise 2. Controls points should be displayed in the same manner.

23. Write a routine to display any specified conic in the $xy$ plane using a rational Bézier-spline representation.

24. Write a routine to display any specified conic in the $xy$ plane using a rational B-spline representation.

25. Develop an algorithm for calculating the normal vector to a Bézier surface at a given point $P(u, v)$.

26. Derive expressions for calculating the forward differences for a given quadratic curve.

27. Derive expressions for calculating the forward differences for a given cubic curve.

**IN MORE DEPTH**

1. In this chapter’s exercises, you will experiment with creating and displaying three-dimensional spline surfaces to represent some of the more complex curved objects in your application. Choose some objects that fit this category in your scene and sketch out either a Bézier spline or B-spline representation of their surfaces using the methods discussed in the chapter. Once you have chosen a representation, use the OpenGL functions for displaying spline surfaces to render the objects in the scene using the default resolution of evaluation points (or a reasonable one, in the case of Bézier surfaces). Then, use the visual rendering of the objects to adjust the spline model to improve the visual accuracy of the objects. Use trimming curves where appropriate to produce the right object shapes.

2. Experiment with varying the resolution of the polygon meshes that serve as the approximations to the spline surfaces that
you defined in the previous exercise. For Bézier surfaces, choose a minimum number of evaluation points in each dimension at which the representation of the objects is minimally acceptable as far as visual appearance goes. Do the same for any B-spline representations, varying the number of sampling points instead. Using this as a baseline, render the scene from the previous exercise several times, each time increasing the number of evaluation or sampling points that define the mesh approximations of the objects by some fixed amount. For each setting of resolution, record the amount of time that it takes to render the scene using shaded fill areas to render the objects. Continue doing this until the resolution produces little or no noticeable difference in approximation quality. Then, make a plot of rendering time as a function of resolution and discuss the properties of the plot. Is there an ideal setting for each object that balances visual quality with performance?
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A major consideration in the generation of realistic graphics displays is determining what is visible within a scene from a chosen viewing position. There are a number of approaches we can take to accomplish this, and numerous algorithms have been devised for efficient identification and display of visible objects for different types of applications. Some methods require more memory, some involve more processing time, and some apply only to special types of objects. Which method we select for a particular application can depend on such factors as the complexity of the scene, type of objects to be displayed, available equipment, and whether static or animated displays are to be generated. The various algorithms are referred to as visible-surface detection methods. Sometimes these methods are also referred to as hidden-surface elimination methods, although there can be subtle differences between identifying visible surfaces and eliminating hidden surfaces. With a wire-frame display, for example, we may not want to eliminate the hidden surfaces, but rather to display them with dashed boundaries or in some other way to retain information about their shape.


1 Classification of Visible-Surface Detection Algorithms

We can broadly classify visible-surface detection algorithms according to whether they deal with the object definitions or with their projected images. These two approaches are called object-space methods and image-space methods, respectively. An object-space method compares objects and parts of objects to each other within the scene definition to determine which surfaces, as a whole, we should label as visible. In an image-space algorithm, visibility is decided point by point at each pixel position on the projection plane. Most visible-surface algorithms use image-space methods, although object-space methods can be used effectively to locate visible surfaces in some cases. Line-display algorithms, for instance, generally use object-space methods to identify visible lines in wire-frame displays, but many image-space visible-surface algorithms can be adapted easily to visible-line detection.

Although there are major differences in the basic approaches taken by the various visible-surface detection algorithms, most use sorting and coherence methods to improve performance. Sorting is used to facilitate depth comparisons by ordering the individual surfaces in a scene according to their distance from the view plane. Coherence methods are used to take advantage of regularities in a scene. An individual scan line can be expected to contain intervals (runs) of constant pixel intensities, and scan-line patterns often change little from one line to the next. Animation frames contain changes only in the vicinity of moving objects. And constant relationships can often be established between the objects in a scene.

2 Back-Face Detection

A fast and simple object-space method for locating the back faces of a polyhedron is based on front-back tests. A point \((x, y, z)\) is behind a polygon surface if

\[Ax + By + Cz + D < 0\]  

where \(A, B, C,\) and \(D\) are the plane parameters for the polygon. When this position is along the line of sight to the surface, we must be looking at the back of the polygon. Therefore, we could use the viewing position to test for back faces.

We can simplify the back-face test by considering the direction of the normal vector \(\mathbf{N}\) for a polygon surface. If \(\mathbf{V}_{\text{view}}\) is a vector in the viewing direction from our camera position, as shown in Figure 1, then a polygon is a back face if

\[\mathbf{V}_{\text{view}} \cdot \mathbf{N} > 0\]  

Furthermore, if object descriptions have been converted to projection coordinates and our viewing direction is parallel to the viewing \(z_v\) axis, then we need to consider only the \(z\) component of the normal vector \(\mathbf{N}\).

In a right-handed viewing system with the viewing direction along the negative \(z_v\) axis (Figure 2), a polygon is a back face if the \(z\) component, \(C\), of its normal vector \(\mathbf{N}\) satisfies \(C < 0\). Also, we cannot see any face whose normal has \(z\) component \(C = 0\), because our viewing direction is grazing that polygon. Thus, in general, we can label any polygon as a back face if its normal vector has a \(z\) component value that satisfies the inequality

\[C \leq 0\]
Similar methods can be used in packages that employ a left-handed viewing system. In these packages, plane parameters $A$, $B$, $C$, and $D$ are calculated from polygon vertex coordinates specified in a clockwise direction (instead of the counterclockwise direction used in a right-handed system). Inequality 1 then remains a valid test for points behind the polygon. Also, back faces have normal vectors that point away from the viewing position and are identified by $C \geq 0$ when the viewing direction is along the positive $z_v$ axis.

By examining parameter $C$ for the different plane surfaces describing an object, we can immediately identify all the back faces. For a single convex polyhedron, such as the pyramid in Figure 2, this test identifies all the hidden surfaces in the scene, because each surface is either completely visible or completely hidden. Also, if a scene contains only nonoverlapping convex polyhedra, then again all hidden surfaces are identified with the back-face method.

For other objects, such as the concave polyhedron in Figure 3, more tests must be carried out to determine whether there are additional faces that are totally or partially obscured by other faces. A general scene can be expected to contain overlapping objects along the line of sight, and we then need to determine where the obscured objects are partly or completely hidden by other objects. In general, back-face removal can be expected to eliminate about half of the polygon surfaces in a scene from further visibility tests.

3 Depth-Buffer Method

A commonly used image-space approach for detecting visible surfaces is the depth-buffer method, which compares surface depth values throughout a scene for each pixel position on the projection plane. Each surface of a scene is processed separately, one pixel position at a time, across the surface. The algorithm is usually applied to scenes containing only polygon surfaces, because depth values can be computed very quickly and the method is easy to implement. But we could also apply the same procedures to nonplanar surfaces. This visibility-detection approach is also frequently alluded to as the z-buffer method, because object depth is usually measured along the $z$ axis of a viewing system. However, rather than using actual $z$ coordinates within the scene, depth-buffer algorithms often compute a distance from the view plane along the $z$ axis.

Figure 4 shows three surfaces at varying distances along the orthographic projection line from position $(x, y)$ on a view plane. These surfaces can be processed in any order. As each surface is processed, its depth from the view plane is compared to previously processed surfaces. If a surface is closer than any previously processed surfaces, its surface color is calculated and saved, along with its
The visible surfaces in a scene are represented by the set of surface colors that have been saved after all surface processing is completed. Implementation of the depth-buffer algorithm is typically carried out in normalized coordinates, so that depth values range from 0 at the near clipping plane (the view plane) to 1.0 at the far clipping plane.

As implied by the name of this method, two buffer areas are required. A depth buffer is used to store depth values for each (x, y) position as surfaces are processed, and the frame buffer stores the surface-color values for each pixel position. Initially, all positions in the depth buffer are set to 1.0 (maximum depth), and the frame buffer (refresh buffer) is initialized to the background color. Each surface listed in the polygon tables is then processed, one scan line at a time, by calculating the depth value at each (x, y) pixel position. This calculated depth is compared to the value previously stored in the depth buffer for that pixel position. If the calculated depth is less than the value stored in the depth buffer, the new depth value is stored. Then the surface color at that position is computed and placed in the corresponding pixel location in the frame buffer.

The depth-buffer processing steps are summarized in the following algorithm.

**Depth-Buffer Algorithm**

1. Initialize the depth buffer and frame buffer so that for all buffer positions (x, y),

   \[ \text{depthBuff} (x, y) = 1.0, \quad \text{frameBuff} (x, y) = \text{backgndColor} \]
2. Process each polygon in a scene, one at a time, as follows:
   - For each projected \((x, y)\) pixel position of a polygon, calculate the depth \(z\) (if not already known).
   - If \(z < \text{depthBuff}(x, y)\), compute the surface color at that position and set
     \[
     \text{depthBuff}(x, y) = z, \quad \text{frameBuff}(x, y) = \text{surfColor}(x, y)
     \]
     After all surfaces have been processed, the depth buffer contains depth values for the visible surfaces and the frame buffer contains the corresponding color values for those surfaces.

Given the depth values for the vertex positions of any polygon in a scene, we can calculate the depth at any other point on the plane containing the polygon. At surface position \((x, y)\), the depth is calculated from the plane equation as

\[
z = -\frac{Ax - By - D}{C} \tag{4}
\]

For any scan line (Figure 5), adjacent horizontal \(x\) positions across the line differ by \(\pm 1\), and vertical \(y\) values on adjacent scan lines differ by \(\pm 1\). If the depth of position \((x, y)\) has been determined to be \(z\), then the depth \(z'\) of the next position \((x + 1, y)\) along the scan line is obtained from Eq. 4 as

\[
z' = \frac{-A(x + 1) - By - D}{C} \tag{5}
\]

or

\[
z' = z - \frac{A}{C} \tag{6}
\]

The ratio \(-A/C\) is constant for each surface, so succeeding depth values across a scan line are obtained from preceding values with a single addition.

Processing pixel positions from left to right across each scan line, we start by calculating the depth on a left polygon edge that intersects that scan line (Figure 6). For each successive position across the scan line, we then calculate the depth value using Eq. 6.

We can implement the depth-buffer algorithm by starting at a top vertex of the polygon. Then, we could recursively calculate the \(x\) coordinate values down a left edge of the polygon. The \(x\) value for the beginning position on each scan line can be calculated from the beginning (edge) \(x\) value of the previous scan line as

\[
x' = x - \frac{1}{m}
\]
where \( m \) is the slope of the edge (Figure 7). Depth values down this edge are obtained recursively as

\[
z' = z + \frac{A/m + B}{C}
\]  

(7)

If we are processing down a vertical edge, the slope is infinite and the recursive calculations reduce to

\[
z' = z + \frac{B}{C}
\]

One slight complication with this approach is that while pixel positions are at integer \((x, y)\) coordinates, the actual point of intersection of a scan line with the edge of a polygon may not be. As a result, it may be necessary to adjust the intersection point by rounding its fractional part up or down, as is done in scan-line polygon fill algorithms.

An alternative approach is to use a midpoint method or Bresenham-type algorithm for determining the starting \(x\) values along edges for each scan line. The method can be applied to curved surfaces by determining depth and color values at each surface projection point.

For polygon surfaces, the depth-buffer method is very easy to implement, and it requires no sorting of the surfaces in a scene. But it does require the availability of a second buffer in addition to the refresh buffer. A system with a resolution of \(1280 \times 1024\), for example, would require over 1.3 million positions in the depth buffer, with each position containing enough bits to represent the number of depth increments needed. One way to reduce storage requirements is to process one section of the scene at a time, using a smaller depth buffer. After each view section is processed, the buffer is reused for the next section.

In addition, the basic depth-buffer algorithm often performs needless calculations. Objects are processed in an arbitrary order, so that a color can be computed for a surface point that is later replaced by a closer surface. To alleviate this problem, some graphics packages provide options that allow a user to adjust the depth range for surface testing. This allows distant objects, for example, to be excluded from the depth tests. Using this option, we could even exclude objects that are very close to the projection plane. Hardware implementations of the depth-buffer algorithm are typically an integral component of sophisticated computer-graphics systems.

4 A-Buffer Method

An extension of the depth-buffer ideas is the A-buffer procedure (at the other end of the alphabet from “z-buffer,” where \(z\) represents depth). This depth-buffer extension is an antialiasing, area-averaging, visibility-detection method developed at Lucasfilm Studios for inclusion in the surface-rendering system called...
REYES (an acronym for “Renders Everything You Ever Saw”). The buffer region for this procedure is referred to as the *accumulation buffer*, because it is used to store a variety of surface data, in addition to depth values.

A drawback of the depth-buffer method is that it identifies only one visible surface at each pixel position. In other words, it deals only with opaque surfaces and cannot accumulate color values for more than one surface, as is necessary if transparent surfaces are to be displayed (Figure 8). The A-buffer method expands the depth-buffer algorithm so that each position in the buffer can reference a linked list of surfaces. This allows a pixel color to be computed as a combination of different surface colors for transparency or antialiasing effects.

Each position in the A-buffer has two fields:

- **Depth field**: Stores a real-number value (positive, negative, or zero).
- **Surface data field**: Stores surface data or a pointer.

If the depth field is nonnegative, the number stored at that position is the depth of a surface that overlaps the corresponding pixel area. The surface data field then stores various surface information, such as the surface color for that position and the percent of pixel coverage, as illustrated in Figure 9(a). If the depth field for a position in the A-buffer is negative, this indicates multiple-surface contributions to the pixel color. The color field then stores a pointer to a linked list of surface data, as in Figure 9(b). Surface information in the A-buffer includes

- RGB intensity components
- Opacity parameter (percent of transparency)
- Depth
- Percent of area coverage
- Surface identifier
- Other surface-rendering parameters

The A-buffer visibility-detection scheme can be implemented using methods similar to those in the depth-buffer algorithm. Scan lines are processed to...
determine how much of each surface covers each pixel position across the individual scan lines. Surfaces are subdivided into a polygon mesh and clipped against the pixel boundaries. Using the opacity factors and percent of surface coverage, the rendering algorithms calculate the color for each pixel as an average of the contributions from the overlapping surfaces.

5 Scan-Line Method

This image-space method for identifying visible surfaces computes and compares depth values along the various scan lines for a scene. As each scan line is processed, all polygon surface projections intersecting that line are examined to determine which are visible. Across each scan line, depth calculations are performed to determine which surface is nearest to the view plane at each pixel position. When the visible surface has been determined for a pixel, the surface color for that position is entered into the frame buffer.

Surfaces are processed using the information stored in the polygon tables. The edge table contains coordinate endpoints for each line in the scene, the inverse slope of each line, and pointers into the surface-facet table to identify the surfaces bounded by each line. The surface-facet table contains the plane coefficients, surface material properties, other surface data, and possibly pointers into the edge table. To facilitate the search for surfaces crossing a given scan line, an active list of edges is formed for each scan line as it is processed. The active edge list contains only those edges that cross the current scan line, sorted in order of increasing \( x \). In addition, we define a flag for each surface that is set to “on” or “off” to indicate whether a position along a scan line is inside or outside the surface. Pixel positions across each scan line are processed from left to right. At the left intersection with the surface projection of a convex polygon, the surface flag is turned on; at the right intersection point along the scan line, it is turned off. For a concave polygon, scan-line intersections can be sorted from left to right, with the surface flag set to “on” between each intersection pair.

Figure 10 illustrates the scan-line method for locating visible portions of surfaces for pixel positions along a scan line. The active list for scan line 1 contains information from the edge table for edges AB, BC, EH, and FG. For positions along this scan line between edges AB and BC, only the flag for surface \( S_1 \) is on. Therefore, no depth calculations are necessary, and color values are calculated from the surface properties and lighting conditions for surface \( S_1 \). Similarly, between edges

**Figure 10**
Scan lines crossing the view-plane projection of two surfaces, \( S_1 \) and \( S_2 \). Dashed lines indicate the boundaries of hidden surface sections.
FIGURE 11
Intersecting and cyclically overlapping surfaces that alternately obscure one another.

EH and FG, only the flag for surface $S_2$ is on. No other positions along scan line 1 intersect surfaces, so the color for those pixels is the background color, which could be loaded into the frame buffer as part of the initialization routine.

For scan lines 2 and 3 in Figure 10, the active edge list contains edges AD, EH, BC, and FG. Along scan line 2 from edge AD to edge EH, only the flag for surface $S_1$ is on. But between edges EH and BC, the flags for both surfaces are on. Therefore, a depth calculation is necessary, using the plane coefficients for the two surfaces, when we encounter edge EH. For this example, the depth of surface $S_1$ is assumed to be less than that of $S_2$, so the color values for surface $S_1$ are assigned to the pixels across the scan line until boundary BC is encountered. Then the surface flag for $S_1$ goes off, and the colors for surface $S_2$ are stored up to edge FG. No other depth calculations are necessary, because we assume that surface $S_2$ remains behind $S_1$ once we have determined the depth relationship at edge EH.

We can take advantage of coherence along the scan lines as we pass from one scan line to the next. In Figure 10, scan line 3 has the same active list of edges as scan line 2. No changes have occurred in line intersections, so it is again unnecessary to make depth calculations between edges EH and BC. The two surfaces must be in the same orientation as determined on scan line 2, so the colors for surface $S_1$ can be entered without further depth calculations.

Any number of overlapping polygon surfaces can be processed with this scan-line method. Flags for the surfaces are set to indicate whether a position is inside or outside, and depth calculations are performed only at the edges of overlapping surfaces. This procedure works correctly only if surfaces do not cut through or otherwise cyclically overlap each other (Figure 11). If any kind of cyclic overlap is present in a scene, we can divide the surfaces to eliminate the overlaps. The dashed lines in this figure indicate where planes could be subdivided to form two distinct surfaces, so that the cyclic overlaps are eliminated.

6 Depth-Sorting Method

Using both image-space and object-space operations, the depth-sorting method performs the following basic functions:

1. Surfaces are sorted in order of decreasing depth.
2. Surfaces are scan-converted in order, starting with the surface of greatest depth.
Sorting operations are carried out in both image and object space, and the scan conversion of the polygon surfaces is performed in image space.

This visibility-detection method is often referred to as the painter’s algorithm. In creating an oil painting, an artist first paints the background colors. Next, the most distant objects are added, then the nearer objects, and so forth. At the final step, the foreground is painted on the canvas over the background and the more distant objects. Each color layer covers up the previous layer. Using a similar technique, we first sort surfaces according to their distance from the view plane. The color values for the farthest surface can then be entered into the refresh buffer. Taking each succeeding surface in turn (in decreasing depth order), we “paint” the surface onto the frame buffer over the colors of the previously processed surfaces.

Painting polygon surfaces into the frame buffer according to depth is carried out in several steps. Assuming we are viewing along the \( z \) direction, surfaces are ordered on the first pass according to the smallest \( z \) value on each surface. The surface \( S \) at the end of the list (with the greatest depth) is then compared to the other surfaces in the list to determine whether there are any depth overlaps. If no depth overlaps occur, \( S \) is the most distant surface and it is scan-converted. Figure 12 shows two surfaces that overlap in the \( xy \) plane but have no depth overlap. This process is then repeated for the next surface in the list. So long as no overlaps occur, each surface is processed in depth order until all have been scan-converted. If a depth overlap is detected at any point in the list, we need to make some additional comparisons to determine whether any of the surfaces should be reordered.

We make the following tests for each surface that has a depth overlap with \( S \). If any one of these tests is true, no reordering is necessary for \( S \) and the surface being tested. The tests are listed in order of increasing difficulty:

1. The bounding rectangles (coordinate extents) in the \( xy \) directions for the two surfaces do not overlap.
2. Surface \( S \) is completely behind the overlapping surface relative to the viewing position.
3. The overlapping surface is completely in front of \( S \) relative to the viewing position.
4. The boundary-edge projections of the two surfaces onto the view plane do not overlap.

We perform these tests in the order listed and proceed to the next overlapping surface as soon as we find that one of the tests is true. If all the overlapping surfaces
pass at least one of these tests, then \( S \) is the most distant surface. No reordering is then necessary, therefore, and \( S \) is scan-converted.

Test 1 is performed in two parts. We check for overlap first in the \( x \) direction, then in the \( y \) direction. If there is no surface overlap in either of these directions, the two planes cannot obscure one other. An example of two surfaces that overlap in the \( z \) direction but not in the \( x \) direction is shown in Figure 13.

We can perform tests 2 and 3 using back-front polygon tests. That is, we substitute the coordinates for all vertices of \( S \) into the plane equation for the overlapping surface and check the sign of the result. If the plane equations are set up so that the front of the surface is toward the viewing position, then \( S \) is behind \( S' \) if all vertices of \( S \) are in back of \( S' \) (Figure 14). Similarly, \( S' \) is completely ahead of \( S \) if all vertices of \( S \) are in front of \( S' \). Figure 15 shows an overlapping surface \( S' \) that is completely in front of \( S \), but surface \( S \) is not completely behind \( S' \) (test 2 is not true).

If tests 1 through 3 have all failed, we perform test 4 to determine whether the two surface projections overlap. As demonstrated in Figure 16, two surfaces may or may not intersect even though their coordinate extents overlap.

Should all four tests fail for an overlapping surface \( S' \), we interchange surfaces \( S \) and \( S' \) in the sorted list. An example of two surfaces that would be reordered with this procedure is given in Figure 17. At this point, we still do not know for certain that we have found the farthest surface from the view plane. Figure 18 illustrates a situation in which we would first interchange \( S \) and \( S'' \). However, \( S'' \) obscures part of \( S' \), so we need to interchange \( S'' \) and \( S' \) to get the three surfaces.

**Figure 13**
Two surfaces with depth overlap but no overlap in the \( x \) direction.

**Figure 14**
Surface \( S \) is completely behind the overlapping surface \( S' \).

**Figure 15**
Overlapping surface \( S' \) is completely in front of surface \( S \), but \( S \) is not completely behind \( S' \).

**Figure 16**
Two polygon surfaces with overlapping bounding rectangles in the \( xy \) plane.

**Figure 17**
Surface \( S \) extends to a greater depth, but it obscures surface \( S' \).

**Figure 18**
Three surfaces that have been entered into the sorted surface list in the order \( S', S'', S \) should be reordered as \( S', S'', S \).
into the correct depth order. Therefore, we need to repeat the testing process for each surface that is reordered in the list.

It is possible for the algorithm just outlined to get into an infinite loop if two or more surfaces alternately obscure each other, as in Figure 11. In such situations, the algorithm would continually rearrange the ordering of the overlapping surfaces. To avoid such loops, we can flag any surface that has been reordered to a farther depth position so that it cannot be moved again. If an attempt is made to switch the surface a second time, we divide it into two parts to eliminate the cyclic overlap. The original surface is then replaced by the two new surfaces, and we continue processing as before.

7 BSP-Tree Method

A binary space-partitioning (BSP) tree is an efficient method for determining object visibility by painting surfaces into the frame buffer from back to front, as in the painter’s algorithm. The BSP tree is particularly useful when the view reference point changes, but the objects in a scene are at fixed positions.

Applying a BSP tree to visibility testing involves identifying surfaces that are behind or in front of the partitioning plane at each step of the space subdivision, relative to the viewing direction. Figure 19 illustrates the basic concept in this algorithm. With plane $P_1$, we first partition the space into two sets of objects. One set of objects is in back of plane $P_1$ relative to the viewing direction, and the other set is in front of $P_1$. Because one object is intersected by plane $P_1$, we divide that object into two separate objects, labeled $A$ and $B$. Objects $A$ and $C$ are in front of $P_1$, and objects $B$ and $D$ are behind $P_1$. Because each object list contains more than one object, we partition the space again with plane $P_2$, recursively.

**Figure 19**
A region of space (a) is partitioned with two planes $P_1$ and $P_2$ to form the BSP tree representation shown in (b).
processing the front and back object lists. This process continues until all object lists contain no more than one object. This partitioning can be easily represented using a binary tree such as the one shown in Figure 19(b). In this tree, the objects are represented as terminal nodes, with front objects occupying the left branches and back objects occupying the right branches. The location of an object in the tree exactly represents its position relative to each of the partitioning planes.

For objects described with polygon facets, we often choose the partitioning planes to coincide with polygon-surface planes. The polygon equations are then used to identify back and front polygons, and the tree is constructed with one partitioning plane for each polygon face. Any polygon intersected by a partitioning plane is split into two parts.

When the BSP tree is complete, we interpret the tree relative to the position of our viewpoint, beginning at the root node. If the viewpoint is in front of that partitioning plane, we recursively process the back subtree, then recursively process the front subtree. If the viewpoint is behind the partitioning plane, we reverse this, and process the front subtree followed by the back subtree. Thus, the surfaces are generated for display in the order back to front, so that foreground objects are painted over the background objects. Fast hardware implementations for constructing and processing BSP trees are used in some systems.

8 Area-Subdivision Method

This technique for hidden-surface removal is essentially an image-space method, but object-space operations can be used to accomplish depth ordering of surfaces. The area-subdivision method takes advantage of area coherence in a scene by locating those projection areas that represent part of a single surface. We apply this method by successively dividing the total view-plane area into smaller and smaller rectangles until each rectangular area contains the projection of part of a single visible surface, contains no surface projections, or the area has been reduced to the size of a pixel.

To implement this method, we need to establish tests that can quickly identify the area as part of a single surface or tell us that the area is too complex to analyze easily. Starting with the total view, we apply the tests to determine whether we should subdivide the total area into smaller rectangles. If the tests indicate that the view is sufficiently complex, we subdivide it. Next, we apply the tests to each of the smaller areas, subdividing these if the tests indicate that visibility of a single surface is still uncertain. We continue this process until the subdivisions are easily analyzed as belonging to a single surface or until we have reached the resolution limit. An easy way to do this is to successively divide the area into four equal parts at each step, as shown in Figure 20. This approach is similar to that used in constructing a quadtree. A viewing area with a pixel resolution of $1024 \times 1024$ could be subdivided ten times in this way before a subarea is reduced to the size of a single pixel.

There are four possible relationships that a surface can have with an area of the subdivided view plane. We can describe these relative surface positions using the following classifications (Figure 21).

**Surrounding Surface:** A surface that completely encloses the area.

**Overlapping Surface:** A surface that is partly inside and partly outside the area.

**Inside Surface:** A surface that is completely inside the area.

**Outside Surface:** A surface that is completely outside the area.
The tests for determining surface visibility within a rectangular area can be stated in terms of the four surface classifications illustrated in Figure 21. No further subdivisions of a specified area are needed if one of the following conditions is true.

**Condition 1:** An area has no inside, overlapping, or surrounding surfaces (all surfaces are outside the area).

**Condition 2:** An area has only one inside, overlapping, or surrounding surface.

**Condition 3:** An area has one surrounding surface that obscures all other surfaces within the area boundaries.

Initially, we can compare the coordinate extents of each surface with the area boundaries. This will identify the inside and surrounding surfaces, but overlapping and outside surfaces usually require intersection tests. If a single bounding rectangle intersects the area in some way, additional checks are used to determine whether the surface is surrounding, overlapping, or outside. Once a single inside, overlapping, or surrounding surface has been identified, the surface color values are stored in the frame buffer.

One method for testing condition 3 is to sort the surfaces according to minimum depth from the view plane. For each surrounding surface, we then compute the maximum depth within the area under consideration. If the maximum depth of one of these surrounding surfaces is closer to the view plane than the minimum depth of all other surfaces within the area, condition 3 is satisfied. Figure 22 illustrates this situation.

Another method for testing condition 3 that does not require depth sorting is to use plane equations to calculate depth values at the four vertices of the area for all surrounding, overlapping, and inside surfaces. If all four depths for one of the surrounding surfaces are less than the calculated depths for all other surfaces, condition 3 is satisfied. Then the area can be displayed with the colors for that surrounding surface.

For some situations, the previous two testing methods may fail to identify correctly a surrounding surface that obscures all the other surfaces. Further testing could be carried out to identify the single surface that covers the area, but it is faster to subdivide the area than to continue with more complex testing. Once a surface has been identified as an outside or surrounding surface for an area, it will remain in that category for all subdivisions of the area. Furthermore, we can expect to eliminate some inside and overlapping surfaces as the subdivision process continues, so that the areas become easier to analyze. In the limiting case, when a subdivision the size of a pixel is produced, we simply calculate the depth of each relevant surface at that point and assign the color of the nearest surface to that pixel.

As a variation on the basic subdivision process, we could subdivide areas along surface boundaries instead of dividing them in half. If the surfaces have...
been sorted according to minimum depth, we can use the surface of smallest depth value to subdivide a given area. Figure 23 illustrates this method for subdividing areas. The projection of the boundary of surface $S$ is used to partition the original area into the subdivisions $A_1$ and $A_2$. Surface $S$ is then a surrounding surface for $A_1$, and visibility conditions 2 and 3 can be tested to determine whether further subdividing is necessary. In general, fewer subdivisions are required using this approach, but more processing is needed to subdivide areas and to analyze the relation of surfaces to the subdivision boundaries.

### 9 Octree Methods

When an octree representation is used for the viewing volume, visible-surface identification is accomplished by searching octree nodes in a front-to-back order. In Figure 24, the foreground of a scene is contained in octants 0, 1, 2, and 3. Surfaces in the front of these octants are visible to the viewer. Any surfaces toward the rear of the front octants or in the back octants (4, 5, 6, and 7) may be hidden by the front surfaces.

We can process the octree nodes of Figure 24 in the order 0, 1, 2, 3, 4, 5, 6, 7. This results in a depth-first traversal of the octree, where the nodes for the four front suboctants of octant 0 are visited before the nodes for the four back...
suboctants. The traversal of the octree continues in this order for each octant subdivision.

When a color value is encountered in an octree node, that color is saved in the quadtree only if no values have previously been saved for the same area. In this way, only the front colors are saved. Nodes that have the value “void” are ignored. Any node that is completely obscured is eliminated from further processing, so that its subtrees are not accessed. Figure 25 depicts the octants in a region of space and the corresponding quadrants on the view plane. Contributions to quadrant 0 come from octants 0 and 4. Color values in quadrant 1 are obtained from surfaces in octants 1 and 5, and values in each of the other two quadrants are generated from the pairs of octants aligned with each of these quadrants.

Effective octree visibility testing is carried out with recursive processing of octree nodes and the creation of a quadtree representation for the visible surfaces. In most cases, both a front and a back octant must be considered in determining the correct color values for a quadrant. But if the front octant is homogeneously filled with some color, we do not process the back octant. For heterogeneous regions, a recursive procedure is called, passing as new arguments the child of the heterogeneous octant and a newly created quadtree node. If the front is empty, it is necessary only to process the child of the rear octant. Otherwise, two recursive calls are made: one for the rear octant and one for the front octant.

Different views of objects represented as octrees can be obtained by applying transformations to the octree representation that reorient the object according to the view selected. Octants can then be renumbered so that the octree representation is always organized with octants 0, 1, 2, and 3 as the front face.

10 Ray-Casting Method

If we consider the line of sight from a pixel position on the view plane through a scene, as in Figure 26, we can determine which objects in the scene (if any) intersect this line. After calculating all ray-surface intersections, we identify the visible surface as the one whose intersection point is closest to the pixel. This visibility-detection scheme uses ray casting procedures. Ray casting, as a visibility-detection tool, is based on geometricoptics methods, which trace the paths of light rays. Because there are an infinite number of light rays in a scene and we are interested only in those rays that pass through pixel positions, we can trace
the light-ray paths backward from the pixels through the scene. The ray-casting approach is an effective visibility-detection method for scenes with curved surfaces, particularly spheres.

We can think of ray casting as a variation on the depth-buffer method (Section 3). In the depth-buffer algorithm, we process surfaces one at a time and calculate depth values for all projection points over the surface. The calculated surface depths are then compared to previously stored depths to determine visible surfaces at each pixel. In ray casting, we process pixels one at a time and calculate depths for all surfaces along the projection path to that pixel.

Ray casting is a special case of ray-tracing algorithms that trace multiple ray paths to pick up global reflection and refraction contributions from multiple objects in a scene. With ray casting, we only follow a ray out from each pixel to the nearest object. Efficient ray-surface intersection calculations have been developed for common objects, particularly spheres.

11 Comparison of Visibility-Detection Methods

The effectiveness of a visible-surface detection method depends on the characteristics of a particular application. If the surfaces in a scene are widely distributed along the viewing direction so that there is very little depth overlap, a depth-sorting or BSP-tree method is often most efficient. When there are few overlaps of the surface projections on the view plane, a scan-line or area-subdivision approach is a fast way to locate visible surfaces.

As a general rule, either the depth-sorting algorithm or the BSP-tree method is a highly effective approach for scenes with only a few surfaces. This is because these scenes usually have few surfaces that overlap in depth. The scan-line method also performs well when a scene contains a small number of surfaces. We can use the scan-line, depth-sorting, or BSP-tree method to identify visible surfaces effectively for scenes with up to several thousand polygon surfaces. With scenes that contain more than a few thousand surfaces, the depth-buffer method or octree approach performs best. The depth-buffer method has a nearly constant processing time, independent of the number of surfaces in a scene. This is because the size of the surface areas decreases as the number of surfaces in the scene increases. Therefore, the depth-buffer method exhibits relatively low performance with simple scenes and relatively high performance with complex scenes. BSP trees are useful when multiple views are to be generated using different view reference points. If a scene contains curved-surface representations, we can use octree or ray-casting methods to identify visible parts of the scene.

When octree representations are used in a system, the visibility-detection process is fast and simple. Only integer additions and subtractions are used in the process, and there is no need to perform sorting or intersection calculations. Another advantage of octrees is that they store more than just the surface geometry. The entire solid region of an object is available for display, which makes the octree representation useful for obtaining cross-sectional slices of three-dimensional objects.

It is possible to combine and implement the different visible-surface detection methods in various ways. In addition, visibility-detection algorithms are often implemented in hardware, and special systems utilizing parallel processing are employed to increase the efficiency of these methods. Special hardware systems are used when processing speed is an especially important consideration, as in the generation of animated views for flight simulators.
12 Curved Surfaces

Effective methods for determining the visibility of objects with curved surfaces include ray casting and octree methods. With ray casting, we calculate ray-surface intersections and locate the smallest intersection distance along the pixel ray. With octrees, we simply search the nodes from front to back to locate the surface color values. Once an octree representation has been established from the input definition of the objects, all visible surfaces are identified with the same processing procedures. No special considerations need be given to different kinds of surfaces, curved or otherwise.

A curved surface can also be approximated as a polygon mesh, and we can then use one of the visible-surface identification methods previously discussed. But for some objects, such as spheres, it could be more efficient as well as more accurate to use ray casting and the equations describing the curved surface.

Curved-Surface Representations

We can represent a surface with an implicit equation of the form \( f(x, y, z) = 0 \) or with a parametric representation. Spline surfaces, for example, are normally described with parametric equations. In some cases, it is useful to obtain an explicit surface equation, such as with a height function over an \( xy \) ground plane:

\[
z = f(x, y)
\]

Many objects of interest, such as spheres, ellipsoids, cylinders, and cones, have quadratic representations. These surfaces are commonly used to model molecular structures, roller bearings, rings, and shafts.

Scan-line and ray-casting algorithms often involve numerical approximation techniques to solve the surface equation at the intersection point with a scan line or with a pixel ray. Various techniques, including parallel calculations and fast hardware implementations, have been developed for solving the curved-surface intersection equations for commonly used objects.

Surface Contour Plots

For many applications in mathematics, physical sciences, engineering, and other fields, it is useful to display a surface function with a set of contour lines that show the surface shape. The surface may be described with an equation or with data tables, such as topographic data on elevations or population density. With an explicit functional representation, we can plot the visible surface contour lines and eliminate those contour sections that are hidden by the visible parts of the surface.

To obtain an \( xy \) plot of a functional surface, we can write the surface representation in the form

\[
y = f(x, z)
\]

A curve in the \( xy \) plane can then be plotted for values of \( z \) within some selected range, using a specified interval \( \Delta z \). Starting with the largest value of \( z \), we plot the curves from “front” to “back” and eliminate hidden sections. We draw the curve sections on the screen by mapping an \( xy \) range for the function into an \( xy \) pixel screen range. Then, unit steps are taken in \( x \) and the corresponding \( y \) value for each \( x \) value is determined from Eq. 8 for a given value of \( z \).

One way to identify the visible curve sections on the surface is to maintain a list of \( y_{\text{min}} \) and \( y_{\text{max}} \) values previously calculated for the pixel \( x \) coordinates on the screen. As we step from one pixel \( x \) position to the next, we check the calculated
Visible-Surface Detection Methods

Scenes usually do not contain isolated line sections, unless we are displaying a graph, diagram, or network layout. But often we want to view a three-dimensional scene in an outline form to obtain a quick display of the object features. The fastest way to generate a wire-frame view of a scene is to display all object edges. However, it may be difficult to determine the front and back features of the objects in such a display. One solution to this problem is to apply depth cueing, so that the displayed intensity of a line is a function of its distance from the viewer. Alternatively, we can apply visibility tests, so that hidden line sections can be either eliminated or displayed differently from the visible edges. Procedures for determining visibility of object edges are referred to as wire-frame visibility methods. They are also called visible-line detection methods or hidden-line detection methods. In addition, some of the visible-surface methods discussed in preceding sections can be used to test for edge visibility.

Wire-Frame Surface-Visibility Algorithms

A direct approach to identifying visible line sections is to compare edge positions with the positions of the surfaces in a scene. This process involves the same methods used in line-clipping algorithms. That is, we test the position of line endpoints with respect to the boundaries of a specified area, but, for visibility testing, we also need to compare edge and surface depth values. When the projected edge endpoints of a line segment are both within the projected area of a surface, we compare the depth of the endpoints to the surface depth at those $(x, y)$ positions. If both endpoints are behind the surface, we have a hidden edge. If both endpoints are in front of the surface, the edge is visible with respect to that surface. Otherwise, we must calculate intersection positions and determine the depth values at those intersection points. If the edge has greater depth than the surface at the perimeter intersections, part of the edge is hidden by the surface, as in Figure 27(a). Another possibility is that an edge has greater depth at one boundary intersection and less depth than the surface at the other boundary intersection (assuming surfaces are convex). In that case, we need to determine where the edge penetrates the surface interior, as in Figure 27(b). Once we have identified a hidden section of an edge, we could eliminate it, display it as a dashed line, or display it in some other way to distinguish it from the visible sections.

Some of the visible-surface detection methods are readily adapted to wire-frame visibility testing of object edges. Using a back-face method, we could

\[ y \text{ value against the stored range, } y_{\text{min}} \text{ and } y_{\text{max}}, \text{ for the next pixel. If } y_{\text{min}} \leq y \leq y_{\text{max}}, \text{ that point on the surface is not visible and we do not plot it. But if the calculated } y \text{ value is outside the stored } y \text{ bounds for that pixel, the point is visible. We then plot the point and reset the bounds for that pixel. Similar procedures can be used to project the contour plot onto the } xz \text{ or } yz \text{ plane. } \]

\[ \text{We can apply the same methods to a discrete set of data points by determining isosurface lines. For example, if we have a discrete set of } z \text{ values for an } n_x \times n_y \text{ grid of } xy \text{ values, we can determine the path for a line of constant } z \text{ over the surface using contour plotting methods. Each selected contour line can then be projected onto a view plane and displayed with straight-line segments. Again, lines can be drawn on the display device in a front-to-back depth order, and we eliminate contour sections that pass behind previously drawn (visible) contour lines.} \]
identify all the back surfaces of an object and display only the boundaries for the visible surfaces. With depth sorting, surfaces can be painted into the refresh buffer so that surface interiors are in the background color while boundaries are in the foreground color. By processing the surfaces from back to front, hidden lines are erased by the nearer surfaces. An area-subdivision method can be adapted to hidden-line removal by displaying only the boundaries of visible surfaces. And scan-line methods can be used to display the scan-line intersection positions at the boundaries of visible surfaces.

**Wire-Frame Depth-Cueing Algorithm**

Another method for displaying visibility information is to vary the brightness of objects in a scene as a function of distance from the viewing position. This depth-cueing method is typically applied using the linear function

\[ f_{\text{depth}}(d) = \frac{d_{\text{max}} - d}{d_{\text{max}} - d_{\text{min}}} \]  

where \( d \) is the distance of a point from the viewing position. Values for minimum and maximum depth, \( d_{\text{min}} \) and \( d_{\text{max}} \), can be set to convenient values for a particular application, or the minimum and maximum depths can be set to the normalization depth range: \( d_{\text{min}} = 0.0 \) and \( d_{\text{max}} = 1.0 \). As each pixel position is processed, its color is multiplied by \( f_{\text{depth}}(d) \). Thus, nearer points are displayed with higher intensities, and the points at the maximum depth have an intensity equal to 0.

The depth-cueing function can be implemented with various options. In some graphics libraries, a general atmosphere function is available, which can combine depth cueing with atmospheric effects to simulate smoke or haze, for example. Thus, an object's color could be modified by the depth-cueing function and then combined with the atmosphere color.
14 OpenGL Visibility-Detection Functions

We can apply both back-face removal and the depth-buffer visibility-testing method to our scenes using functions that are provided in the basic library of OpenGL. In addition, we can use OpenGL functions to construct a wire-frame display of a scene with the hidden lines removed, and we can display scenes with depth cueing.

OpenGL Polygon-Culling Functions

Back-face removal is accomplished with the functions

```c
glEnable (GL_CULL_FACE);
glcullFace (mode);
```

where parameter `mode` is assigned the value `GL_BACK`. In fact, we could use this function to remove the front faces instead, or we could even remove both front and back faces. If our viewing position is inside a building, for example, then we want to see only the back faces (the inside of the rooms). In this case, we could either set parameter `mode` to `GL_FRONT`, or we could change the definition of front-facing polygons using the `glFrontFace` function. Then, if the viewing position moves outside the building, we can cull the back faces from the display; and in some applications, we might want to view only other primitives in a scene, such as point sets and individual straight-line segments. So, to eliminate all surfaces in a scene, we set parameter `mode` to the OpenGL symbolic constant `GL_FRONT_AND_BACK`.

By default, parameter `mode` in the `glCullFace` function has the value `GL_BACK`. Therefore, if we activate culling with the `glEnable` function without explicitly invoking function `glCullFace`, the back faces in a scene will be removed. The culling routine is turned off with

```c
glDisable (GL_CULL_FACE);
```

OpenGL Depth-Buffer Functions

To use the OpenGL depth-buffer visibility-detection routines, we first need to modify the GL Utility Toolkit (GLUT) initialization function for the display mode to include a request for the depth buffer, as well as for the refresh buffer. We do this, for example, with the statement

```c
glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB | GLUT_DEPTH);
```

Depth buffer values can then be initialized with

```c
glClear (GL_DEPTH_BUFFER_BIT);
```

Normally, the depth buffer is initialized with the same statement that initializes the refresh buffer to the background color. But we do need to clear the depth buffer each time we want to display a new frame. In OpenGL, depth values are normalized in the range from 0.0 to 1.0, so that the preceding initialization sets all depth-buffer values to the maximum value 1.0 by default.

The OpenGL depth-buffer visibility-detection routines are activated with the following function:

```c
glEnable (GL_DEPTH_TEST);
```
And we deactivate the depth-buffer routines with
\[
glDisable (GL\_DEPTH\_TEST);
\]

We can also apply depth-buffer visibility testing using some other initial value
for the maximum depth, and this initial value is chosen with the OpenGL function:
\[
glClearDepth (\text{maxDepth});
\]

Parameter \text{maxDepth} can be set to any value between 0.0 and 1.0. To load this
initialization value into the depth buffer, we next must invoke the \text{glClear}
(GL\_DEPTH\_BUFFER\_BIT) function. Otherwise, the depth buffer is initialized
with the default value (1.0). Because surface-color calculations and other process-
ing are not performed for objects that are beyond the specified maximum depth,
this function can be used to speed up the depth-buffer routines when a scene
contains many distant objects that are behind the foreground objects.

Projection coordinates in OpenGL are normalized to the range from \(-1.0\) to
1.0, and the depth values between the near and far clipping planes are further
normalized to the range from 0.0 to 1.0. The value 0.0 corresponds to the near
clipping plane (the projection plane), and the value 1.0 corresponds to the far
clipping plane. As an option, we can adjust these normalization values with
\[
glDepthRange (\text{nearNormDepth}, \text{farNormDepth});
\]

By default, \text{nearNormDepth} = 0.0 and \text{farNormDepth} = 1.0. But with the
\text{glDepthRange} function, we can set these two parameters to any values within
the range from 0.0 to 1.0, including \text{nearNormDepth} > \text{farNormDepth}. Using the
\text{glDepthRange} function, we can restrict the depth-buffer testing to any region of
the view volume, and we can even reverse the positions of the near and far planes.

Another option available in OpenGL is the test condition that is to be used for
the depth-buffer routines. We specify a test condition with the following function:
\[
glDepthFunc (\text{testCondition});
\]

Parameter \text{testCondition} can be assigned any one of the following
eight symbolic constants: \text{GL\_LESS}, \text{GL\_GREATER}, \text{GL\_EQUAL}, \text{GL\_NOTEQUAL},
\text{GL\_LEQUAL}, \text{GL\_GEQUAL}, \text{GL\_NEVER} (no points are processed), and \text{GL\_ALWAYS}
(all points are processed). These different tests can be useful in various applica-
tions to reduce calculations in depth-buffer processing. The default value for
parameter \text{testCondition} is \text{GL\_LESS}, so that a depth value is processed if it
has a value that is less than the current value in the depth buffer for that pixel
position.

We can also set the status of the depth buffer so that it is in a read-only state
or in a read-write state. This is accomplished with
\[
glDepthMask (\text{writeStatus});
\]

When \text{writeStatus} = \text{GL\_TRUE} (the default value), we can both read from
and write to the depth buffer. With \text{writeStatus} = \text{GL\_FALSE}, the write mode
for the depth buffer is disabled and we can retrieve values only for comparison
in depth testing. This feature is useful when we want to use the same complica-
ted background with displays of different foreground objects. After storing the
background in the depth buffer, we disable the write mode and process the fore-
ground. This allows us to generate a series of frames with different foreground
objects or with one object in different positions for an animation sequence. Thus,
only the depth values for the background are saved. Another application of the 
glDepthMask function is in displaying transparency effects. In this case, we want 
to save only the depths of opaque objects for visibility testing, not the depths of 
the transparent-surface positions. So the write mode for the depth buffer is turned 
off when a transparent surface is processed. Similar commands are available for setting the write status for the other buffers (color, index, and stencil).

**OpenGL Wire-Frame Surface-Visibility Methods**

A wire-frame display of a standard graphics object can be obtained in OpenGL 
by requesting that only its edges are to be generated. We do this by setting the 
polygon-mode function as, for example:

```c
glPolygonMode (GL_FRONT_AND_BACK, GL_LINE);
```

But this displays both visible and hidden edges.

To eliminate the hidden lines in a wire-frame display, we can employ the 
depth-offset method. That is, we first specify the wireframe version of the object 
using the foreground color, then we specify an interior fill version using a depth 
offset and the background color for the interior fill. The depth offset ensures that 
the background-color fill will not interfere with the display of the visible edges. As 
an example, the following code segment generates a wire-frame display of an 
object using a white foreground color and a black background color:

```c
glEnable (GL_DEPTH_TEST);
glPolygonMode (GL_FRONT_AND_BACK, GL_LINE);
g1Color3f (1.0, 1.0, 1.0);
/* Invoke the object-description routine. */

// The wireframe version with a white foreground and black background

// The interior fill overlay version with a depth offset

// The interior fill version using a depth offset and black background color

// Invoke the object-description routine again. */

g1Disable (GL_POLYGON_OFFSET_FILL);
```

**OpenGL Depth-Cueing Function**

We can vary the brightness of an object as a function of its distance from the 
viewing position with

```c
glEnable (GL_FOG);
g1Fogi (GL_FOG_MODE, GL_ LINEAR);
```

This applies the linear depth function in Eq. 9 to object colors using \( d_{\text{min}} = 0.0 \) 
and \( d_{\text{max}} = 1.0 \). But we can set different values for \( d_{\text{min}} \) and \( d_{\text{max}} \) with the following 
function calls:

```c
gfog (GL_FOG_START, minDepth);
g1Fogf (GL_FOG_END, maxDepth);
```
In these two functions, parameters minDepth and maxDepth are assigned floating-point values, although integer values can be used if we change the function suffix to i.

In addition, we can use the glFog function to set an atmosphere color that is to be combined with the color of an object after applying the linear depth-cueing function.

## 15 Summary

The simplest visibility test is the back-face detection algorithm, which is fast and effective as an initial screening to eliminate many polygons from further visibility tests. For a single convex polyhedron, back-face detection eliminates all hidden surfaces, but, in general, back-face detection cannot completely identify all hidden surfaces.

A commonly used method for identifying all visible surfaces in a scene is the depth-buffer algorithm. When applied to standard graphics objects, this procedure is highly efficient, but it does have extra storage requirements. Two buffers are needed: one to store pixel colors and one to store the depth values for the pixel positions. Fast, incremental, scan-line methods are used to process each polygon in a scene to calculate surface depths. As each surface is processed, the two buffers are updated. An extension of the depth-buffer approach is the A-buffer, which provides additional information for displaying antialiased and transparent surfaces.

Several other visibility-detection methods have been devised. The scan-line method processes all surfaces at once for each scan line. With the depth-sorting method (painter’s algorithm), objects are “painted” into the refresh buffer according to their distances from the viewing position. Subdivision schemes for identifying visible parts of a scene include the BSP-tree method, area subdivision, and octree representations. Visible surfaces can also be detected using ray-casting methods, which project lines from the pixel plane into a scene to determine object intersection positions along these projected lines. Ray-casting methods are an integral part of ray-tracing algorithms, which allow scenes to be displayed with global-illumination effects.

Visibility-detection methods are also used in displaying three-dimensional line drawings. With curved surfaces, we can display contour plots. For wire-frame displays of polyhedrons, we search for the various edge sections of the surfaces in a scene that are visible from the viewing position.

We can implement any visibility-detection scheme in an application program by creating our own routines, but graphics libraries commonly provide functions only for back-face removal and the depth-buffer method. In high-end computer-graphics systems, the depth-buffer routines are hardware-implemented.

Functions for polygon culling and for depth-buffer visibility determinations are available in the OpenGL core library. With the polygon-culling routines, we can remove the back faces of standard graphics objects, their front faces, or both. With the depth-buffer routines, we can set the range for the depth tests and the type of depth testing that is to be performed. Wire-frame displays are obtained using the OpenGL polygon-mode and polygon-offset operations. And we can also generate OpenGL scenes using depth-cueing effects. In Table 1, we summarize the OpenGL functions for visibility testing.
TABLE 1
Summary of OpenGL Visibility-Detection Functions

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glCullFace</td>
<td>Specifies front or back planes of polygons for culling operations when activated with glEnable (GL_CULL_FACE).</td>
</tr>
<tr>
<td>glutInitDisplayMode</td>
<td>Specifies depth-buffer operations using argument GLUT_DEPTH.</td>
</tr>
<tr>
<td>glClear (GL_DEPTH_BUFFER_BIT)</td>
<td>Initializes depth-buffer values to the default (1.0) or a value specified by the glClearDepth function.</td>
</tr>
<tr>
<td>glClearDepth</td>
<td>Specifies an initial depth-buffer value.</td>
</tr>
<tr>
<td>glEnable (GL_DEPTH_TEST)</td>
<td>Activates depth-testing operations.</td>
</tr>
<tr>
<td>glDepthRange</td>
<td>Specifies a range for normalizing depth values.</td>
</tr>
<tr>
<td>glDepthFunc</td>
<td>Specifies a depth-testing condition.</td>
</tr>
<tr>
<td>glDepthMask</td>
<td>Sets write status for the depth buffer.</td>
</tr>
<tr>
<td>glPolygonOffset</td>
<td>Specifies an offset to eliminate hidden lines in a wire-frame display when a background fill color is applied.</td>
</tr>
<tr>
<td>glFog</td>
<td>Specifies linear depth-cueing operations and values for minimum and maximum depth in the depth-cueing calculations.</td>
</tr>
</tbody>
</table>

REFERENCES


EXERCISES
1 Set up a back-face detection procedure that will identify all the visible faces of any input convex polyhedron that has different-colored surfaces. The polyhedron is to be defined in a right-handed viewing system, and the viewing direction is specified as user input.

2 Implement the procedure in the preceding exercise using an orthographic parallel projection to view visible faces of the input convex polyhedron. Assume that all parts of the object are in front of the view plane.
Implement the procedure in Exercise 1 using a perspective projection to view visible faces of the input convex polyhedron. Assume that all parts of the object are in front of the view plane.

Write a program to produce an animation of a convex polyhedron. The object is to be rotated incrementally about an axis that passes through the object and is parallel to the view plane. Assume that the object lies completely in front of the view plane. Use an orthographic parallel projection to map the views successively onto the view plane.

Modify the program in the preceding exercise to allow the user to switch between an orthographic parallel projection and a perspective projection using keyboard input.

Write a routine to implement the depth-buffer method for the display of the visible surfaces of any input polyhedron. The array for the depth-buffer can be set to any convenient size on your system, such as 500 × 500. How can the storage requirements for the depth buffer be determined from the definition of the objects to be displayed?

Modify the procedure in the preceding exercise to display the visible surfaces in a scene containing any number of polyhedrons. Set up efficient methods for storing and processing the various objects in the scene.

Write a program using the procedure in the previous exercise that takes as input a set of polyhedrons contained within a (conceptual) sphere of a given radius centered at the origin. Each time a certain key is pressed, the program should generate a new random camera position outside of the sphere and a random look-at point somewhere inside the sphere. The view up vector should always be the positive y unit vector. The program should then display the visible surfaces of the objects in the scene from that viewpoint.

Develop a program to implement the scan-line algorithm for displaying the visible surfaces of a given polyhedron. Use polygon tables to store the definition of the object, and use coherence techniques to evaluate points along and between scan lines.

Write a program to implement the scan-line algorithm for a scene containing several polyhedrons. Use polygon tables to store the definition of the object, and use coherence techniques to evaluate points along and between scan lines.

Set up a program to display the visible surfaces of a convex polyhedron using the painter’s algorithm. That is, surfaces are to be sorted on depth and painted on the screen from back to front.

Write a program that uses the depth-sorting method to display the visible surfaces of any given object with plane faces.

Develop a depth-sorting program to display the visible surfaces in a scene containing several polyhedrons.

Write a program to display the visible surfaces of a convex polyhedron using the BSP-tree method.

Give examples of situations where the two methods discussed for condition 3 in the area-subdivision algorithm will fail to identify correctly a surrounding surface that obscures all other surfaces.

Develop an algorithm that would test a given plane surface against a rectangular area to decide whether it is a surrounding, overlapping, inside, or outside surface.

Develop an algorithm for generating a quadtree representation for the visible surfaces of an object by applying the area-subdivision tests to determine the values of the quadtree elements.

Set up an algorithm to store a quadtree representation of an object in a frame buffer.

Set up a procedure to display the visible surfaces of an object that is described with an octree representation.

Use the procedure developed in the previous exercise to write a program that displays the visible surfaces of a set of objects represented as octree structures. The viewing parameters should be taken in as input.

Devises an algorithm for viewing a single sphere using the ray-casting method.

Discuss how antialiasing methods can be incorporated into the various hidden-surface elimination algorithms.
26 Write a routine to produce a surface contour plot for a given surface function \( f(x, y) \).

27 Develop an algorithm for detecting visible line sections in a scene by comparing each line in the scene to each polygon surface facet.

28 Discuss how wire-frame displays might be generated with the various visible-surface detection methods discussed in this chapter.

29 Set up a procedure for generating a wire-frame display of a polyhedron with the hidden edges of the object shown as dashed lines.

30 Write a program using the procedure developed in the previous exercise that takes a set of polyhedrons contained within a (conceptual) sphere of a given radius centered at the origin as input and displays them as wireframe objects with the hidden edges of each object shown as dashed lines. Each time a key is pressed, the program should generate a new random camera position outside of the sphere and a random look-at point somewhere inside the sphere. The view up vector should always be the positive y unit vector.

31 Write a program to display a polyhedron with selected faces removed, using the OpenGL polygon-culling functions. Each face of the polygon is to be given a different color, and a face is to be selected for removal with user input. Also, a viewing position and other viewing parameters are to be specified as input values.

32 Modify the program in the preceding exercise to view the polyhedron from any position, using the depth-buffer routines instead of the polygon-culling routines.

33 Modify the program in the preceding exercise so that the depth range and the depth test condition can also be specified as user input.

34 Generate a wire-frame display of a polyhedron using the `glPolygonMode` and `glPolygonOffset` functions as discussed in Section 14.

35 Modify the program of the preceding exercise to display the polyhedron using the depth-cueing function `glFogi`.

36 Modify the program of the preceding exercise to display several polyhedrons that are distributed in depth. The depth-cueing range is to be set with user input.

37 Modify the program in the previous exercise to allow the change the camera position by moving it around the surface of a sphere whose radius is defined as the distance from the camera position to the look-at point, which is assumed to be a point within the coordinate extents of the set of objects in the scene. The distance from the camera to the look-at point is assumed to be large enough to make all objects lie in front of the view plane for any camera position on the sphere.

**IN MORE DEPTH**

1. Choose a visible surface algorithm in this chapter based on the properties of your application and the strengths and weaknesses of each of the algorithms in terms of their computational complexity. Implement the algorithm and use it to render the visible surfaces of the objects in your scene.

2. Compare the rendering times for your scene with and without visible surface detection using the algorithm that you developed in the previous exercise. Then, do the same using the built-in back-face culling routines provided in OpenGL. Is there any improvement over the built-in routines that you obtain by tailoring the detection algorithm to your specific application? Discuss any further improvements you could implement in the algorithm or modifications that you could make to the object representations to increase rendering performance.
This page intentionally left blank
Realistic displays of a scene are obtained by generating perspective projections of objects and applying natural lighting effects to the visible surfaces. An illumination model, also called a lighting model (and sometimes referred to as a shading model), is used to calculate the color of an illuminated position on the surface of an object. A surface-rendering method uses the color calculations from an illumination model to determine the pixel colors for all projected positions in a scene. The illumination model can be applied to every projection position, or the surface rendering can be accomplished by interpolating colors on the surfaces using a small set of illumination-model calculations. Scan-line, image-space algorithms typically use interpolation schemes. Sometimes, a surface-rendering procedure is called a shading method that calculates surface colors using a shading model, but this can lead to some confusion between the two terms. To avoid possible misinterpretations due to the use of similar terminology, we refer to the model for calculating the light intensity at a single surface point.
Illumination Models and Surface-Rendering Methods

as an *illumination model* or a *lighting model*, and we use the term *surface rendering* to mean a procedure for applying a lighting model to obtain pixel colors for all projected surface positions.

Among other things, photorealism in computer graphics involves two elements: accurate representations of surface properties and good physical descriptions of the lighting effects in a scene. These surface lighting effects include light reflections, transparency, surface texture, and shadows.

In general, modeling the lighting effects that we see on an object is a complex process, involving principles of both physics and psychology. Fundamentally, lighting effects are described with models that consider the interaction of electromagnetic energy with the object surfaces in a scene. Once light reaches our eyes, it triggers perception processes that determine what we actually “see.” Physical illumination models involve a number of factors, such as material properties, object position relative to light sources and other objects, and the features of the light sources. Objects can be composed of opaque materials, or they can be more or less transparent. In addition, they can have shiny or dull surfaces, and they can have a variety of surface-texture patterns. Light sources of varying shapes, colors, and positions can be used to provide the illumination for a scene.

Given the parameters for the optical properties of surfaces, the relative positions of the surfaces in a scene, the color and positions of the light sources, the characteristics of the light sources, and the position and orientation of the viewing plane, illumination models calculate the light intensity projected from a particular surface position in a specified viewing direction.

Illumination models in computer graphics are often approximations of the physical laws that describe surface-lighting effects. To reduce computations, most packages use empirical models based on simplified photometric calculations. In the following sections, we take a look at the basic lighting models often used in computer-graphics systems, and we explore the various surface-rendering algorithms for applying the lighting models to obtain effective displays of natural scenes.

## 1 Light Sources

Any object that is emitting radiant energy is a *light source* that contributes to the lighting effects for other objects in a scene. We can model light sources with a variety of shapes and characteristics, and most emitters serve only as a source of illumination for a scene. In some applications, however, we may want to create an object that is both a light source and a light reflector. For example, a plastic globe surrounding a light bulb both emits and reflects light from the surface of the globe. We could also model the globe as a semitransparent surface around a light source. However, for some objects, such as a large fluorescent light panel, it might be more convenient to describe the surface simply as a combination emitter and reflector.

A light source can be defined with a number of properties. We can specify its position, the color of the emitted light, the emission direction, and its shape. If the source is also to be a light-reflecting surface, we need to give its reflectivity properties. In addition, we could set up a light source that emits different colors in different directions. For example, we could define a light source that emits a red light on one side and a green light on the other side.
In most applications, and particularly for real-time graphics displays, a simple light-source model is used to avoid excessive computations. We assign light-emitting properties using a single value for each of the red, green, and blue (RGB) color components, which we can describe as the amount, or the “intensity,” of that color component.

**Point Light Sources**

The simplest model for an object that is emitting radiant energy is a **point light source** with a single color, specified with three RGB components. We define a point source for a scene by giving its position and the color of the emitted light. As shown in Figure 1, light rays are generated along radially diverging paths from the single-color source position. This light-source model is a reasonable approximation for sources whose dimensions are small compared to the size of objects in the scene. We can also simulate larger sources as point emitters if they are not too close to a scene. We use the position of a point source in an illumination model to determine which objects in the scene are illuminated by that source and to calculate the light direction to a selected object surface position.

**Infinitely Distant Light Sources**

A large light source, such as the sun, that is very far from a scene can also be approximated as a point emitter, but there is little variation in its directional effects. In contrast to a light source in the middle of a scene, which illuminates objects on all sides of the source, a remote source illuminates the scene from only one direction. The light path from a distant light source to any position in the scene is nearly constant, as illustrated in Figure 2.

We can simulate an infinitely distant light source by assigning it a color value and a fixed direction for the light rays emanating from the source. The vector for the emission direction and the light-source color are needed in the illumination calculations, but not the position of the source.

**Radial Intensity Attenuation**

As radiant energy from a light source travels outwards through space, its amplitude at any distance $d$ from the source is attenuated by the factor $1/d^2$. This means that a surface close to the light source receives a higher incident light...
intensity from that source than a more distant surface. Therefore, to produce realistic lighting effects, we should take this intensity attenuation into account. Otherwise, all surfaces are illuminated with the same intensity from a light source, and undesirable display effects can result. For example, if two surfaces with the same optical parameters project to overlapping positions, they would be indistinguishable from one another. Thus, regardless of their relative distances from the light source, the two surfaces would appear to be one surface.

In practice, however, using an attenuation factor of $1/d^2$ with a point source does not always produce realistic pictures. The factor $1/d^2$ tends to produce too much intensity variation for objects that are close to the light source, and very little variation when $d$ is large. This is because actual light sources are not infinitesimal points, and illuminating a scene with point emitters is only a simple approximation of true lighting effects. To generate more realistic displays using point sources, we can attenuate light intensities with an inverse quadratic function of $d$ that includes a linear term:

$$f_{\text{radatten}}(d) = \frac{1}{a_0 + a_1 d + a_2 d^2}$$

The numerical values for the coefficients, $a_0$, $a_1$, and $a_2$, can then be adjusted to produce optimal attenuation effects. For instance, we can assign a large value to $a_0$ when $d$ is very small to prevent $f_{\text{radatten}}(d)$ from becoming too large. As an additional option, often available in graphics packages, a different set of values for the attenuation coefficients could be assigned to each point light source in the scene.

We cannot apply intensity-attenuation calculation 1 to a point source at “infinity,” because the distance to the light source is indeterminate. Also, all points in the scene are at a nearly equal distance from a far-off source. To accommodate both remote and local light sources, we can express the intensity-attenuation function as

$$f_{l,\text{radatten}} = \begin{cases} 
1.0, & \text{if source is at infinity} \\
\frac{1}{a_0 + a_1 d + a_2 d^2}, & \text{if source is local}
\end{cases}$$

Directional Light Sources and Spotlight Effects

A local light source can be modified easily to produce a directional, or spotlight, beam of light. If an object is outside the directional limits of the light source, we exclude it from illumination by that source. One way to set up a directional light source is to assign it a vector direction and an angular limit $\theta_l$ measured from that vector direction, in addition to its position and color. This defines a conical region of space with the light-source vector direction along the axis of the cone (Figure 3). A multicolor point light source could be modeled in this way using multiple direction vectors and a different emission color for each direction.

We can denote $V_{\text{light}}$ as the unit vector in the light-source direction and $V_{\text{obj}}$ as the unit vector in the direction from the light position to an object position. Then

$$V_{\text{obj}} \cdot V_{\text{light}} = \cos \alpha$$

where angle $\alpha$ is the angular distance of the object from the light direction vector. If we restrict the angular extent of any light cone so that $0^\circ < \theta_l \leq 90^\circ$, then the object is within the spotlight if $\cos \alpha \geq \cos \theta_l$, as shown in Figure 4. If $V_{\text{obj}} \cdot V_{\text{light}} < \cos \theta_l$, however, the object is outside the light cone.
Angular Intensity Attenuation

For a directional light source, we can attenuate the light intensity angularly about the source as well as radially out from the point-source position. This allows us to simulate a cone of light that is most intense along the axis of the cone, with the intensity decreasing as we move farther from the cone axis. A commonly used angular intensity-attenuation function for a directional light source is

\[ f_{\text{angatten}}(\phi) = \cos^{a_l} \phi, \quad 0^\circ \leq \phi \leq \theta \]  

where the attenuation exponent \( a_l \) is assigned some positive value and angle \( \phi \) is measured from the cone axis. Along the cone axis, \( \phi = 0^\circ \) and \( f_{\text{angatten}}(\phi) = 1.0 \). The greater the value for the attenuation exponent \( a_l \), the smaller the value of the angular intensity-attenuation function for a given value of angle \( \phi > 0^\circ \).
There are several special cases to consider in the implementation of the angular-attenuation function. There is no angular attenuation if the light source is not directional (not a spotlight). Also, an object is not illuminated by the light source if it is anywhere outside the cone of the spotlight. To determine the angular attenuation factor along a line from the light position to a surface position in a scene, we can compute the cosine of the direction angle from the cone axis using the dot product calculation in Equation 3. We designate $\mathbf{V}_{\text{light}}$ as the unit vector in the light-source direction (along the cone axis) and $\mathbf{V}_{\text{obj}}$ as the unit vector in the direction from the light source to an object position. Using these two unit vectors and assuming that $0^\circ < \theta_l \leq 90^\circ$, we can express the general equation for angular attenuation as

$$f_{l,\text{angatten}} = \begin{cases} 
1.0, & \text{if source is not a spotlight} \\
0.0, & \text{if } \mathbf{V}_{\text{obj}} \cdot \mathbf{V}_{\text{light}} = \cos \alpha < \cos \theta_l \\
(\mathbf{V}_{\text{obj}} \cdot \mathbf{V}_{\text{light}})^{\theta_l}, & \text{otherwise}
\end{cases} \quad (5)$$

**Extended Light Sources and the Warn Model**

When we want to include a large light source at a position close to the objects in a scene, such as the long neon lamp in Figure 5, we can approximate it as a light-emitting surface. One way to do this is to model the light surface as a grid of directional point emitters. We can set the direction for the point sources so that objects behind the light-emitting surface are not illuminated. We could also include other controls to restrict the direction of the emitted light near the edges of the source.

The Warn model provides a method for producing studio lighting effects using sets of point emitters with various parameters to simulate the barn doors, flaps, and spotlighting controls employed by photographers. Spotlighting is achieved with the cone of light discussed earlier, and the flaps and barn doors provide additional directional control. For instance, two flaps can be set up for each of the $x$, $y$, and $z$ directions to further restrict the path of the emitted light rays. This light-source simulation is implemented in some graphics packages.

## 2 Surface Lighting Effects

An illumination model computes the lighting effects for a surface using the various optical properties that have been assigned to that surface. These properties include degree of transparency, color reflectance coefficients, and various surface-texture parameters.

When light is incident on an opaque surface, part of it is reflected and part is absorbed. The amount of incident light reflected by the surface depends on...
the type of material. Shiny materials reflect more of the incident light, and dull surfaces absorb more of the incident light. For a transparent surface, some of the incident light is also transmitted through the material.

Surfaces that are rough or grainy tend to scatter the reflected light in all directions. This scattered light is called diffuse reflection. A very rough, matte surface produces primarily diffuse reflections, so the surface appears equally bright from any viewing angle. Figure 6 illustrates diffuse light scattering from a surface. What we call the color of an object is the color of the diffuse reflection when the object is illuminated with white light, which is composed of a combination of all colors. A blue object, for example, reflects the blue component of the white light and absorbs all the other color components. If the blue object is viewed under a red light, it appears black because all the incident light is absorbed.

In addition to diffuse light scattering, some of the reflected light is concentrated into a highlight, or bright spot, called specular reflection. This highlighting effect is more pronounced on shiny surfaces than on dull surfaces, and we can see the specular reflection when we look at an illuminated shiny surface, such as polished metal, an apple, or a person’s forehead, only when we view the surface from a particular direction. A representation of specular reflection is shown in Figure 7.

Another factor that must be considered in an illumination model is the background light or ambient light in a scene. A surface that is not directly exposed to a light source may still be visible due to the reflected light from nearby objects that are illuminated. Thus, the ambient light for a scene is the illumination effect produced by the reflected light from the various surfaces in the scene. Figure 8 illustrates this background lighting effect. The total reflected light from a surface is the sum of the contributions from light sources and from the light reflected by other illuminated objects.

3 Basic Illumination Models

Accurate surface lighting models compute the results of interactions between incident radiant energy and the material composition of an object. To simplify the surface-illumination calculations, we can use approximate representations for the physical processes that produce the lighting effects discussed in the previous section. The empirical model described in this section produces reasonably good results, and it is implemented in most graphics systems.

Light-emitting objects in a basic illumination model are generally limited to point sources. However, many graphics packages provide additional functions for dealing with directional lighting (spotlights) and extended light sources.

Ambient Light

In our basic illumination model, we can incorporate background lighting by setting a general brightness level for a scene. This produces a uniform ambient lighting that is the same for all objects, and it approximates the global diffuse reflections from the various illuminated surfaces.

Assuming that we are describing only monochromatic lighting effects, such as shades of gray, we designate the level for the ambient light in a scene with an intensity parameter $I_a$. Each surface in the scene is then illuminated with this background light. Reflections produced by ambient-light illumination are simply a form of diffuse reflection, and they are independent of the viewing direction and the spatial orientation of a surface. However, the amount of the incident ambient
light that is reflected depends on surface optical properties, which determine how much of the incident energy is reflected and how much is absorbed.

**Diffuse Reflection**

We can model diffuse reflections from a surface by assuming that the incident light is scattered with equal intensity in all directions, independent of the viewing position. Such surfaces are called **ideal diffuse reflectors**. They are also referred to as **Lambertian reflectors**, because the reflected radiant light energy from any point on the surface is calculated with **Lambert’s cosine law**. This law states that the amount of radiant energy coming from any small surface area \( dA \) in a direction \( \phi_N \) relative to the surface normal is proportional to \( \cos \phi_N \) (Figure 9). The intensity of light in this direction can be computed as the ratio of the magnitude of the radiant energy per unit time divided by the projection of the surface area in the radiation direction:

\[
\text{Intensity} = \frac{\text{radiant energy per unit time}}{\text{projected area}} \propto \frac{\cos \phi_N}{dA \cos \phi_N} = \text{constant}
\]

Thus, for Lambertian reflection, the intensity of light is the same over all viewing directions.

Assuming that every surface is to be treated as an ideal diffuse reflector (Lambertian), we can set a parameter \( k_d \) for each surface that determines the fraction of the incident light that is to be scattered as diffuse reflections. This parameter is called the **diffuse-reflection coefficient** or the **diffuse reflectivity**. The diffuse reflection in any direction is then a constant, which is equal to the incident light intensity multiplied by the diffuse-reflection coefficient. For a monochromatic light source, parameter \( k_d \) is assigned a constant value in the interval 0.0 to 1.0, according to the reflecting properties we want the surface to have. If we want a highly reflective surface, we set the value of \( k_d \) near 1.0. This produces a brighter surface with the intensity of the reflected light near that of the incident light. If we want to simulate a surface that absorbs most of the incident light, we set the reflectivity to a value near 0.0.

For the background lighting effects, we can assume that every surface is fully illuminated by the ambient light \( I_a \) that we assigned to the scene. Therefore, the ambient contribution to the diffuse reflection at any point on a surface is simply

\[
I_{\text{ambdiff}} = k_d I_a
\]

Ambient light alone, however, produces a flat uninteresting shading for a surface (Color Plate 12), so scenes are rarely rendered using only ambient light. At least
one light source is included in a scene, often as a point source at the viewing position.

When a surface is illuminated by a light source with an intensity $I_l$, the amount of incident light from the source depends on the orientation of the surface relative to the light source direction. A surface that is oriented nearly perpendicular to the illumination direction receives more light from the source than a surface that is tilted at an oblique angle to the direction of the incoming light. This illumination effect can be observed on a white sheet of paper or smooth cardboard that is placed parallel to a sunlit window. As the sheet is slowly rotated away from the window direction, the surface appears less bright. Figure 10 illustrates this effect, showing a beam of light rays incident on two equal-area plane surface elements with different spatial orientations relative to the illumination direction from a distant source (parallel incoming rays).

From Figure 10, we see that the number of light rays intersecting a surface element is proportional to the area of the surface projection perpendicular to the incident light direction. If we denote the angle of incidence between the incoming light direction and the surface normal as $\theta$ (Figure 11), then the projected area of a surface element perpendicular to the light direction is proportional to $\cos \theta$. Therefore, we can model the amount of incident light on a surface from a source with intensity $I_l$ as

$$I_{l, \text{incident}} = I_l \cos \theta$$  

(8)

Using Equation 8, we can model the diffuse reflections from a light source with intensity $I_l$ using the calculation

$$I_{l, \text{diff}} = k_d I_{l, \text{incident}} = k_d I_l \cos \theta$$  

(9)

When the incoming light from the source is perpendicular to the surface at a particular point, $\theta = 90^\circ$ and $I_{l, \text{diff}} = k_d I_l$. As the angle of incidence increases, the illumination from the light source decreases. Furthermore, a surface is illuminated by a point source only if the angle of incidence is in the range $0^\circ$ to $90^\circ$ ($\cos \theta$ is in the interval from 0.0 to 1.0). When $\cos \theta < 0.0$, the light source is behind the surface.

At any surface position, we can denote the unit normal vector as $\mathbf{N}$ and the unit direction vector to a point source as $\mathbf{L}$, as in Figure 12. Then, $\cos \theta = \mathbf{N} \cdot \mathbf{L}$ and the diffuse reflection equation for single point-source illumination at a surface position can be expressed in the form

$$I_{l, \text{diff}} = \begin{cases} k_d I_l (\mathbf{N} \cdot \mathbf{L}), & \text{if } \mathbf{N} \cdot \mathbf{L} > 0 \\ 0.0, & \text{if } \mathbf{N} \cdot \mathbf{L} \leq 0 \end{cases}$$  

(10)

The unit direction vector $\mathbf{L}$ to a nearby point light source is calculated using the surface position and the light-source position:

$$\mathbf{L} = \frac{\mathbf{P}_{\text{source}} - \mathbf{P}_{\text{surf}}}{|\mathbf{P}_{\text{source}} - \mathbf{P}_{\text{surf}}|}$$  

(11)
A light source at “infinity,” however, has no position, only a propagation direction. In that case, we use the negative of the assigned light-source emission direction for the direction of vector \(\mathbf{L}\).

Color Plate 13 illustrates the application of Equation 10 to positions over the surface of a sphere, using selected values for parameter \(k_d\) between 0 and 1. At \(k_d = 0\), no light is reflected and the object surface appears black. Increasing values for \(k_d\) increase the intensity of the diffuse reflections, producing lighter shades of gray. Each projected pixel position for the surface is assigned an intensity value as calculated by the diffuse reflection equation. The surface renderings in this figure illustrate single point-source lighting with no other lighting effects. This is what we might expect to see if we shined a very small flashlight, such as a penlight, on the object in a completely darkened room. For general scenes, however, we expect some surface reflections due to the ambient light in addition to the illumination effects produced by a light source.

We can combine the ambient and point-source intensity calculations to obtain an expression for the total diffuse reflection at a surface position. In addition, many graphics packages introduce an ambient-reflection coefficient \(k_a\) that can be assigned to each surface to modify the ambient-light intensity \(I_a\). This simply provides us with an additional parameter for adjusting the lighting effects in our empirical model. Using parameter \(k_a\), we can write the total diffuse-reflection equation for a single point source as

\[
I_{\text{diff}} = \begin{cases} 
  k_d I_a + k_d I_l (\mathbf{N} \cdot \mathbf{L}), & \text{if } \mathbf{N} \cdot \mathbf{L} > 0 \\
  k_d I_a, & \text{if } \mathbf{N} \cdot \mathbf{L} \leq 0
\end{cases}
\]

where both \(k_a\) and \(k_d\) depend on surface material properties and are assigned values in the range from 0 to 1.0 for monochromatic lighting effects.

### Specular Reflection and the Phong Model

The bright spot, or specular reflection, that we can see on a shiny surface is the result of total, or near total, reflection of the incident light in a concentrated region around the specular-reflection angle. Figure 13 shows the specular reflection direction for a position on an illuminated surface. The specular reflection angle equals the angle of the incident light, with the two angles measured on opposite sides of the unit normal surface vector \(\mathbf{N}\). In this figure, \(\mathbf{R}\) represents the unit vector in the direction of ideal specular reflection, \(\mathbf{L}\) is the unit vector directed toward the point light source, and \(\mathbf{V}\) is the unit vector pointing to the viewer from the selected surface position. Angle \(\phi\) is the viewing angle relative to the specular-reflection direction \(\mathbf{R}\). For an ideal reflector (a perfect mirror), incident light is reflected only in the specular-reflection direction, and we would see reflected light only when vectors \(\mathbf{V}\) and \(\mathbf{R}\) coincide (\(\phi = 0\)).

Objects other than ideal reflectors exhibit specular reflections over a finite range of viewing positions around vector \(\mathbf{R}\). Shiny surfaces have a narrow specular reflection range, and dull surfaces have a wider reflection range. An empirical model for calculating the specular reflection range, developed by Phong Bui Tuong and called the Phong specular-reflection model or simply the Phong model, sets the intensity of specular reflection proportional to \(\cos^n \phi\). Angle \(\phi\) can be assigned values in the range 0° to 90°, so that \(\cos \phi\) varies from 0 to 1.0. The value assigned to the specular-reflection exponent \(n_s\) is determined by the type of surface that we want to display. A very shiny surface is modeled with a large value for \(n_s\) (say, 100 or more), and smaller values (down to 1) are used for duller surfaces. For a perfect reflector, \(n_s\) is infinite. For a rough surface, such as chalk or cinderblock, \(n_s\) is assigned a value near 1. Figures 14 and 15
show the effect of $n_s$ on the angular range for which we can expect to see specular reflections.

The intensity of specular reflection depends on the material properties of the surface and the angle of incidence, as well as other factors such as the polarization and color of the incident light. We can approximately model monochromatic specular intensity variations using a **specular-reflection coefficient**, $W(\theta)$, for each surface. Figure 16 shows the general variation of $W(\theta)$ over the range $\theta = 0^\circ$ to $\theta = 90^\circ$ for a few materials. In general, $W(\theta)$ tends to increase as the angle of incidence increases. At $\theta = 90^\circ$, all the incident light is reflected ($W(\theta) = 1$). The
variation of specular intensity with angle of incidence is described by *Fresnel's Laws of Reflection*. Using the spectral-reflection function $W(\theta)$, we can write the Phong specular-reflection model as

$$I_{\text{spec}} = W(\theta) I_l \cos^n \phi$$

where $I_l$ is the intensity of the light source, and $\phi$ is the viewing angle relative to the specular-reflection direction $R$.

As seen in Figure 16, transparent materials, such as glass, exhibit appreciable specular reflections only as $\theta$ approaches $90^\circ$. At $\theta = 0^\circ$, about 4 percent of the incident light on a glass surface is reflected, and for most of the range of $\theta$, the reflected intensity is less than 10 percent of the incident intensity. However, for many opaque materials, specular reflection is nearly constant for all incidence angles. In this case, we can reasonably model the specular effects by replacing $W(\theta)$ with a constant specular-reflection coefficient $k_s$. We then simply set $k_s$ equal to some value in the range from 0 to 1.0 for each surface.

Because $V$ and $R$ are unit vectors in the viewing and specular-reflection directions, we can calculate the value of $\cos \phi$ with the dot product $V \cdot R$. In addition, no specular effects are generated for the display of a surface if $V$ and $L$ are on the same side of the normal vector $N$ or if the light source is behind the surface. Thus, assuming the specular-reflection coefficient is a constant for any material, we can determine the intensity of the specular reflection due to a point light source at a surface position with the calculation

$$I_{\text{spec}} = \begin{cases} k_s I_l (V \cdot R)^n, & \text{if } V \cdot R > 0 \quad \text{and} \quad N \cdot L > 0 \\ 0, & \text{if } V \cdot R \leq 0 \quad \text{or} \quad N \cdot L \leq 0 \end{cases}$$

The direction for $R$, the reflection vector, can be computed from the directions for vectors $L$ and $N$. As seen in Figure 17, the projection of $L$ onto the direction of the normal vector has a magnitude equal to the dot product $N \cdot L$, which is also equal to the magnitude of the projection of unit vector $R$ onto the direction of $N$. Therefore, from this diagram, we see that

$$R + L = (2N \cdot L)N$$

and the specular-reflection vector is obtained as

$$R = (2N \cdot L)N - L$$

We calculate $V$ using the surface position and the viewing position, in same way that we obtained the unit vector $L$ (Eq. 11). But if a fixed viewing direction
is to be used for all positions in a scene, we can set \( V = (0.0, 0.0, 1.0) \), which is a unit vector in the positive \( z \) direction. Specular calculations take less time to calculate using a constant \( V \), but the displays are not as realistic.

A somewhat simplified Phong model is obtained using the halfway vector \( H \) between \( L \) and \( V \) to calculate the range of specular reflections. If we replace \( V \cdot R \) in the Phong model with the dot product \( N \cdot H \), this simply replaces the empirical \( \cos \phi \) calculation with the empirical \( \cos \alpha \) calculation (Figure 18). The halfway vector is obtained as

\[
H = \frac{L + V}{|L + V|}
\]

(16)

For nonplanar surfaces, \( N \cdot H \) requires less computation than \( V \cdot R \) because the calculation of \( R \) at each surface point involves the variable vector \( N \). Also, if both the viewer and the light source are sufficiently far from the surface, vectors \( V \) and \( L \) are each constants, and thus \( H \) is also constant for all surface points. If the angle between \( H \) and \( N \) is greater than 90°, \( N \cdot H \) is negative and we set the specular-reflection contribution to 0.0.

Vector \( H \) is the orientation direction for the surface that would produce maximum specular reflection in the viewing direction, for a given position of a point light source. For this reason, \( H \) is sometimes referred to as the surface orientation direction for maximum highlights. Also, if vector \( V \) is coplanar with vectors \( L \) and \( R \) (and thus \( N \)), angle \( \alpha \) has the value \( \phi/2 \). When \( V \), \( L \), and \( N \) are not coplanar, \( \alpha > \phi/2 \), depending on the spatial relationship of the three vectors.

### Combined Diffuse and Specular Reflections

For a single point light source, we can model the combined diffuse and specular reflections from a position on an illuminated surface as

\[
I = I_{\text{diff}} + I_{\text{spec}} = k_a I_a + k_d I_l (N \cdot L) + k_s I_l (N \cdot H)^n
\]

(17)

The surface is illuminated only with ambient light if the light source is behind the surface, and there are no specular effects if \( V \) and \( L \) are on the same side of the normal vector \( N \). Color Plate 12 illustrates surface lighting effects produced by the various terms in Equation 17.

### Diffuse and Specular Reflections from Multiple Light Sources

We can place any number of light sources in a scene. For multiple point light sources, we compute the diffuse and specular reflections as a sum of the contributions from the various sources, as follows:

\[
I = I_{\text{ambdiff}} + \sum_{i=1}^{n} [I_{i,\text{diff}} + I_{i,\text{spec}}] = k_a I_a + \sum_{i=1}^{n} I_l [k_d (N \cdot L) + k_s (N \cdot H)^n]
\]

(18)

### Surface Light Emissions

Some surfaces in a scene could be emitting light, as well as reflecting light from their surfaces. For example, a room scene can contain lamps or overhead lighting, and outdoor night scenes could include streetlights, store signs, and automobile headlights. We can empirically model surface light emissions by simply including an emission term \( I_{\text{surfemission}} \) in the illumination model in the same way that we...
simulated background lighting using an ambient light level. This surface emission is then added to the surface reflections resulting from the light-source and the background-lighting illumination.

To illuminate other objects from a light-emitting surface, we could position a directional light source behind the surface to produce a cone of light through the surface. Alternatively, we could simulate the emission with a set of point light sources distributed over the surface. In general, however, an emitting surface is usually not used in the basic illumination model to illuminate other surfaces because of the added calculation time. Rather, surface emissions are used as a simple means for approximating the appearance of the surface of an extended light-source. This produces a glowing effect for the surface.

### Basic Illumination Model with Intensity

#### Attenuation and Spotlights

We can formulate a general, monochromatic illumination model for surface reflections that includes multiple point light sources, attenuation factors, directional light effects (spotlight), infinite sources, and surface emissions as

\[ I = I_{\text{surf emission}} + I_{\text{amb diff}} + \sum_{l=1}^{n} f_l \cdot \text{radatten} \cdot f_l \cdot \text{angatten} \cdot (I_l \cdot \text{diff} + I_l \cdot \text{spec}) \]  \hspace{1cm} (19)

The radial attenuation function \( f_l \cdot \text{radatten} \) is evaluated using Equation 2, and the angular attenuation function is evaluated using Equation 5. For each light source, we calculate the diffuse reflection from a surface point as

\[ I_{l, \text{diff}} = \begin{cases} 0.0, & \text{if } N \cdot L_l \leq 0.0 \text{ (light source behind object)} \\ k_d I_l (N \cdot L_l), & \text{otherwise} \end{cases} \]  \hspace{1cm} (20)

The specular reflection term, due to a point-source illumination, is calculated with similar expressions:

\[ I_{l, \text{spec}} = \begin{cases} 0.0, & \text{if } N \cdot L_l \leq 0.0 \\ k_s I_l \max\{0.0, (N \cdot H_l)^n\}, & \text{otherwise} \end{cases} \]  \hspace{1cm} (21)

To ensure that any pixel intensity does not exceed the maximum allowable value, we can apply some type of normalization procedure. A simple approach is to set a maximum magnitude for each term in the intensity equation. If any calculated term exceeds the maximum, we simply set it to the maximum value. Another way to compensate for intensity overflow is to normalize the individual terms by dividing each by the magnitude of the largest term. A more complicated procedure is to calculate all pixel intensities for the scene, then scale this set of intensities onto the intensity range from 0.0 to 1.0.

Also, the values for the coefficients in the radial attenuation function, and the optical surface parameters for a scene, can be adjusted to prevent calculated intensities from exceeding the maximum allowable value. This is an effective method for limiting intensity values when a single light source illuminates a scene. In general, however, calculated intensities are never allowed to exceed the value 1.0, and negative intensity values are adjusted to the value 0.0.
RGB Color Considerations

For an RGB color description, each intensity specification in the illumination model is a three-element vector that designates the red, green, and blue components of that intensity. Thus, for each light source, \( I_l = (I_{lR}, I_{lG}, I_{lB}) \). Similarly, the reflection coefficients are also specified with RGB components: \( k_s = (k_{sR}, k_{sG}, k_{sB}), k_d = (k_{dR}, k_{dG}, k_{dB}), \) and \( k_a = (k_{aR}, k_{aG}, k_{aB}) \). Each component of the surface color is then calculated with a separate expression. For example, the blue component of the diffuse and specular reflections for a point source are computed from modified expressions 20 and 21 as

\[
I_{lB,\text{diff}} = k_{dB} I_{lB} (N \cdot L) \tag{22}
\]

and

\[
I_{lB,\text{spec}} = k_{dB} I_{lB} \max\{0, (N \cdot H)^n\} \tag{23}
\]

Surfaces are most often illuminated with white light sources, but, for special effects or indoor lighting, we might use other colors for the light sources. We then set the reflectivity coefficients to model a particular surface color. For example, if we want an object to have a blue surface, we select a nonzero value in the range from 0.0 to 1.0 for the blue reflectivity component, \( k_{dB} \), while the red and green reflectivity components are set to zero \( (k_{dR} = k_{dG} = 0.0) \). Any nonzero red or green components in the incident light are absorbed, and only the blue component is reflected.

In his original specular-reflection model, Phong set parameter \( k_s \) to a constant value independent of the surface color. This produces specular reflections that are the same color as the incident light (usually white), which gives the surface a plastic appearance. For a nonplastic material, the color of the specular reflection is actually a function of the surface properties and may be different from both the color of the incident light and the color of the diffuse reflections. We can approximate specular effects on such surfaces by making the specular-reflection coefficient color-dependent, as in Equation 23. Color Plate 14 illustrates color reflections from a matte surface, and Color Plates 15 and 16 show color reflections from metal surfaces.

Another method for setting surface color is to specify the components of diffuse and specular color vectors for each surface, while retaining the reflectivity coefficients as single-valued constants. For an RGB color representation, for instance, the components of these two surface-color vectors could be denoted as \((S_{dR}, S_{dG}, S_{dB})\) and \((S_{sR}, S_{sG}, S_{sB})\). The blue component of the diffuse reflection (Eq. 22) is then calculated as

\[
I_{lB,\text{diff}} = k_d S_{dB} I_{lB} (N \cdot L) \tag{24}
\]

This approach provides somewhat greater flexibility, because surface color parameters and reflectivity values can be set independently.

In some graphics packages, additional lighting parameters are supplied by allowing a light source to be assigned multiple colors, where each color contributes to one of the surface lighting effects. For example, one of the colors can be used as a contribution to the general background lighting in a scene. Similarly, another light-source color can be used as the light intensity for the diffuse-reflection calculations, and a third light-source color can be used in the specular-reflection calculations.
Other Color Representations

We can describe colors using a variety of models other than the RGB representation. For example, a color can be represented using cyan, magenta, and yellow components, or a color could be described in terms of a particular hue along with the perceived brightness and saturation of the color. We can incorporate any of these representations, including color specifications with more than three components, into an illumination model. As an example, Equation 24 can be expressed in terms of any spectral color with wavelength $\lambda$ as

$$I_{\lambda, \text{diff}} = k_d S_d I_{\lambda}(N \cdot L_d)$$  

(25)

Luminance

Another characteristic of color is luminance, which is sometimes also called luminous energy. Luminance provides information about the lightness or darkness level of a color, and it is a psychological measure of our perception of brightness that varies with the amount of illumination we are viewing.

Physically, color is described in terms of the frequency range for visible radiant energy (light), and luminance is calculated as a weighted sum of the intensity components in a particular illumination. Because any actual illumination contains a continuous range of frequencies, a luminance value is computed as

$$\text{luminance} = \int_{\text{visible}} \text{p}(f) I(f) \, df$$  

(26)

Parameter $I(f)$ in this calculation represents the intensity of the light component with a frequency $f$ that is radiating in a particular direction. Parameter $p(f)$ is an experimentally determined proportionality function that varies with both frequency and illumination level. The integration is performed for all intensities over the frequency range contained in the light.

For grayscale and monochromatic displays, we need only the luminance values to describe object lighting. And some graphics packages do allow the lighting parameters to be expressed in terms of luminance. Green components of a light source contribute most to the luminance, and blue components contribute least. Therefore, the luminance of an RGB color source is typically computed as

$$\text{luminance} = 0.299R + 0.587G + 0.114B$$  

(27)

Sometimes, better lighting effects are achieved by increasing the contribution for the green component of each RGB color. One recommendation for this calculation is $0.2125R + 0.7154G + 0.0721B$. The luminance parameter is most often represented with the symbol $Y$, which corresponds to the $Y$ component in the $XYZ$ color model.

4 Transparent Surfaces

We describe an object, such as a glass windowpane, as transparent if we can see things that are behind that object. Similarly, if we cannot see things that are behind an object, it is opaque. In addition, some transparent objects, such as frosted glass and certain plastic materials, are translucent so that the transmitted light is diffused in all directions. Objects viewed through translucent materials appear blurred and are often not clearly identifiable.
A transparent surface, in general, produces both reflected and transmitted light. The light transmitted through the surface is the result of emissions and reflections from the objects and sources behind the transparent object. Figure 19 illustrates the intensity contributions to the surface lighting for a transparent object that is in front of an opaque object.

Translucent Materials

Both diffuse and specular transmission can take place at the surfaces of a transparent object. Diffuse effects are important when translucent materials are to be modeled. Light passing through a translucent material is scattered so that the background objects are seen as blurred images. We can simulate diffuse transmissions by distributing intensity contributions from background objects over a finite area, or we can use ray-tracing methods to simulate translucency. These manipulations are time-consuming, and basic illumination models ordinarily compute only specular-transparency effects.

Light Refraction

Realistic displays of a transparent material are obtained by modeling the refraction path of a ray of light through the material. When a light beam is incident upon a transparent surface, part of it is reflected and part is transmitted through the material as refracted light, as shown in Figure 20. Because the speed of light is different in different materials, the path of the refracted light is different from that of the incident light. The direction of the refracted light, specified by the angle of refraction, is a function of the index of refraction of the material and the incoming direction of the incident light. Index of refraction is defined as the ratio of the speed of light in a vacuum to the speed of light in the material. Angle of refraction $\theta_r$ is calculated from Snell’s law as

$$\sin \theta_r = \frac{n_i}{n_r} \sin \theta_i$$

where $\theta_i$ is the angle of incidence, $n_i$ is the index of refraction for the incident material, and $n_r$ is the index of refraction for the refracting material.

Actually, the index of refraction also depends on other factors, such as the temperature of the material and the wavelength of the incident light. Thus, the various color components of incident white light, for example, are refracted at different angles, which vary with temperature. Furthermore, within anisotropic materials such as crystalline quartz, the speed of light depends on direction, and some transparent materials exhibit double refraction, in which two refracted light rays are generated. For most applications, however, we can use a single average index of refraction for each material, as listed in Table 1. Using the index of refraction for air (approximately 1.0) surrounding a pane of heavy crown glass (refractive index $\approx 1.61$) in Equation 28, with an angle of incidence of $30^\circ$, we obtain a refraction angle of about $18^\circ$ for the light passing through the crown glass.

Refraction occurs whenever a ray moves through the boundary between materials, so in a situation where the ray passes completely through an object, the ray will be refracted twice—one refraction for each boundary transition. Figure 21 illustrates the refraction changes for a ray of light passing through a thin sheet of glass. The overall effect of the refraction is to shift the incident light to a parallel path as it emerges from the material. Because the evaluations for the trigonometric functions in Equation 28 are time-consuming, these refraction effects could be approximated by simply shifting the path of the incident light by an appropriate amount for a given material.
Table 1: Average Index of Refraction for Common Materials

<table>
<thead>
<tr>
<th>Material</th>
<th>Index of Refraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vacuum, air or other gas</td>
<td>1.00</td>
</tr>
<tr>
<td>Ordinary crown glass</td>
<td>1.52</td>
</tr>
<tr>
<td>Heavy crown glass</td>
<td>1.61</td>
</tr>
<tr>
<td>Ordinary flint glass</td>
<td>1.61</td>
</tr>
<tr>
<td>Heavy flint glass</td>
<td>1.92</td>
</tr>
<tr>
<td>Rock salt</td>
<td>1.55</td>
</tr>
<tr>
<td>Quartz</td>
<td>1.54</td>
</tr>
<tr>
<td>Water</td>
<td>1.33</td>
</tr>
<tr>
<td>Ice</td>
<td>1.31</td>
</tr>
</tbody>
</table>

From Snell’s law and the diagram in Figure 20, we can obtain the unit transmission vector $T$ in the refraction direction $\theta_r$ as follows:

$$T = \left( \frac{n_i}{n_r} \cos \theta_i - \cos \theta_r \right) N - \frac{n_i}{n_r} L \quad (29)$$

where $N$ is the unit surface normal and $L$ is the unit vector in the direction from the surface position to the light source. Transmission vector $T$ can be used to locate intersections of the refraction path with objects behind the transparent surface. Including refraction effects in a scene can produce highly realistic displays, but the determination of refraction paths and object intersections requires considerable computation. Most scan-line image-space methods model light transmission with approximations that reduce processing time. Accurate refraction effects are displayed using ray-tracing algorithms.

Basic Transparency Model

A simpler procedure for modeling transparent objects is to ignore the path shifts due to refraction. In effect, this approach assumes there is no change in the index of refraction from one material to another, so that the angle of refraction is always the same as the angle of incidence. This method speeds up the calculation of intensities and can produce reasonable transparency effects for thin, polygonal surfaces.

We can combine the transmitted intensity $I_{\text{trans}}$ through a transparent surface from a background object with the reflected intensity $I_{\text{refl}}$ from the surface (Figure 22) using a transparency coefficient $k_t$. We assign parameter $k_t$ a value between 0.0 and 1.0 to specify how much of the background light is to be transmitted. Total surface intensity is then calculated as

$$I = (1 - k_t)I_{\text{refl}} + k_t I_{\text{trans}} \quad (30)$$

The term $(1 - k_t)$ is the opacity factor. For example, if the transparency factor is assigned the value 0.3, then 30 percent of the background light is combined with 70 percent of the reflected surface illumination.
This procedure can be used to combine the lighting effects from any number of transparent and opaque objects, so long as we process the surfaces in a depth-first order (i.e., back to front). For example, looking through an empty drinking glass, we can see opaque objects that are behind its two transparent surfaces. Similarly, when we look through the windshield of an automobile, objects inside the car are visible, as well as objects that may be behind the back window.

For highly transparent objects, we assign $k_t$ a value near 1.0. Nearly opaque objects transmit very little light from background objects, and we can set $k_t$ to a value near 0.0 for these materials. It is also possible to allow $k_t$ to be a function of position over the surface such that different parts of an object can transmit more or less of the light from the background surfaces.

A depth-sorting visibility algorithm can be modified to handle transparency by first sorting surfaces in depth order, then determining whether any visible surface is transparent. If it is, its reflected surface intensity is combined with the surface intensity of objects behind it to obtain the pixel intensity at each projected surface point.

Transparency effects could also be implemented using a modified depth-buffer approach. We can divide the surfaces in a scene into two groups so that all the opaque surfaces are processed first. At this point, the frame buffer contains the intensities of the visible surfaces, and the depth buffer contains their depths. Then, the depth positions of the transparent objects are compared to the values previously stored in the depth buffer. If any transparent surface is visible, its reflected intensity is calculated and combined with the opaque surface intensity previously stored in the frame buffer. This method can be modified to produce more accurate displays by using additional storage for the depth and other parameters of the transparent surfaces. This allows depth values for the transparent surfaces to be compared to each other, as well as to the depth values of the opaque surfaces. Visible transparent surfaces are then rendered by combining their surface intensities with those of the visible and opaque surfaces behind them.

Another approach is the A-buffer method. For each pixel position in the A-buffer, surface patches for all overlapping surfaces are saved and sorted in depth order. Then, intensities for the transparent and opaque surface patches that overlap in depth are combined in the proper visibility order to produce the final averaged intensity for the pixel.

### 5 Atmospheric Effects

Another factor that is sometimes included in an illumination model is the effect of the atmosphere on an object’s color. A hazy atmosphere makes colors fade and objects appear dimmer. Thus, we could specify a function to modify surface colors according to the amount of dust, smoke, or smog that we want to simulate in the atmosphere. The hazy-atmosphere effect is often simulated with an exponential attenuation function such as

$$f_{\text{atmo}}(d) = e^{-\rho d} \quad (31)$$

or

$$f_{\text{atmo}}(d) = e^{-(\rho d)^2} \quad (32)$$

The value assigned to $d$ is the distance of the object from the viewing position. In addition, we use parameter $\rho$ in either of these exponential functions to set a positive density value for the atmosphere. Higher values for $\rho$ produce a denser atmosphere and cause surface colors to be more muted. After the surface color of an object has been computed, we multiply that color by one of the atmosphere illumination models and surface-rendering methods.
functions to decrease its intensity by an amount that depends on the value we set for the density of the atmosphere.

Instead of an exponential function, we could simplify the atmospheric attenuation calculations by using the linear depth-cueing function. This decreases the intensity of surface colors for distant objects, but we then have no provision for varying the density of the atmosphere.

Sometimes we might also want to simulate an atmosphere color. For example, the air in a smoky room could be modeled with a slate-gray color, or perhaps a pale blue. The following calculation could then be used to combine the atmosphere color with an object’s color:

\[ I = f_{atmo}(d)I_{obj} + [1 - f_{atmo}(d)]I_{atmo} \]  \hspace{1cm} (33)

where \( f_{atmo} \) is an exponential or linear atmosphere-attenuation function.

---

6 Shadows

Visibility detection methods can be used to locate regions that are not illuminated by light sources. With the viewing position at the location of a light source, we can determine which surface sections in the scene are not visible. These are the shadow areas. Once we have determined the shadow areas for all light sources, the shadows could be treated as surface patterns and stored in pattern arrays.

Shadow patterns generated by a visible-surface detection method are valid for any selected viewing position, so long as the light-source positions are not changed. Surfaces that are visible from the view position are shaded according to the lighting model, which can be combined with texture patterns. We can display shadow areas with ambient light intensity only, or we could combine the ambient light with specified surface textures.

---

7 Camera Parameters

The viewing and illumination procedures we have considered so far produce sharp images that are equivalent to photographing a scene with a pinhole camera. When we photograph an actual scene, however, we can adjust the camera so that only selected objects are in focus. Other objects are then more or less out of focus, depending on the depth distribution of the objects in the scene. We can simulate the appearance of out-of-focus positions in a computer-graphics program, by projecting each position to an area covering multiple pixel positions, with the object colors merged into other objects to produce a blurred projection pattern. This procedure is similar to the methods used in antialiasing, and we can incorporate the camera effects into either a scan-line or a ray-tracing algorithm. Computer-generated scenes appear more realistic when focusing effects are included, but the focusing calculations are time-consuming.

---

8 Displaying Light Intensities

A surface intensity calculated by an illumination model can have any value in the range from 0.0 to 1.0, but a computer-graphics system can display only a limited set of intensities. Therefore, a calculated intensity value must be converted to one of
the allowable system values. In addition, the allowable number of system intensity levels can be distributed so that they correspond to the way that our eyes perceive intensity differences. When we display scenes on a bilevel system, we could convert calculated intensities into halftone patterns, as discussed in Section 9.

**Distributing System Intensity Levels**

For any system, the allowable number of intensity levels can be distributed over the range from 0.0 to 1.0 so that this distribution corresponds to our perception of equal intensity intervals between levels. We perceive relative light intensities the same way that we perceive relative sound intensities: on a logarithmic scale. This means that if the ratio of two intensity values is the same as the ratio of two other intensities, we perceive the difference between each pair of intensities to be the same. For example, we perceive the difference between intensities 0.20 and 0.22 to be the same as the difference between 0.80 and 0.88. Therefore, to display \( n + 1 \) successive intensity levels with equal perceived brightness, the intensity levels on the monitor should be spaced so that the ratio of successive intensities is constant, as follows:

\[
\frac{I_1}{I_0} = \frac{I_2}{I_1} = \cdots = \frac{I_n}{I_{n-1}} = r
\]

(34)

where \( I \) represents the intensity of one of the color components of a light. The lowest level that can be displayed is represented as \( I_0 \) and the highest is represented as \( I_n \). Any intermediate intensity can then be expressed in terms of \( I_0 \) as

\[
I_k = r^k I_0
\]

(35)

We can calculate the value of \( r \), given the values of \( I_0 \) and \( n \) for a particular system, by substituting \( k = n \) in the previous expression. Because \( I_n = 1.0 \), we have

\[
r = \left( \frac{1.0}{I_0} \right)^{1/n}
\]

(36)

Thus, the calculation for \( I_k \) in Equation 35 can be rewritten as

\[
I_k = I_0^{(n-k)/n}
\]

(37)

For example, if \( I_0 = \frac{1}{8} \) for a system with \( n = 3 \), we have \( r = 2 \) and the four intensity values are \( \frac{1}{8}, \frac{1}{4}, \frac{1}{2}, \) and \( 1.0 \).

The lowest intensity value \( I_0 \) depends on the characteristics of the monitor and is typically in the range from 0.005 to around 0.025. This residual intensity on a video monitor is due to reflected light from the screen phosphors. Therefore, a “black” region on the screen will always have some intensity value above 0.0. For a grayscale display with 8 bits per pixel (\( n = 255 \)) and \( I_0 = 0.01 \), the ratio of successive intensities is approximately \( r = 1.0182 \). The approximate values for the 256 intensities on this system are 0.0100, 0.0102, 0.0104, 0.0106, 0.0107, 0.0109, \ldots, 0.9821, and 1.0000.

Similar methods are used with RGB color components. For example, we can express the intensity of the blue component of a color at level \( k \) in terms of the lowest attainable blue value as

\[
I_{Bk} = r_B^k I_{B0}
\]

(38)

where

\[
r_B = \left( \frac{1.0}{I_{B0}} \right)^{1/n}
\]

(39)

and \( n \) is the number of intensity levels.
**Gamma Correction and Video Lookup Tables**

When we display color or monochromatic images on a video monitor, the perceived brightness variations are nonlinear, but illumination models produce a linear variation for intensity values. The RGB color (0.25, 0.25, 0.25) obtained from a lighting model represents one-half the intensity of the color (0.5, 0.5, 0.5). Usually, these calculated intensities are then stored in an image file as integer values ranging from 0 to 255, with one byte for each of the three RGB components. This intensity file is also linear, so a pixel with the value (64, 64, 64) represents half the intensity of a pixel with the value (128, 128, 128). The electron-gun voltages, which control the number of electrons striking the phosphor screen, produce brightness levels as determined by the monitor response curve shown in Figure 23. Therefore, the displayed intensity value (64, 64, 64) would not appear to be half as bright as the value (128, 128, 128).

To compensate for monitor nonlinearities, graphics systems use a video lookup table that adjusts the linear input intensity values. The monitor response curve is described with the exponential function

$$I = aV^\gamma$$  \hspace{1cm} (40)

Parameter $I$ is displayed intensity and parameter $V$ is the corresponding electron-gun voltage. Values for parameters $a$ and $\gamma$ depend on the characteristics of the monitor used in the graphics system. Thus, if we want to display a particular intensity value $I$, the voltage value to produce this intensity is

$$V = \left(\frac{I}{a}\right)^{1/\gamma}$$  \hspace{1cm} (41)

This calculation is referred to as the gamma correction of intensity, and gamma values are typically in the range from about 1.7 to 2.3. The National Television System Committee (NTSC) signal standard is $\gamma = 2.2$. Figure 24 shows a gamma-correction curve using the NTSC gamma value with both intensity and voltage normalized on the interval from 0 to 1.0. Equation 41 is used to set up

![Figure 23](image1.png)  \hspace{1cm} ![Figure 24](image2.png)

---

**Figure 23**
A typical monitor response curve, showing the variation in displayed intensity (or “brightness”) as a function of the normalized electron-gun voltage.

**Figure 24**
A video lookup correction curve for mapping a normalized intensity value to a normalized electron-gun voltage, using gamma correction with $\gamma = 2.2$.  

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**514**
the video lookup table that converts integer intensity values in an image file to values that control the electron-gun voltages.

We can combine gamma correction with logarithmic intensity mapping to produce a lookup table. If \( I \) is an input intensity value from an illumination model, we first locate the nearest intensity \( I_k \) from a table of values created with Equation 34 or Equation 37. Alternatively, we could determine the level number for this intensity value with the calculation

\[
k = \text{round}\left[ \log_r \left( \frac{I}{I_0} \right) \right]
\]  

(42)

then we compute the intensity value at this level using Equation 37. Once we have the intensity value \( I_k \), we can calculate the electron-gun voltage as

\[
V_k = \left( \frac{I_k}{a} \right)^{1/\gamma}
\]  

(43)

Values \( V_k \) can then be placed in the lookup tables, with values for \( k \) stored in the frame-buffer pixel positions. If a particular system has no lookup table, computed values for \( V_k \) could be stored directly in the frame buffer. The combined conversion to a logarithmic intensity scale followed by calculation of the \( V_k \) using Equation 43 is also sometimes referred to as gamma correction.

If the video amplifiers of a monitor are designed to convert the linear intensity values to electron-gun voltages, we cannot combine the two intensity conversion processes. In this case, gamma correction is built into the hardware, and the logarithmic values \( I_k \) must be precomputed and stored in the frame buffer (or the color table).

### Displaying Continuous-Tone Images

High-quality computer graphics systems generally provide 256 intensity levels for each color component, but acceptable displays can be obtained for many applications with fewer levels. A four-level system provides minimum shading capability for continuous-tone images, while photo-realistic images can be generated on systems that are capable of from 32 to 256 intensity levels per pixel.

Figure 25 shows a continuous-tone photograph displayed with various intensity levels. When a small number of intensity levels are used to reproduce a continuous-tone image, the borders between the different intensity regions (called contours) are clearly visible. In the 2-level reproduction, the facial features in the photograph are just barely identifiable. Using 4 intensity levels, we begin to identify the original shading patterns, but the contouring effects are glaring. With 8 intensity levels, contouring effects are still obvious, but we begin to have a better indication of the original shading. At 16 or more intensity levels, contouring effects diminish and the reproductions are very close to the original. Reproductions of continuous-tone images using more than 32 intensity levels show only very subtle differences from the original.

### 9 Halftone Patterns and Dithering Techniques

With a system that has very few available intensity levels, we can create an apparent increase in the number of available intensities by incorporating multiple pixel positions into the display of each intensity value for a scene. When we view a small region consisting of several pixel positions, our eyes tend to integrate
or average the fine detail into an overall intensity. Bilevel monitors and printers, in particular, can take advantage of this visual effect to produce pictures that appear to be displayed with multiple intensity values.

Continuous-tone photographs are reproduced for publication in newspapers, magazines, and books with a printing process called halftoning, and the reproduced pictures are called halftones. For a black-and-white photograph, each constant intensity area is reproduced as a set of small black circles on a white background. The diameter of each circle is proportional to the darkness required for that intensity region. Darker regions are printed with larger circles, and lighter regions are printed with smaller circles (more white space). Figure 26 shows an enlarged section of a grayscale halftone reproduction. Color halftones are printed using small circular dots of various sizes and colors. Book and magazine halftones are printed on high-quality paper using approximately 60 to 80 circles of varying diameter per centimeter. Newspapers use lower-quality paper and lower resolution (about 25 to 30 dots per centimeter).
Halftone Approximations

In computer graphics, halftone reproductions are simulated using rectangular pixel regions that are called halftone approximation patterns, or just pixel patterns. The number of intensity levels that we can display with this method depends on how many pixels we include in the rectangular grids and how many levels a system can display. With $n \times n$ pixels for each grid on a bilevel system, we can represent $n^2 + 1$ intensity levels. Figure 27 shows one way to set up pixel patterns to represent five intensity levels that could be used with a bilevel system. In pattern 0, all pixels are turned off; in pattern 1, one pixel is turned on; and in pattern 4, all four pixels are turned on. An intensity value $I$ in a scene is mapped to a particular pattern according to the range listed below each grid shown in the figure. Pattern 0 is used for $0.0 \leq I < 0.2$, pattern 1 for $0.2 \leq I < 0.4$, and pattern 4 is used for $0.8 \leq I \leq 1.0$.

With $3 \times 3$ pixel grids on a bilevel system, we can display 10 intensity levels. One way to set up the 10 pixel patterns for these levels is shown in Figure 28. Pixel positions are chosen at each level so that the patterns approximate the increasing circle sizes used in halftone reproductions. That is, the “on” pixel positions are near the center of the grid for lower intensity levels and expand outward as the intensity level increases.

For any pixel-grid size, we can represent the pixel patterns for the various possible intensities with a mask (matrix) of pixel position numbers. For example, the following mask can be used to generate the nine $3 \times 3$ grid patterns for intensity levels above 0 shown in Figure 28:

$$
\begin{bmatrix}
8 & 3 & 7 \\
5 & 1 & 2 \\
4 & 9 & 6 \\
\end{bmatrix}
$$

To display a particular intensity with level number $k$, we turn on each pixel whose position number is less than or equal to $k$.

Figure 27 illustrates a set of $2 \times 2$ pixel grid patterns that can be used to display five intensity levels on a bilevel system, showing the “on” pixels as green circles. The intensity values that are mapped to each of the grid patterns are listed below the pixel arrays.

Figure 28 shows a set of $3 \times 3$ pixel grid patterns that can be used to display 10 intensities on a bilevel system, showing the “on” pixels as green circles. The intensity values that are mapped to each of the grid patterns are listed below the pixel arrays.
Although the use of $n \times n$ pixel patterns increases the number of intensities that can be represented, the resolution of the display area is reduced by a factor of $1/n$ in the $x$ and $y$ directions. Using $2 \times 2$ grid patterns on a $512 \times 512$ screen area, for instance, reduces the resolution to $256 \times 256$ intensity positions; and with $3 \times 3$ patterns, we reduce the resolution of the $512 \times 512$ area to $128 \times 128$.

Another problem with pixel grids is that subgrid patterns become apparent as the grid size increases. The grid size that can be used without distorting the intensity variations depends on the size of a displayed pixel. Therefore, for systems with lower resolution (fewer pixels per centimeter), we must be satisfied with fewer intensity levels. On the other hand, high-quality displays require at least 64 intensity levels. This means that we need $8 \times 8$ pixel grids. And to achieve a resolution equivalent to that of halftones in books and magazines, we must display 60 dots per centimeter. Thus, we need to be able to display $60 \times 8 = 480$ dots per centimeter. Some devices, such as high-quality film recorders, can display this resolution.

Pixel-grid patterns for halftone approximations must also be constructed to minimize contouring and other visual effects not present in the original scene. We can minimize contouring by evolving each successive grid pattern from the previous pattern. That is, we form the pattern at level $k$ by adding an “on” position to the grid pattern used for level $k - 1$. Thus, if a pixel position is on for one grid level, it is on for all higher levels (Figs. 27 and 28). We can minimize the introduction of other visual effects by avoiding symmetrical patterns. With a $3 \times 3$ pixel grid, for instance, the third intensity level above zero would be better represented by the pattern in Figure 29(a) than by any of the symmetrical arrangements in Figure 29(b). The symmetrical patterns in this figure would produce either vertical, horizontal, or diagonal streaks in any large area shaded with intensity level 3. For hardcopy output on devices such as film recorders and some printers, isolated pixels are not effectively reproduced. Therefore a grid pattern with a single “on” pixel or with isolated “on” pixels, as in Figure 30, should be avoided.

Halftone-approximation methods can be applied also to increase the number of intensity options on systems that are capable of displaying more than two intensities per pixel. For example, on a grayscale system that can display four intensity values per pixel, we can use $2 \times 2$ pixel grids to represent 13 different intensity levels. Figure 31 illustrates one way to set up the 13 pixel-grid patterns, where each pixel can be set to intensity level 0, 1, 2, or 3.

Similarly, we can use pixel-grid patterns to increase the number of intensities that can be represented on a color system. A three-bit-per-pixel RGB system, for example, uses one bit per pixel for each color gun. Thus, a pixel is displayed with three phosphor dots, so that the pixel can be assigned any one of eight different

**FIGURE 29**
For a $3 \times 3$ pixel grid, the pattern in (a) is better than any of the symmetrical patterns in (b) for representing the third intensity level above 0.

(a) ![Pattern](image1)

(b) ![Pattern](image2)

**FIGURE 30**
Halftone grid patterns with isolated pixels that cannot be reproduced effectively on some hardcopy devices.
colors (including black and white). But with $2 \times 2$ pixel-grid patterns, we have 12 phosphor dots that we can use to represent a color, as shown in Figure 32. The red electron gun can activate any combination of the four red dots in the grid pattern, and this provides five possible settings for the red color of the pattern. The same is true for the green and blue guns, which gives us a total of 125 different color combinations that can be represented with our $2 \times 2$ grid patterns.

Dithering Techniques

The term dithering is used in various contexts. Primarily, it refers to techniques for approximating halftones without reducing resolution, as pixel-grid patterns do. However, dithering is sometimes used also as a synonym for any halftone-approximation scheme, and sometimes it is used as another term for color halftone approximations.

Random values added to pixel intensities to break up contours are often referred to as dither noise. Various algorithms have been used to generate the random distributions. The effect is to add noise over an entire picture, which tends to soften intensity boundaries.

A method called ordered dither generates intensity variations with a one-to-one mapping of points in a scene to pixel positions using a dither matrix $D_n$ to select an intensity level. Matrix $D_n$ contains $n \times n$ elements that are assigned distinct positive integer values in the range from 0 to $n^2 - 1$. For example, we can generate four intensity levels with

$$D_2 = \begin{bmatrix} 3 & 1 \\ 0 & 2 \end{bmatrix}$$

and we can generate nine intensity levels with

$$D_3 = \begin{bmatrix} 7 & 2 & 6 \\ 4 & 0 & 1 \\ 3 & 8 & 5 \end{bmatrix}$$

The matrix elements for $D_2$ and $D_3$ are in the same order as the pixel mask for setting up $2 \times 2$ and $3 \times 3$ pixel grids, respectively. With a bilevel system, we determine the display intensity values by comparing input intensities to the matrix elements. Each input intensity is first scaled to the range $0 \leq I \leq n^2$. If the intensity $I$ is to be applied to screen position $(x, y)$, we calculate the reference position (row and column) in the dither matrix as

$$j = (x \mod n) + 1, \quad k = (y \mod n) + 1$$

If $I > D_n(j, k)$, we turn on the pixel at position $(x, y)$. Otherwise, the pixel is off. For RGB color applications, this procedure is implemented for the intensity of each of the individual color components (red, green, and blue).
Elements of the dither matrix are assigned in accordance with the guidelines discussed for pixel grids. That is, we want to minimize artificial visual effects, such as contouring. Order dither produces constant intensity areas identical to those generated with pixel-grid patterns when the values of the matrix elements correspond to those in the halftone-approximation grid mask. Variations from the pixel-grid displays occur at the boundary of two different intensity areas.

Typically, the number of intensity levels is taken to be a multiple of 2. Higher-order dither matrices, \( n \geq 4 \), are then obtained from lower-order matrices using the recurrence relation

\[
D_n = \begin{bmatrix}
4D_{n/2} + D_2(1, 1)U_{n/2} & 4D_{n/2} + D_2(1, 2)U_{n/2} \\
4D_{n/2} + D_2(2, 1)U_{n/2} & 4D_{n/2} + D_2(2, 2)U_{n/2}
\end{bmatrix}
\]  

(48)

Parameter \( U_{n/2} \) represents the “unity” matrix (all elements are 1). For example, if \( D_2 \) is specified as in Equation 45, then recurrence relation 48 yields

\[
D_4 = \begin{bmatrix}
15 & 7 & 13 & 5 \\
3 & 11 & 1 & 9 \\
12 & 4 & 10 & 6 \\
0 & 8 & 2 & 10
\end{bmatrix}
\]  

(49)

Another method for mapping a picture with \( m \times n \) points to a display area with \( m \times n \) pixels is error diffusion. Here, the error between an input intensity value and the selected intensity level at a given pixel position is dispersed, or diffused, to pixel positions to the right and below the current pixel position. Starting with a matrix \( \mathbf{M} \) of intensity values obtained by scanning a photograph, we want to construct an array \( \mathbf{I} \) of pixel intensity values for an area of the screen. We do this by first scanning across the rows of \( \mathbf{M} \), from left to right, starting with the top row, and determining the nearest available pixel-intensity level for each element of \( \mathbf{M} \). Then the error between the value stored in matrix \( \mathbf{M} \) and the displayed intensity level at each pixel position is distributed to neighboring elements using the following simplified algorithm:

```c
for (j = 0; j < m; j++)
    for (k = 0; k < n; k++) {
        /* Determine the available system intensity value 
           * that is closest to the value of M [j][k] and 
           * assign this value to I [j][k]. 
        */
        error = M [j][k] - I [j][k];
        I [j][k+1] = M [j][k+1] + alpha * error;
        I [j+1][k] = M [j+1][k] + gamma * error;
        I [j+1][k+1] = M [j+1][k+1] + delta * error;
    }
```

Once the elements of matrix \( \mathbf{I} \) have been assigned intensity-level values, we then map the matrix to an area of a display device such as a printer or video monitor. Of course, we cannot disperse the error past the last matrix column (\( k = n \)) or below the last matrix row (\( j = m \)), and for a bilevel system, the system intensity values are just 0 and 1. Parameters for distributing the error can be chosen to satisfy the following relationship:

\[
\alpha + \beta + \gamma + \delta \leq 1
\]  

(50)
One choice for the error-diffusion parameters that produces fairly good results is \((\alpha, \beta, \gamma, \delta) = \left(\frac{7}{16}, \frac{3}{16}, \frac{5}{16}, \frac{1}{16}\right)\). Figure 33 illustrates the error distribution using these parameter values. Error diffusion sometimes produces “ghosts” in a picture by repeating, or echoing, certain parts of the picture, particularly with facial features such as hairlines and nose outlines. Ghosting can often be reduced in these cases by choosing values for the error-diffusion parameters that sum to a value less than 1 and by rescaling the matrix values after the dispersion of errors. One way to rescale is to multiply all matrix elements by 0.8 and then add 0.1. Another method for improving picture quality is to alternate the scanning of matrix rows from right to left and left to right.

A variation on the error-diffusion method is dot diffusion. In this method, the \(m \times n\) array of intensity values is divided into 64 classes numbered from 0 to 63, as shown in Figure 34. The error between a matrix value and the displayed intensity is then distributed only to those neighboring matrix elements that have a larger class number. Distribution of the 64 class numbers is based on minimizing the number of elements that are completely surrounded by elements with a lower class number, because this would tend to direct all errors of the surrounding elements to that one position.

### 10 Polygon Rendering Methods

Intensity calculations from an illumination model can be applied to surface rendering in various ways. We could use an illumination model to determine the surface intensity at every projected pixel position, or we could apply the illumination model to a few selected points and approximate the intensity at the other surface positions. Graphics packages typically perform surface rendering using scan-line algorithms that reduce processing time by dealing only with polygon surfaces and by calculating surface intensity only at the vertices. The vertex intensities are then
interpolated to the other positions on the polygon surface. Other, more accurate polygon scan-line rendering methods have been developed, and ray-tracing algorithms calculate the intensity at each projected surface point for curved or planar surfaces. In this section, we consider the scan-line surface-rendering schemes that are applied to polygons.

Constant-Intensity Surface Rendering

The simplest method for rendering a polygon surface is to assign the same color to all projected surface positions. In this case, we use the illumination model to determine the intensity for the three RGB color components at a single surface position, such as a vertex or the polygon centroid. This approach, called constant-intensity surface rendering or flat surface rendering, provides a fast and simple method for displaying polygon facets on an object, which can be useful for quickly generating the general appearance of a curved surface, as in Color Plate 17(b). Flat rendering is also useful in design or other applications where we might want quickly to identify the individual polygonal facets used to model a curved surface.

In general, flat surface rendering of a polygon provides an accurate display of the surface if all of the following assumptions are valid:

- The polygon is one face of a polyhedron and not a section of a curved-surface approximation mesh.
- All light sources illuminating the polygon are sufficiently far from the surface that \( \mathbf{N} \cdot \mathbf{L} \) and the attenuation function are constant over the area of the polygon.
- The viewing position is sufficiently far from the polygon that \( \mathbf{V} \cdot \mathbf{R} \) is constant over the area of the polygon.

Even if some of these conditions are not true, we can still reasonably approximate surface lighting effects using constant-intensity surface rendering if the polygon facets of an object are small.

Gouraud Surface Rendering

The Gouraud surface rendering scheme, devised by Henri Gouraud and also referred to as intensity-interpolation surface rendering, linearly interpolates vertex intensity values across the polygon faces of an illuminated object. Developed for rendering a curved surface that is approximated with a polygon mesh, the Gouraud method smoothly transitions the intensity values for each polygon facet into the values for adjacent polygons along the common edges. This interpolation of intensities across the polygon area eliminates the intensity discontinuities that can occur in flat surface rendering.

Each polygon section of a tessellated curved surface is processed by the Gouraud surface-rendering method using the following procedures:

1. Determine the average unit normal vector at each vertex of the polygon.
2. Apply an illumination model at each polygon vertex to obtain the light intensity at that position.
3. Interpolate the vertex intensities linearly over the projected area of the polygon.
At each polygon vertex, we obtain a normal vector by averaging the normal vectors of all polygons in the surface mesh that share that vertex, as illustrated in Figure 35. Thus, for any vertex position \( V \), we obtain the unit vertex normal with the calculation

\[
N_V = \frac{\sum_{k=1}^{n} N_k}{\left| \sum_{k=1}^{n} N_k \right|}
\]  

(51)

Once we have obtained the normal vector at a vertex, we invoke the illumination model to obtain the surface intensity at that point.

After all vertex intensities have been computed for a polygonal facet, we can interpolate the vertex values to obtain the intensities at positions along scan lines that intersect the projected area of the polygon, as demonstrated in Figure 36. For each scan line, the intensity at the intersection of the scan line with a polygon edge is linearly interpolated from the intensities at the endpoints of that edge. For the example in Figure 36, the polygon edge with endpoint vertices at positions 1 and 2 is intersected by the scan line at point 4. A fast method for obtaining the intensity at point 4 is to interpolate between the values at vertices 1 and 2 using only the vertical displacement of the scan line, as follows:

\[
I_4 = \frac{y_4 - y_2}{y_1 - y_2} I_1 + \frac{y_1 - y_4}{y_1 - y_2} I_2
\]  

(52)

In this expression, the symbol \( I \) represents the intensity for one of the RGB color components. Similarly, the intensity at the right intersection of this scan line (point 5) is interpolated from intensity values at vertices 2 and 3. From these two boundary intensities, we linearly interpolate to obtain the pixel intensities for positions across the scan line. The intensity for one of the RGB color components at point \( p \) in Figure 36, for instance, is calculated from the intensities at points 4 and 5 as

\[
I_p = \frac{x_5 - x_p}{x_5 - x_4} I_4 + \frac{x_p - x_4}{x_5 - x_4} I_5
\]  

(53)

In the implementation of Gouraud rendering, we can perform the intensity calculations represented by Equations 52 and 53 efficiently by using...
incremental methods. Starting from a scan line that intersects one of the polygon vertices, we can incrementally obtain intensity values for other scan lines that intersect an edge that is connected to that vertex. Assuming that the polygon facets are convex, each scan line crossing the polygon has two edge intersections, such as points 4 and 5 in Figure 36. Once we have obtained the intensities at the two edge intersections for a scan line, we apply the incremental procedures to obtain pixel intensities across the scan line.

As an example of the incremental calculation of intensities, we consider scan lines $y$ and $y-1$ in Figure 37, which intersect the left edge of a polygon. If scan line $y$ is the next scan line below the vertex at $y_1$ with intensity $I_1$, that is $y = y_1 - 1$, then we can compute the intensity $I$ on scan line $y$ from Equation 52 as

$$I = I_1 + \frac{I_2 - I_1}{y_1 - y_2}$$  (54)

Continuing down the polygon edge, the intensity along this edge for the next scan line, $y - 1$, is

$$I' = I + \frac{I_2 - I_1}{y_1 - y_2}$$  (55)

Thus, each successive intensity value down the edge is computed simply by adding the constant term $(I_2 - I_1)/(y_1 - y_2)$ to the previous intensity value. Similar incremental calculations are used to obtain intensities at successive horizontal pixel positions along each scan line.

Gouraud surface rendering can be combined with a hidden-surface algorithm to fill in the visible polygons along each scan line. An example of a three-dimensional object rendered with the Gouraud method appears in Color Plate 17(c).

This intensity-interpolation method eliminates the discontinuities associated with flat rendering, but it has some other deficiencies. Highlights on the surface are sometimes displayed with anomalous shapes, and the linear intensity interpolation can cause bright or dark intensity streaks, called Mach bands, to appear on the surface. These effects can be reduced by dividing the surface into a greater number of polygon faces or by using more precise intensity calculations.

**Phong Surface Rendering**

A more accurate interpolation method for rendering a polygon mesh was subsequently developed by Phong Bui Tuong. This approach, called Phong surface rendering or normal-vector interpolation rendering, interpolates normal vectors instead of intensity values. The result is a more accurate calculation of intensity values, a more realistic display of surface highlights, and a great reduction in the
Mach-band effect. However, the Phong method requires more computation than the Gouraud method.

Each polygon section of a tessellated curved surface is processed by the Phong surface-rendering method using the following procedures:

1. Determine the average unit normal vector at each vertex of the polygon.
2. Interpolate the vertex normals linearly over the projected area of the polygon.
3. Apply an illumination model at positions along scan lines to calculate pixel intensities using the interpolated normal vectors.

Interpolation procedures for normal vectors in the Phong method are the same as those for the intensity values in the Gouraud method. The normal vector $\mathbf{N}$ in Figure 38 is interpolated vertically from the normal vectors at vertices 1 and 2 as

$$
\mathbf{N} = \frac{y - y_2}{y_1 - y_2} \mathbf{N}_1 + \frac{y_1 - y}{y_1 - y_2} \mathbf{N}_2
$$

(56)

This result must be re-normalized before we perform our shading calculations. We apply the same incremental methods for obtaining normal vectors on successive scan lines and at successive pixel positions along scan lines. The difference between the two surface-rendering approaches is that we must now apply the illumination model at every projected pixel position along the scan lines to obtain the surface intensity values.

**Fast Phong Surface Rendering**

We can reduce processing time in the Phong-rendering method by approximating some of the illumination-model calculations. **Fast Phong surface rendering** performs the intensity calculations using a truncated Taylor-series expansion and limiting the polygon facets to triangular surface patches.

Because the Phong method interpolates normal vectors from the vertex normals, we can write the expression for calculating the surface normal $\mathbf{N}$ at position $(x, y)$ in a triangular patch as

$$
\mathbf{N} = A x + B y + C
$$

(57)

where vectors $A$, $B$, and $C$ are determined from the three vertex equations:

$$
\mathbf{N}_k = A x_k + B y_k + C, \quad \text{for } k = 1, 2, 3
$$

(58)

with $(x_k, y_k)$ denoting a projected triangle vertex position on the pixel plane.
Omitting the reflectivity and attenuation parameters, we can write the calculation for light-source diffuse reflection from a surface point \((x, y)\) as

\[
I_{\text{diff}}(x, y) = \frac{L \cdot N}{||L|| \cdot ||N||} = \frac{L \cdot (Ax + By + C)}{||L|| \cdot ||Ax + By + C||} = \frac{(L \cdot A)x + (L \cdot B)y + L \cdot C}{||L|| \cdot ||Ax + By + C||}
\]  

(59)

This expression can be written in the form

\[
I_{\text{diff}}(x, y) = \frac{ax + by + c}{\sqrt{dx^2 + ey^2 + fx + gy + h^2}}
\]  

(60)

where parameters such as \(a, b, c,\) and \(d\) are used to represent the various dot products. For example,

\[
a = \frac{L \cdot A}{||L||}
\]  

(61)

Finally, we can express the denominator in Equation 60 as a Taylor series expansion and retain terms up to the second degree in \(x\) and \(y\). This yields

\[
I_{\text{diff}}(x, y) = T_5 x^2 + T_4 xy + T_3 y^2 + T_2 x + T_1 y + T_0
\]  

(62)

where each \(T_k\) is a function of the various parameters in Equation 60, such as \(a, b,\) and \(c\).

Using forward differences, we then evaluate Equation 62 with only two additions for each pixel position \((x, y)\) once the initial forward-difference parameters have been evaluated. Although the simplifications in the fast-Phong approach reduce the Phong surface-rendering calculations, it still takes approximately twice as long to render a surface with the fast-Phong method as it does with Gouraud surface rendering. And the basic Phong method, using forward-difference calculations, takes about 6 to 7 times longer than Gouraud rendering.

Fast-Phong rendering for diffuse reflection can be extended to include specular reflections, using similar approximations for evaluating the specular terms such as \((N \cdot H)^n\). In addition, we can generalize the algorithm to include a finite viewing position and polygons other than triangles.

### 11 OpenGL Illumination and Surface-Rendering Functions

A variety of routines are available in OpenGL for setting up point light sources, selecting surface-reflection coefficients, and choosing values for other parameters in the basic illumination model. In addition, we can simulate transparency, and objects can be displayed using either flat surface rendering or Gouraud surface rendering.

**OpenGL Point Light-Source Function**

Multiple point light sources can be included in an OpenGL scene description, and various properties, such as position, type, color, attenuation, and spotlight effects,
are associated with each light source. We set a property value for a light source with the function

\[ \text{glLight* (lightName, lightProperty, propertyValue);} \]

A suffix code of i or f is appended to the function name, depending on the data type of the property value. For vector data, the suffix code v is also appended and parameter \text{propertyValue} is then a pointer to an array. Each light source is referenced with an identifier, and parameter \text{lightName} is assigned one of the OpenGL symbolic identifiers \text{GL_LIGHT0}, \text{GL_LIGHT1}, \text{GL_LIGHT2}, \ldots, \text{GL_LIGHT7}, although some implementations of OpenGL may allow more than 8 light sources. Similarly, parameter \text{lightProperty} must be assigned one of the 10 OpenGL symbolic property constants. After all properties have been assigned to a light source, we turn on that light with the command

\[ \text{glEnable (lightName);} \]

However, we also need to activate the OpenGL lighting routines, and we do that with the command

\[ \text{glEnable (GL_LIGHTING);} \]

Object surfaces are then rendered using lighting calculations that include contributions from each light source that has been enabled.

**Specifying an OpenGL Light-Source Position and Type**

The OpenGL symbolic property constant for designating a light-source position is \text{GL_POSITION}. Actually, this symbolic constant is used to set two light-source properties at the same time: the light-source position and the \text{light-source type}. Two general classifications of light sources are available in OpenGL to illuminate a scene. A point light source can be classified as near the objects to be illuminated (a local source), or it can be treated as if it were infinitely far from the scene. And this classification is independent of the position that we assign to a light source. For a nearby light source, the emitted light radiates in all directions, and the position of the light source is included in the lighting calculations. However, the emitted light from a distant source is allowed to emanate in one direction only, and this direction is applied to all surfaces in the scene, independently of the assigned light-source position. The direction for the emitted rays from a light that is classified as a distant source is calculated as the direction from the assigned position of the light source to the coordinate origin.

A four-element floating-point vector is used to designate both the type of light and the coordinate values for the light position. The first three elements of this vector give the world-coordinate position, and the fourth element is used to designate the light-source type. If we assign the value 0.0 to the fourth element of the position vector, the light is considered to be a very distant source (referred to in OpenGL as a “directional” light), and the light-source position is then used only to determine the light direction. Otherwise, the light is taken to be a local point source (referred to in OpenGL as a “positional” light), and the light position is used by the lighting routines to determine the light direction to each object in the scene. In the following code example, light 1 is designated as a local source
at location (2.0, 0.0, 3.0), and light 2 is a distant source with light emission in the
negative $y$ direction:

```c
GLfloat light1PosType[] = {2.0, 0.0, 3.0, 1.0};
GLfloat light2PosType[] = {0.0, 1.0, 0.0, 0.0};

glLightfv(GL_LIGHT1, GL_POSITION, light1PosType);
glEnable(GL_LIGHT1);

glLightfv(GL_LIGHT2, GL_POSITION, light2PosType);
glEnable(GL_LIGHT2);
```

If we do not specify a position and type for a light source, the default values are
(0.0, 0.0, 1.0, 0.0), which indicates a distant source with light rays traveling in the
negative $z$ direction.

The position of a light source is included in the scene description, and it is
transformed to viewing coordinates along with the object positions by the
OpenGL geometric-transformation and viewing-transformation matrices. Therefore,
if we want to keep the light source at a fixed position relative to the objects
in a scene, we set its position after the specification of the geometric and viewing
transformations in the program. However, if we want the light source to move
as the viewpoint moves, we set its position before the specification of the viewing
transformation. Also, we can apply a translation or rotation to a light source to
move it around in a stationary scene.

### Specifying OpenGL Light-Source Colors

Unlike an actual light source, an OpenGL light has three different color proper-
ties. In this empirical scheme, the three light-source colors provide options for
varying the lighting effects in a scene. We set these colors using the symbolic
color-property constants `GL_AMBIENT`, `GL_DIFFUSE`, and `GL_SPECULAR`. Each
of these colors is assigned by specifying a four-element floating-point set of val-
ues representing the red, green, blue, and alpha (RGBA) components of the color,
specified in that order. The alpha component controls color-blending, and is used
only if the OpenGL color-blending routines are activated. As we might guess
from the names of the symbolic color-property constants, one of the light-source
colors contributes to the background (ambient) light in a scene, another color
is used in diffuse-lighting calculations, and the third color is used to compute
specular-lighting effects for a surface. Realistically, a light source has just one
color, but we can use the three OpenGL light-source colors to create various light-
ing effects. In the following code example, we set the ambient color for a local light
source, labeled `GL_LIGHT3`, to black, and we set the diffuse and specular colors to
white:

```c
GLfloat blackColor[] = {0.0, 0.0, 0.0, 1.0};
GLfloat whiteColor[] = {1.0, 1.0, 1.0, 1.0};

glLightfv(GL_LIGHT3, GL_AMBIENT, blackColor);
gllightfv(GL_LIGHT3, GL_DIFFUSE, whiteColor);
gllightfv(GL_LIGHT3, GL_SPECULAR, whiteColor);
```

The default colors for light source 0 are black for the ambient color and white for
the diffuse and specular colors. All the other light sources have a default color of
black for each of the ambient, diffuse, and specular color properties.
Specifying Radial-Intensity Attenuation Coefficients for an OpenGL Light Source

We can apply radial-intensity attenuation to the light emitted from an OpenGL local light source, and the OpenGL lighting routines calculate this attenuation using Equation 2, with \( d \) as the distance from a light-source position to an object position. The three OpenGL property constants for radial intensity attenuation are \text{GL\_CONSTANT\_ATTENUATION}, \text{GL\_LINEAR\_ATTENUATION}, and \text{GL\_QUADRATIC\_ATTENUATION}, which correspond to the coefficients \( a_0 \), \( a_1 \), and \( a_2 \) in Equation 2. Either a positive integer value or a positive floating-point value can be used to set each attenuation coefficient. For example, we could assign the radial-attenuation coefficient values as

\[
\begin{align*}
\text{glLightf (GL\_LIGHT6, GL\_CONSTANT\_ATTENUATION, 1.5);}
\text{glLightf (GL\_LIGHT6, GL\_LINEAR\_ATTENUATION, 0.75);}
\text{glLightf (GL\_LIGHT6, GL\_QUADRATIC\_ATTENUATION, 0.4);}
\end{align*}
\]

Once the values for the attenuation coefficients have been set, the radial attenuation function is applied to all three colors (ambient, diffuse, and specular) of the light source. Default values for the attenuation coefficients are \( a_0 = 1.0, a_1 = 0.0, \) and \( a_2 = 0.0 \). Thus, the default is no radial attenuation: \( f_{\text{radatten}} = 1.0 \). Although radial attenuation can produce more realistic displays, the calculations are time-consuming.

OpenGL Directional Light Sources (Spotlights)

For local light sources (those not considered to be at infinity), we can also specify a directional, or spotlight, effect. This limits the light that is emitted from a source to a cone-shaped region of space. We define the conical region with a direction vector along the axis of the cone and an angular spread \( \theta_l \) from the cone axis, as shown in Figure 39. In addition, we can specify an angular-attenuation exponent \( a_l \) for the light source that determines how much the light intensity decreases as we move from the center of the cone toward the cone surface. Along any direction within the light cone, the angular attenuation factor is \( \cos^a \alpha \) (Eq. 5), where \( \cos \alpha \) is calculated as the dot product of the cone axis vector and the vector from the light source to an object position. We compute the value for each of the ambient, diffuse, and specular light colors at angle \( \alpha \) by multiplying the intensity components by this angular attenuation factor. If \( \alpha > \theta_l \), the object is outside the light-source cone, and the object is not illuminated by this light source. For light rays within the cone, we can also attenuate the intensity values radially.

There are three OpenGL property constants for directional effects: \text{GL\_SPOT\_DIRECTION}, \text{GL\_SPOT\_CUTOFF}, and \text{GL\_SPOT\_EXPONENT}. We specify the light direction as either an integer or a floating-point world-coordinate vector. The cone angle \( \theta_l \) is given as an integer or floating-point value in degrees, and this angle can be either 180° or any value in the range from 0° to 90°. When the cone angle is set to 180°, the light source emits rays in all directions (360°). We set the exponent value for intensity attenuation either as an integer or floating-point number in the range from 0 to 128. The following statements set the directional effects for light source 3 so that the cone axis is in the positive x direction, the cone angle \( \theta_l \) is 30°, and the attenuation exponent is 2.5:

\[
\begin{align*}
\text{GLfloat dirVector [ ] = \{1.0, 0.0, 0.0\};}
\text{glLightfv (GL\_LIGHT3, GL\_SPOT\_DIRECTION, dirVector);}
\text{glLightf (GL\_LIGHT3, GL\_SPOT\_CUTOFF, 30.0);}
\text{glLightf (GL\_LIGHT3, GL\_SPOT\_EXPONENT, 2.5);}
\end{align*}
\]
FIGURE 39
A circular cone of light emitted from an OpenGL light source. The angular extent of the light cone, measured from the cone axis, is $\theta$, and the angle from the axis to an object direction vector is labeled as $\alpha$.

If we do not specify a direction for a light source, the default direction is parallel to the negative $z$ axis; that is, $(0.0, 0.0, -1.0)$. Also, the default cone angle is $180^\circ$ and the default attenuation exponent is 0. Thus, the default is a point light source that radiates in all directions, with no angular attenuation.

OpenGL Global Lighting Parameters

Several OpenGL lighting parameters can be specified at the global level. These values are used to control the way that some lighting calculations are performed, and a global parameter value is set with the following function:

```c
glLightModel* (paramName, paramValue);
```

We append a suffix code of $i$ or $f$, depending on the data type of the parameter value. In addition, for vector data, we append the suffix code $v$. Parameter `paramName` is assigned an OpenGL symbolic constant that identifies the global property to be set, and parameter `paramValue` is assigned a single value or set of values. Using the `glLightModel` function, we can set a global ambient-light level, we can specify how specular highlights are to be calculated, and we can choose to apply the illumination model to the back faces of polygon surfaces.

In addition to the ambient color for individual light sources, we can set an independent value for the OpenGL background lighting as a global value. This provides just one more option in the empirical lighting calculations. To set this option, we use the symbolic constant `GL_LIGHT_MODEL_AMBIENT`. The following statement, for example, sets the general background lighting for a scene to a low-intensity (dark) blue color, with an alpha value of 1.0:

```c
globalAmbient [ ] = {0.0, 0.0, 0.3, 1.0};

glLightModelfv (GL_LIGHT_MODEL_AMBIENT, globalAmbient);
```
If we do not set a global ambient-light level, the default is the low-intensity white (dark gray) color (0.2, 0.2, 0.2, 1.0).

Specular-reflection calculations require the determination of several vectors, including the vector \( \mathbf{V} \) from a surface position to the viewing position. To speed up specular calculations, the OpenGL lighting routines can use a constant direction for vector \( \mathbf{V} \), regardless of the surface position relative to the view point. This constant unit vector is in the positive \( z \) direction, (0.0, 0.0, 1.0), and this is the default value for \( \mathbf{V} \). However, if we want to turn off this default and use the actual viewing position (which is the viewing-coordinate origin) to calculate \( \mathbf{V} \), we issue the following command:

\[
\text{glLightModeli (GL_LIGHTMODEL_LOCALVIEWER, GL_TRUE);}
\]

Although the specular calculations take more time when we use the actual viewing position to calculate \( \mathbf{V} \), we do obtain more realistic displays. We turn off the surface calculations for vector \( \mathbf{V} \) when we use the default value GL_FALSE (or 0, or 0.0) for the local-viewer parameter.

When surface textures are added to the OpenGL lighting calculations, surface highlights can be dulled and the texture patterns may be distorted by the specular terms. Therefore, as an option, texture patterns can be applied only to the nonspecular terms that contribute to a surface color. These nonspecular terms include ambient effects, surface emissions, and diffuse reflections. Using this option, the OpenGL lighting routines generate two colors for each surface lighting calculation: a specular color and the nonspecular color contributions. Texture patterns are combined only with the nonspecular color, and then the two colors are combined. We select this two-color option with

\[
\text{glLightModeli (GL_LIGHTMODEL_COLORCONTROL,}
\text{ GL_SEPARATE_SPECULAR_COLOR);}
\]

We need not separate the color terms if we are not using texture patterns, and the lighting calculations are performed more efficiently if this option is not invoked. The default value for this property is GL_SINGLE_COLOR, which does not separate the specular color from the other surface-color components.

In some applications, we may want to display back-facing surfaces of an object. An example is the inside, cutaway view of a solid, in which some back-facing surfaces, in addition to the front-facing surfaces, are to be displayed. However, by default, the lighting calculations use the assigned material properties only for the front faces. To apply the lighting calculations to both the front and back faces, using the corresponding front-face and back-face material properties, we issue the command

\[
\text{glLightModeli (GL_LIGHTMODEL_TWOSIDE, GL_TRUE);}
\]

The surface normal vectors for the back faces are then reversed, and the lighting calculations are applied using the material properties that have been assigned to the back faces. To turn off the two-sided lighting calculations, we use the value GL_FALSE (or 0, or 0.0) in the glLightModel function, which is the default.

**OpenGL Surface-Property Function**

Reflection coefficients and other optical properties for surfaces are set using the function

\[
\text{glMaterial* (surfFace, surfProperty, propertyValue);}\]
A suffix code of $i$ or $f$ is appended to the function, depending on the data type for the property value, and we also append the code $v$ when we supply vector-valued properties. Parameter `surfFace` is assigned one of the symbolic constants `GL_FRONT`, `GL_BACK`, or `GL_FRONT_AND_BACK`; parameter `surfProperty` is a symbolic constant identifying a surface parameter such as $I_{surf}$, $k_a$, $k_d$, $k_s$, or $n_s$; and parameter `propertyValue` is set to the corresponding value. All properties except the specular-reflection exponent $n_s$ are specified as vector values. We use a sequence of `glMaterial` functions to set all the illumination properties for an object before we issue the commands that describe the object geometry.

An RGBA value for the surface emission color, $I_{surf}$, is selected using the OpenGL symbolic surface-property constant `GL_EMISSION`. For example, the following statement sets the emission color for front surfaces to a light gray (off-white):

$$\text{surfEmissionColor} = \{0.8, 0.8, 0.8, 1.0\};$$

$$\text{glMaterialfv} (\text{GL_FRONT}, \text{GL_EMISSION}, \text{surfEmissionColor});$$

The default emission color for a surface is black (0.0, 0.0, 0.0, 1.0). Although an emission color can be assigned to a surface, this emission does not illuminate other objects in the scene. To do that, we must define the surface as a light source using the methods discussed in Section 3.

We use the OpenGL symbolic property names `GL_AMBIENT`, `GL_DIFFUSE`, and `GL_SPECULAR` to set values for the surface reflection coefficients. Realistically, the ambient and diffuse coefficients should be assigned the same vector values, and we can do that using the symbolic constant `GL_AMBIENT_AND_DIFFUSE`. The default values for the ambient coefficient are (0.2, 0.2, 0.2, 1.0), the default values for the diffuse coefficient are (0.8, 0.8, 0.8, 1.0), and the default values for the specular coefficient are (1.0, 1.0, 1.0, 1.0). To set the specular-reflection exponent, we use the constant `GL_SHININESS`. We can assign any value in the range from 0 to 128 to this property, and the default value is 0. For example, the following statements set the values for the three reflection coefficients and the specular exponent. The diffuse and ambient coefficients are set so that the surface is displayed as a light-blue color when it is illuminated with white light; specular reflection is the color of the incident light; and the specular exponent is assigned a value of 25.0.

$$\text{diffuseCoeff} = \{0.2, 0.4, 0.9, 1.0\};$$

$$\text{specularCoeff} = \{1.0, 1.0, 1.0, 1.0\};$$

$$\text{glMaterialfv} (\text{GL_FRONT_AND_BACK}, \text{GL_AMBIENT_AND_DIFFUSE}, \text{diffuseCoeff});$$

$$\text{glMaterialfv} (\text{GL_FRONT_AND_BACK}, \text{GL_SPECULAR}, \text{specularCoeff});$$

$$\text{glMaterialf} (\text{GL_FRONT_AND_BACK}, \text{GL_SHININESS}, 25.0);$$

Components for the reflection coefficients can also be set using color-table values, and the OpenGL symbolic constant `GL_COLOR_INDEXES` is provided for this purpose. We assign the color-table indices as a three-element integer or floating-point array, and the default is (0, 1, 1).

**OpenGL Illumination Model**

Surface lighting effects are calculated by OpenGL using the basic illumination model 19, with some variations in the way that the parameters are
specified. The ambient light level is the sum of the light-source ambient components and the global ambient setting. Diffuse-reflection calculations use the diffuse-intensity component of the light sources, and specular-reflection calculations use the specular-intensity component of each light source.

Also, the unit vector \( \mathbf{V} \), specifying the direction from a surface position to a viewing position, can be set to the constant value (0, 0, 0) if the local-viewer option is not used. For a light source positioned at “infinity,” the unit light-direction vector \( \mathbf{L} \) is in the opposite direction to the assigned direction for the light rays from that source.

OpenGL Atmospheric Effects

After the OpenGL illumination model has been applied to obtain surface colors, we can assign a color to the atmosphere in a scene and combine the surface colors with the atmosphere color. Also, we can use an atmosphere intensity-attenuation function to simulate viewing the scene through a hazy or smoky atmosphere. The various atmosphere parameters are set using the `glFog` function:

```c
glEnable (GL_FOG);

glFog* (atmoParameter, paramValue);
```

A suffix code of `i` or `f` is appended to indicate data-value type, and the suffix code `v` is used with vector data.

To set an atmosphere color, we assign the OpenGL symbolic constant `GL_FOG_COLOR` to parameter `atmoParameter`. For example, we can designate the atmosphere as having a bluish-gray color with

```c
GLfloat atmoColor[4] = {0.8, 0.8, 1.0, 1.0};

glFogfv (GL_FOG_COLOR, atmoColor);
```

The default value for the atmosphere color is black (0.0, 0.0, 0.0, 0.0).

We can next choose the atmosphere-attenuation function that is to be used to combine the object color with the atmosphere color. This is accomplished using the symbolic constant `GL_FOG_MODE`:

```c
glFogi (GL_FOG_MODE, atmoAttenFunc);
```

If parameter `atmoAttenFunc` is assigned the value `GL_EXP`, Equation 31 is used as the atmosphere-attenuation function. With the value `GL_EXP2`, we select Equation 32 as the atmosphere-attenuation function. For either of the exponential functions, we select an atmosphere density value with

```c
glFog (GL_FOG_DENSITY, atmoDensity);
```

A third option for atmospheric attenuation is the linear depth-cueing function. In this case, parameter `atmoAttenFunc` is assigned the value `GL_LINEAR`. The default value for parameter `atmoAttenFunc` is `GL_EXP`.

Once an atmosphere-attenuation function has been selected, this function is used to calculate a blended atmosphere-surface color for the object. Equation 33 is used by the OpenGL atmosphere routines to calculate this blended color.
OpenGL Transparency Functions

Some simulated transparency effects are possible in OpenGL using color-blending routines. However, the implementation of transparency in an OpenGL program, in general, is not straightforward. We can combine object colors for a simple scene containing a few opaque and transparent surfaces by using the alpha blending value to specify the degree of transparency and by processing surfaces in a depth-first order. The OpenGL color-blending operations ignore refraction effects, however, and dealing with transparent surfaces in complex scenes with a variety of lighting conditions or animations can be formidable. Also, OpenGL provides no direct provisions for simulating the surface appearance of a translucent object (such as a grainy sheet of plastic or a pane of frosted glass), which diffusely scatters the light transmissions through the semitransparent material. Thus, to display translucent surfaces or the lighting effects resulting from refraction, we would need to write our own routines. To simulate lighting effects through a translucent object, we could use a combination of values for surface texture and material properties. For refraction effects, we could shift the pixel positions for surfaces behind a transparent object using Equation 29 to calculate the amount of offset needed.

We designate objects in a scene as transparent using the alpha parameter in the OpenGL RGBA surface-color commands such as glMaterial and glColor. A surface alpha parameter can be set to the value of the transparency coefficient (Eq. 30) for that object. For example, if we specify the color for a transparent surface with the function

```
setColor4f(R, G, B, A);
```

then we set the alpha parameter to the value $A = k_t$. A completely transparent surface is assigned the alpha value $A = 1.0$, and an opaque surface has the alpha value $A = 0.0$.

Once we have assigned the transparency values, we activate the color-blending features of OpenGL and process the surfaces, starting with the most distant objects and proceeding in order to the objects closest to the viewing position. With color blending activated, each surface color is combined with any overlapping surfaces that are already in the frame buffer, using the assigned surface alpha values.

We set the color-blending factors so that all color components of the current surface (the “source” object) are multiplied by $(1 - A) = (1 - k_t)$, and all color components of the corresponding frame-buffer positions (the “destination”) are multiplied by the factor $A = k_t$:

```
setEnabled (GL_BLEND);

blendFunc (GL_ONE_MINUS_SRC_ALPHA, GL_SRC_ALPHA);
```

The two colors are then blended using Equation 30 with the alpha parameter set to $k_t$, where the frame-buffer colors are those for a surface that is behind the transparent object being processed. For instance, if $A = 0.3$, then the new frame-buffer color is the sum of 30 percent of the current frame-buffer color and 70 percent of the object reflection color, for each surface position. (Alternatively, we could use the alpha color parameter as an opacity factor, instead of a transparency factor. If we set $A$ to an opacity value, though, we also must interchange the two arguments in the function glColor.)

Visibility testing can be accomplished using the OpenGL depth-buffer functions. As each visible opaque surface is processed, both the surface colors and the
surface depth values are stored. However, when we process a visible transparent surface, we want to save only its colors because the surface does not obscure background surfaces. Therefore, as we process each transparent surface, we put the depth buffer into a read-only status using the `glDepthMask` function.

If we process all objects in depth order, the depth-buffer write mode is turned off and then back on again as we process each transparent surface. Alternatively, we could separate the two object classes, as in the following code outline:

```c
glEnable (GL_DEPTH_TEST);
/* Process all opaque surfaces. */

glEnable (GL_BLEND);
glDepthMask (GL_FALSE);
glBlendFunc (GL_ONE_MINUS_SRC_ALPHA, GL_SRC_ALPHA);
/* Process all transparent surfaces. */

glDepthMask (GL_TRUE);
glDisable (GL_BLEND);

glutSwapBuffers ( );
```

If the transparent objects are not processed in a strictly back-to-front order, this approach will not accumulate surface colors accurately for all cases. But for simple scenes, this is a fast and effective method for generating an approximate representation for the transparency effects.

### OpenGL Surface-Rendering Functions

Surfaces can be displayed with OpenGL routines using either constant-intensity surface rendering or Gouraud surface rendering. No OpenGL routines are provided for applying Phong surface rendering, ray tracing, or radiosity methods. A rendering method is selected with

```c
glShadeModel (surfRenderingMethod);
```

We select constant-intensity surface rendering by assigning the symbolic value `GL_FLAT` to parameter `surfRenderingMethod`. For Gouraud shading (the default), we use the symbolic constant `GL_SMOOTH`.

When the `glShadeModel` function is applied to a tessellated curved surface, such as a sphere that is approximated with a polygon mesh, the OpenGL rendering routines use the surface-normal vectors at the polygon vertices to calculate the polygon color. The Cartesian components of a surface-normal vector in OpenGL are specified with the command

```c
glNormal3* (Nx, Ny, Nz);
```

Suffix codes for this function are b (byte), s (short), i (integer), f (float), and d (double). In addition, we append the suffix code v when the vector components are designated with an array. Byte, short, and integer values are converted to floating-point values in the range from $-1.0$ to $1.0$. The `glNormal1` function sets the components for the surface-normal vector as state values that apply to all subsequent `glVertex` commands, and the default normal vector is in the positive z direction: $(0.0, 0.0, 1.0)$. 
For flat surface rendering, we need only one surface normal for each polygon. Thus, we can set each polygon normal as, for example,

```c
GLfloat normalVector[3];

glNormal3fv (normalVector);
```

```c
glBegin (GL_TRIANGLES);
    glVertex3fv (vertex1);
    glVertex3fv (vertex2);
    glVertex3fv (vertex3);
glEnd ( );
```

If we want to apply the Gouraud surface-rendering procedure to the above triangle, we need to designate a normal vector for each vertex as follows:

```c
glBegin (GL_TRIANGLES);
    glNormal3fv (normalVector1);
    glVertex3fv (vertex1);
    glNormal3fv (normalVector2);
    glVertex3fv (vertex2);
    glNormal3fv (normalVector3);
    glVertex3fv (vertex3);
glEnd ( );
```

Although normal vectors need not be specified as unit vectors, we can reduce computations if do state all surface normals as unit vectors. Any non-unit surface normal is converted to a unit normal automatically if we have issued the command

```c
glEnable (GL_NORMALIZE);
```

This command also renormalizes surface vectors if they have been modified by geometric transformations such as scaling or shear.

Another available option is the designation of a list of normal vectors that are to be combined or associated with a vertex array. The statements for creating an array of normal vectors are

```c
glEnableClientState (GL_NORMAL_ARRAY);

glNormalPointer (dataType, offset, normalArray);
```

Parameter `dataType` is assigned the constant value `GL_BYTE`, `GL_SHORT`, `GL_INT`, `GL_FLOAT` (the default value), or `GL_DOUBLE`. The number of bytes between successive normal vectors in the array `normalArray` is given by parameter `offset`, which has a default value of 0.

**OpenGL Halftoning Operations**

A variety of colors and gray-scale effects are possible on some systems using OpenGL halftone routines. The halftone-approximation patterns and operations are hardware dependent, and they typically have no effect on systems with full-color graphics capabilities. However, when a system has only a small number of bits per pixel, RGBA color settings can be approximated with halftone patterns. We activate the halftone routines with

```c
glEnable (GL_DITHER);
```

which is the default, and the halftoning routines are deactivated with the function

```c
glDisable (GL_DITHER);
```
12 Summary

In general, an object is illuminated with radiant energy from light emitters and from the reflective surfaces in a scene. Light sources can be modeled as point objects or they can have an extended size. In addition, light sources can be directional, and they can be treated as infinitely distant sources or as local light sources. Radial attenuation is typically applied to transmitted light using an inverse quadratic function of distance, and spotlights can be angularly attenuated as well. Reflecting surfaces in a scene are opaque, completely transparent, or partially transparent; and lighting effects are described in terms of diffuse and specular components for both reflections and refractions.

Light intensity at a surface position is calculated using an illumination model, and the basic illumination model in most graphics packages uses simplified approximations of physical laws. These lighting calculations provide a light-intensity value for each RGB component of the reflected light from a surface position, and for the transmitted light through a transparent object. The basic illumination model typically accommodates multiple light sources as point emitters, but they can be distant sources, local sources, or spotlights. Ambient light for a scene is described with a fixed intensity for each RGB color component and for all surfaces. Diffuse-intensity reflections from a surface are taken to be proportional to the cosine of the angular distance from the direction of the surface normal. Specular-intensity reflections are computed using the Phong model. In addition, transparency effects are usually approximated using a simple transparency coefficient for a material, although accurate refraction effects can be modeled using Snell's law. Shadow effects from the individual light sources can be added by identifying the regions in a scene that are not visible from the light source. Also, the calculations necessary for obtaining light reflections and transmission effects for translucent materials are not usually part of a basic illumination model, but we can model them using methods that disperse the diffuse light components.

Intensity values calculated with an illumination model are mapped to the intensity levels available on the display system in use. A logarithmic intensity scale is used by systems to provide a set of intensity levels that increase with equal perceived brightness differentials. Gamma correction is applied to intensity values to correct for the nonlinearity of display devices. With bilevel monitors, we can use halftone patterns and dithering techniques to simulate a range of intensity values. Halftone approximations can also be used to increase the number of intensity options on systems that are capable of displaying more than two intensities per pixel. Ordered-dither, error-diffusion, and dot-diffusion methods are used to simulate a range of intensities when the number of points to be plotted in a scene is equal to the number of pixels on the display device.

Surface rendering in graphics packages is accomplished by applying the calculations from the basic illumination model to scan-line procedures that extrapolate the intensity values from a few surface points to all projected pixel positions of a surface. With constant-intensity surface rendering, also called flat rendering, we use one calculated color to display all points of a surface. Flat surface rendering is accurate for polyhedrons or curved-surface polygon meshes when the viewing and light-source positions are far from the objects in a scene. Gouraud surface rendering approximates light reflections from tessellated curved surfaces by calculating intensity values at polygon vertices and linearly interpolating these intensity values across the polygon facets. A more accurate, but slower, surface-rendering procedure is Phong surface rendering, which interpolates the average normal vectors for polygon vertices over the polygon facets. Then, the basic
illumination model is employed to compute surface intensities at each projected surface position, using the interpolated values for the surface normal vectors. Fast Phong surface rendering uses Taylor series approximations to reduce processing time for the intensity calculations.

The core library of OpenGL contains an extensive set of functions for setting up point light sources, specifying the various parameters in the basic illumination model, selecting a surface-rendering method, activating halftone-approximation routines, and for applying texture array patterns to objects. Table 2 provides a summary of these OpenGL illumination and surface-rendering functions.

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>glLight</code></td>
<td>Specifies a light-source property value.</td>
</tr>
<tr>
<td><code>glEnable (lightName)</code></td>
<td>Activates a light source.</td>
</tr>
<tr>
<td><code>glLightModel</code></td>
<td>Specifies global-lighting parameter values.</td>
</tr>
<tr>
<td><code>glMaterial</code></td>
<td>Specifies a value for an optical surface parameter.</td>
</tr>
<tr>
<td><code>glFog</code></td>
<td>Specifies a value for an atmosphere parameter; activates atmospheric effects with the <code>glEnable</code> function.</td>
</tr>
<tr>
<td><code>glColor4f (R, G, B, A)</code></td>
<td>Specifies an alpha value for a surface to simulate transparency. In the function <code>glBlendFunc</code>, sets the source blending factor to <code>GL_SRC_ALPHA</code> and the destination blending factor to <code>GL_ONE_MINUS_SRC_ALPHA</code>.</td>
</tr>
<tr>
<td><code>glShadeModel</code></td>
<td>Specifies either Gouraud surface rendering or single-color surface rendering.</td>
</tr>
<tr>
<td><code>glNormal3</code></td>
<td>Specifies a surface-normal vector.</td>
</tr>
<tr>
<td><code>glEnable (GL_NORMALIZE)</code></td>
<td>Specifies that surface normals are to be converted to unit vectors.</td>
</tr>
<tr>
<td><code>glEnableClientState</code></td>
<td>Activates processing routines for an array of surface-normal vectors.</td>
</tr>
<tr>
<td><code>glNormalPointer</code></td>
<td>Creates a list of surface-normal vectors that are to be used with a vertex array.</td>
</tr>
<tr>
<td><code>glEnable (GL_DITHER)</code></td>
<td>Activates operations for applying surface rendering as halftone approximation patterns.</td>
</tr>
</tbody>
</table>
REFERENCES

Additional programming examples using OpenGL illumination and rendering functions are given in Woo, et al. (1999). Programming examples for the OpenGL lighting and rendering functions are also available at Nate Robins’s tutorial website: http://www.xmission.com/~nate/opengl.html. Finally, a complete listing of OpenGL illumination and rendering functions is provided in Shreiner (2000).

EXERCISES
1 Write a routine to implement Equation 12 for diffuse reflection using a single point light source and constant surface rendering for the faces of a tetrahedron. The object description is to be given in polygon tables, including surface normal vectors for each of the polygon faces. Additional input includes values for the ambient intensity, light-source intensity, surface reflection coefficients, and specular-reflection parameter. All coordinate information can be specified directly in the viewing reference frame.

2 Modify the routine in preceding exercise to display the polygon facets using Phong surface rendering.

3 Modify the routine in the preceding exercise to include a linear intensity attenuation function.

4 Modify the routine in the preceding exercise to include two light sources in the scene.

5 Discuss the differences you might expect to see in the appearance of specular reflections modeled with \((\mathbf{N} \cdot \mathbf{H})^n\) compared to specular reflections modeled with \((\mathbf{V} \cdot \mathbf{R})^n\).

6 Verify that \(2\alpha = \phi\) in Figure 18 when all vectors are coplanar, but that, in general, \(2\alpha \neq \phi\).

7 Discuss how the different visible-surface detection methods can be combined with an intensity model for displaying a set of polyhedrons with opaque surfaces.

8 Discuss how the various visible-surface detection methods can be modified to process transparent objects. Are there any visible-surface detection methods that cannot handle transparent surfaces?

9 Set up an algorithm, based on one of the visible-surface detection methods, that will identify shadow areas in a scene illuminated by a distant point source.

10 How many intensity levels can be displayed with halftone approximations using \(n \times n\) pixel grids, where each pixel can be displayed with \(m\) different intensities?

11 How many different color combinations can be generated using halftone approximations on a four-level RGB system with a 3 x 3 pixel grid?

12 How many different color combinations can be generated using halftone approximations on a two-level RGB system with a 4 x 4 pixel grid?

13 Write a routine to display a given set of surface-intensity variations using halftone approximations with 4 x 4 pixel grids and two intensity levels (0 and 1) per pixel.

14 Write a routine to generate ordered-dither matrices using the recurrence relation in Equation 48.
21 Write a procedure to display a given array of intensity values using the ordered-dither method.
22 Write a procedure to implement the error-diffusion algorithm for a given \( m \times n \) array of intensity values.
23 Write an OpenGL program to display a scene containing a sphere, a cube, and a tetrahedron illuminated by two light sources: one is to be a local green source and the other a distant white-light source. Set surface parameters for both diffuse and specular reflections with Gouraud surface rendering, and apply a linear intensity-attenuation function.
24 Modify the program in the preceding exercise so that the single local green source is replaced with two spotlights: one green and one blue.
25 Modify the program in the preceding exercise so that a smoky atmosphere is added to the scene.
26 Modify the program in the preceding exercise so that the scene is viewed through a semitransparent pane of glass.

**IN MORE DEPTH**

1 Using the techniques presented in this chapter, choose a lighting scheme appropriate to your application and write out its specification in detail. Decide what types of light sources are most appropriate (point sources, directional sources, ambient sources, etc.) and what their colors, positions, and orientations should be, if applicable. In addition, include appropriate lighting effects, such as intensity attenuation, shadows, or atmospheric effects where appropriate. Next, choose appropriate surface properties for the objects in your scene based on the material that they represent and include these properties in your specification. If transparency is an important aspect in your scene, be sure to specify the transparency properties of the objects in your scene as well.

2 Implement the specification that you developed in the previous exercise using the OpenGL illumination and surface-rendering functions. Create and position/orient light sources accordingly within the scene and turn on any atmospheric effects if necessary. If you are using attenuation functions, then experiment with the different models and parameters that define their visual appearance. Next, set the material properties of the surfaces in your scene, including diffuse and specular reflection parameters, and turn on appropriate color-blending routines for transparent surfaces if applicable. Finally, render the scene using Gouraud shading and experiment with modifying each of the parameters discussed here to produce the most visually appealing result.
A wire-frame scene (a) is displayed in (b) using ambient lighting only, with a different color for each object. Diffuse reflections resulting from illumination with ambient light and a single point source are illustrated in (c). For this display, $k_s = 0$ for all surfaces. In (d), both diffuse and specular reflections are shown for the illumination from a point source and the background lighting.

Diffuse reflections from a spherical surface illuminated by a point source emitting white light, with values of the diffuse reflectivity coefficient in the interval $0 \leq k_d \leq 1$. 

Color Plate 14
Light reflections from the surface of a black nylon cushion, modeled as woven cloth patterns and rendered using Monte-Carlo ray-tracing methods. (Courtesy of Stephen H. Westin, Program of Computer Graphics, Cornell University.)

Color Plate 15
Light reflections from a teapot with reflectance parameters set to simulate brushed aluminum surfaces and rendered using Monte-Carlo ray-tracing methods. (Courtesy of Stephen H. Westin, Program of Computer Graphics, Cornell University.)

Color Plate 16
Light reflections from trumpets with reflectance parameters set to simulate shiny brass surfaces. (Courtesy of SOFTIMAGE, Inc.)

Color Plate 17
A polygon mesh approximation of an object (a) is displayed using flat surface rendering in (b) and using Gouraud surface rendering in (c).
So far, we have discussed rendering techniques for displaying smooth object surfaces. However, most objects do not have smooth, even surfaces. We need surface texture to model accurately such objects as brick walls, gravel roads, shag carpets, wood, and human skin. In addition, some surfaces contain patterns that must be taken into account in the rendering procedures. The surface of a vase could contain a painted design; a water glass might have the family crest engraved into the surface; a tennis court contains markings for the alleys, service areas, and baseline; and a four-lane highway has dividing lines and other markings, such as oil spills and tire skids.

Color Plate 18 illustrates the basic stages in modeling and rendering an object that is to contain surface detail. First, a wire-frame display of the object can be used to adjust the overall design. Next, surface layers are fitted over the object outline to produce a rendered, smooth-surface view of the structure. Finally, surface detail could be added to the frame in the image to simulate wood grain or a...
machined metal surface, or a texture could be “painted” onto the material within the
frame to simulate the ball passing through and tearing a picture. Additional examples of
scenes rendered with surface detail are given in Color Plate 19.

We can add detail to surfaces using a variety of methods, including the following:

- Pasting small objects, such as buds, flowers, or spines, onto a larger surface
- Modeling surface patterns with small polygon areas
- Mapping texture arrays or intensity-modifying procedures onto a surface
- Modifying the surface normal vector to create localized bumps
- Modifying both the surface normal vector and the surface tangent vector to
display directional patterns on wood and other materials

1 Modeling Surface Detail with Polygons

A simple method for adding surface detail is to model patterns or other surface
characteristics using polygon facets. For large-scale detail, polygon modeling can
give good results. Some examples of such large-scale detail are squares on a
checkerboard, dividing lines on a highway, tile patterns on a linoleum floor, floral
designs in a smooth low-pile rug, panels in a door, and lettering on the side of a
panel truck. Also, we could model an irregular surface with small, randomly
oriented polygon facets, provided the facets are not too small.

Surface-pattern polygons are generally overlaid on a larger surface polygon
and processed along with the parent surface. The visible-surface detection algo-
rithms process only the parent polygon, but the illumination parameters for the
surface-detail polygons take precedence over the parent polygon. When intricate
or fine surface detail is to be modeled, polygon methods are not practical. For
example, it would be difficult to model the surface structure of a raisin accurately
with polygon facets.

2 Texture Mapping

A common method for adding detail to an object is to map patterns onto the
geometric description of the object. The texture pattern may be defined either in
an array of color values or as a procedure that modifies object colors. This method
for incorporating object detail into a scene is called texture mapping or pattern
mapping, and the textures can be defined as one-dimensional, two-dimensional,
or three-dimensional patterns. Any texture specification is referred to as a texture
space, which is referenced with texture coordinates in the range from 0 to 1.0.

Texture functions in a graphics package often allow the number of color com-
ponents for each position in a pattern to be specified as an option. For example,
each color specification in a texture pattern could consist of four red, green, blue,
and alpha (RGBA) components, three RGB components, a single intensity value
for a shade of blue, an index into a color table, or a single luminance value (a
weighted average of the RGB components of a color). A component of a texture
description is frequently alluded to as a “texel,” but there is some confusion in the
use of the term. Sometimes a position in texture space corresponding to a set of
color components, such as an RGB triple, is called a texel, and sometimes a single
texture-array element, such as the value for the red component of an RGB color,
is also called a texel.
Linear Texture Patterns

A one-dimensional texture pattern can be specified in a single-subscript array of color values, which defines a sequence of colors in a linear texture space. For example, we could set up a list of 32 RGB colors, referenced with subscript values ranging from 0 to 95. The first three elements of the array store the RGB components of the first color, the next three elements store the RGB components of the second color, and so forth. This set of colors, or any contiguous subset of the colors, could then be used to form a patterned stripe across a polygon, a band around a cylinder, or a color pattern for displaying an isolated line segment.

For a linear pattern, the texture space is referenced with a single \( s \)-coordinate value. For RGB color specifications, the value \( s = 0.0 \) designates the first three-element RGB color in the array, the value \( s = 1.0 \) designates the last three RGB color components, and the value \( s = 0.5 \) references the middle three RGB color elements in the array. As an example, if the name of the texture array is \( \text{colorArray} \), then the value \( s = 0.0 \) references the three array values \( \text{colorArray}[0] \), \( \text{colorArray}[1] \), and \( \text{colorArray}[2] \).

To map a linear texture pattern into a scene, we assign an \( s \)-coordinate value to one spatial position and another \( s \)-coordinate value to a second spatial position. The section of the color array corresponding to the specified \( s \)-coordinate range is then used to generate a multicolored line between the two spatial positions. A texture-mapping procedure typically uses a linear function to calculate the array positions that are to be assigned to the pixels along a line segment. When the number of texture colors specified for the line is small, each color may be assigned to a large block of pixels, depending on the length of the line. For example, if the specified \( s \)-coordinate range spans a single RGB color (three RGB color elements) in the texture array, all pixels on the line are displayed in that color. But if many colors are to be mapped to the positions along the line, then fewer pixels are assigned to each color. Also, because some pixels may map to array positions between RGB colors, various schemes can be used to determine the color that is to be assigned to each pixel. A simple color-mapping method is to assign the nearest array color to each pixel. Alternatively, if a pixel is mapped to a position between the starting array elements for two colors, the pixel color could be computed as a linear combination of the nearest two color elements in the array.

Some texture-mapping procedures allow values for texture coordinates that are outside the range from 0 to 1.0. These situations might arise when we want to map multiple copies of a texture onto an object or when calculated \( s \) values could be outside the unit interval. If we want to allow texture-coordinate values outside the range from 0 to 1.0, we could just ignore the integer part of any \( s \) value. In this case, the value of \(-3.6\), for instance, would reference the same position in texture space as the value 0.6 or the value 12.6. However, if we do not want to allow values outside the range from 0 to 1.0, then we could just clamp values to this unit interval. Any computed value less than 0 is then reset to 0, and any computed value greater than 1.0 is reset to 1.0.

Surface Texture Patterns

A texture for a surface area is commonly defined with a rectangular color pattern, and positions in this texture space are referenced with two-dimensional \((s, t)\) coordinate values. Specifications for each color in the texture pattern can be stored in a three-subscript array. If a texture pattern is defined with \(16 \times 16\) RGB colors, for instance, then the array for this pattern contains \(16 \times 16 \times 3 = 768\) elements.

Figure 1 illustrates a two-dimensional texture space. Values for both \( s \) and \( t \) vary from 0 to 1.0. The first row of the array lists the color values across the
Two-dimensional texture-space coordinates that reference positions in an array of color values containing \(m\) rows and \(n\) columns. Each position in the array can reference multiple color components.

Bottom of the rectangular texture pattern, and the last row of the array lists the color values across the top of the pattern. Coordinate position \((0, 0)\) in texture space references the first set of color components in the first position of the first row, and position \((1.0, 1.0)\) references the last set of color components at the last position in the last row of the array. Of course, we could list the colors in the texture array in other ways. If we listed the colors in a top-to-bottom order, the origin of the two-dimensional texture space would be at the top-left corner of the rectangular pattern. But placing the texture-space origin at the lower-left corner usually simplifies the mapping procedures to the spatial-coordinate reference for a scene.

We specify a surface-texture mapping for an object using the same procedures we used for specifying a linear-texture mapping to a scene. The \((s, t)\) texture-space coordinates for the four corners of the texture pattern (Figure 1) can be assigned to four spatial positions in the scene, and a linear transformation is used to assign color values to the projected pixel positions over the designated spatial area. Other mappings are possible. For instance, we could assign three texture-space coordinates to the vertices of a triangle.

Surface positions on an object, such as a cubic-spline patch or a sphere section, can be described with \(uv\) object-space coordinates, and projected pixel positions are referenced in \(xy\) Cartesian coordinates. Surface texture mapping can be accomplished in one of two ways. Either we can map the texture pattern to an object surface, then to the projection plane; or we can map each pixel area onto the object surface, and then map this surface area to texture space. Mapping a texture pattern to pixel coordinates is sometimes called texture scanning, while the mapping from pixel coordinates to texture space is referred to as pixel-order scanning, inverse scanning, or image-order scanning. Figure 2 diagrams the two possible transformation sequences between the three spaces.

Parametric linear transformations provide a simple method for mapping positions in texture space to object space:

\[
\begin{align*}
    u &= u(s, t) = a_u s + b_u t + c_u \\
    v &= v(s, t) = a_v s + b_v t + c_v
\end{align*}
\]
The object-to-image-space transformation is accomplished with the concatenation of the viewing and projection transformations. A disadvantage of mapping from texture space to pixel space is that a selected texture patch usually does not match up with the pixel boundaries, which requires calculations to determine the fractional area of pixel coverage. Therefore, mapping from pixel space to texture space (Figure 3) is the most commonly used texture-mapping method. This avoids pixel subdivision calculations, and allows antialiasing (filtering) procedures to be easily applied. An effective antialiasing procedure is to project a slightly larger pixel area that includes the centers of neighboring pixels, as shown in Figure 4, and applying a pyramid function to weight the intensity values in the texture pattern. But the mapping from image space to texture space does require calculation of the inverse viewing-projection transformation $M_{VP}^{-1}$ and the inverse texture-map transformation $M_{T}^{-1}$. In the following example, we illustrate this approach by mapping a defined pattern onto a cylindrical surface.

**Example 1  Surface Texture Mapping**

To illustrate the steps in surface-texture mapping, we consider the transfer of the pattern shown in Figure 5(a) to a cylindrical surface. The surface parameters are the cylindrical coordinates

$$ u = \theta, \quad v = z $$

with

$$ 0 \leq \theta \leq \pi/2, \quad 0 \leq z \leq 1 $$
In addition, the parametric representation for the surface in the Cartesian reference frame is

\[ x = r \cos u, \quad y = r \sin u, \quad z = v \]

We can map the array pattern to the surface using the following linear transformation, which transforms the texture-space coordinates \((s, t) = (0, 0)\) to the lower-left corner of the surface element \((x, y, z) = (r, 0, 0)\).

\[ u = s\pi/2, \quad v = t \]

Next, we select a viewing position and perform the inverse viewing transformation from pixel coordinates to the Cartesian reference for the cylindrical surface. Then, Cartesian surface coordinates are transferred to \(uv\) surface parameters with the calculations

\[ u = \tan^{-1}(y/x), \quad v = z \]

and projected pixel positions are mapped to texture space with the inverse transformation

\[ s = 2u/\pi, \quad t = v \]

Color values in the pattern array covered by each projected pixel area are then averaged to obtain the pixel color.

**Volume Texture Patterns**

In addition to linear and surface patterns, we can designate a set of colors for positions throughout a three-dimensional region of space. These textures are often referred to as **volume texture patterns** or **solid textures**. We reference a solid texture using three-dimensional texture-space coordinates \((s, t, r)\). Also, the three-dimensional texture space is defined within the unit cube, with texture coordinates ranging from 0 to 1.0.

A volume texture pattern can be stored in a four-subscript array, where the first three subscripts denote a row position, a column position, and a depth position. The fourth subscript is used to reference a component of a particular color in the pattern. For example, an RGB solid texture pattern with 16 rows, 16 columns, and 16 depth planes could be stored in an array with \(16 \times 16 \times 16 \times 3 = 12,288\) elements.

To map the entire texture space to a three-dimensional block, we assign the coordinates for the eight corners of the texture space to eight spatial positions in a scene. Alternatively, we could map a plane section of texture space, such as a depth plane or one face of the texture cube, to a planar area in the scene. A variety of other solid-texture mapping applications are also possible.

Solid texturing allows internal views, such as cut-away displays and cross-sectional slices, for three-dimensional objects to be displayed with texture.
patterns. Thus bricks, cinder blocks, or wood materials can have the same texture patterns applied throughout the spatial extent of the objects. Color Plate 20 shows a scene displayed using solid textures to obtain wood grain and other material patterns.

**Texture Reduction Patterns**

In animations and other applications, the size of an object often changes. For objects displayed with texture patterns, we then need to apply the texture-mapping procedures to the altered dimensions of the object. When the size of a textured object is reduced, the texture pattern is applied to a smaller region and this can lead to texture distortions. To avoid this, we can create a series of **texture reduction patterns** that are to be used when the displayed size of the object is scaled down.

Typically, each reduction pattern is half the size of the previous pattern. For example, if we have a two-dimensional 16 × 16 pattern, then we could set up four additional patterns at the reduced sizes of 8 × 8, 4 × 4, 2 × 2, and 1 × 1. For any view of an object, we can then apply an appropriate reduction pattern to minimize distortions. These reduction patterns are often referred to as MIP maps or mip maps, where the term mip is an acronym for the Latin phrase *multum in parvo*, which can be translated as “much on a small object.”

**Procedural Texturing Methods**

Another method for adding a texture pattern to an object is to use a procedural definition for the color variations that are to be applied. This approach avoids the transformation calculations involved in mapping array patterns to object descriptions. And procedural texturing eliminates the storage requirements that are necessary when many large texture patterns, particularly solid textures, are to be applied to a scene.

We generate a procedural texture by calculating variations for the properties or characteristics of an object. Wood-grain or marble patterns, for example, can be created throughout an object using harmonic functions (sine curves) defined in a region of three-dimensional space. Random perturbations are then superimposed on the harmonic variations to break up the symmetric patterns.

### 3 Bump Mapping

Although texture arrays can be used to add fine surface detail, they usually are not effective for modeling the rough surface appearance of objects such as oranges, strawberries, and raisins. The light-intensity detail that is provided in a texture array for these objects is set independently of illumination parameters such as the light-source direction. A better method for modeling surface bumpiness is to apply a perturbation function to the surface normal and then use the perturbed normal vector in the illumination-model calculations. This technique is called **bump mapping**.

If \( P(u, v) \) represents a position on a parametric surface, we can obtain the surface normal at that point with the calculation

\[
N = P_u \times P_v
\]
where $P_u$ and $P_v$ are the partial derivatives of $P$ with respect to parameters $u$ and $v$. To provide variations in the surface normal, we can modify the surface position vector by adding a small perturbation function, called a bump function:

$$P'(u, v) = P(u, v) + b(u, v) \cdot n$$

(3)

This adds bumps to the surface in the direction of the unit surface normal $n = N/|N|$. The perturbed surface normal is then obtained as

$$N' = P'_u \times P'_v$$

(4)

The partial derivative of $P'$ with respect to $u$ is

$$P'_u = \frac{\partial}{\partial u} (P + b \cdot n)$$

$$= P_u + b_u \cdot n + b_n \cdot u$$

(5)

Assuming that the magnitude of the bump function $b$ is small, we can neglect the last term in this expression, so that

$$P'_u \approx P_u + b_u \cdot n$$

(6)

Similarly,

$$P'_v \approx P_v + b_v \cdot n$$

(7)

In addition, the perturbed surface normal is

$$N' = P_u \times P_v + b_v (P_u \times n) + b_u (n \times P_v) + b_u b_v (n \times n)$$

However, $n \times n = 0$, so that

$$N' = N + b_v (P_u \times n) + b_u (n \times P_v)$$

(8)

The final step is to normalize $N'$ for use in the illumination-model calculations.

There are several ways in which we can specify the bump function $b(u, v)$. We could set up an analytic expression, but computations are reduced if we simply obtain bump values using table lookups. With a bump table, values for $b$ are quickly determined using linear interpolation and incremental calculations. Then, the partial derivatives $b_u$ and $b_v$ are approximated with finite differences. The bump table can be set up with random patterns, regular grid patterns, or character shapes. Random patterns are useful for modeling an irregular surface, such as a raisin, while a repeating pattern could be used to model the surface of an orange, for example. To antialias, we subdivide pixel areas and average the computed subpixel intensities.

Color Plate 21 shows examples of surfaces rendered with bump mapping.

### 4 Frame Mapping

This method for adding surface detail is an extension of bump mapping. In frame mapping, we perturb both the surface normal vector $N$ and a local coordinate system (Figure 6) attached to $N$. The local coordinates are defined with a surface tangent vector $T$ and a binormal vector $B = T \times N$.

Frame mapping is used to model anisotropic surfaces. We orient $T$ along the “grain” of the surface and apply directional perturbations, in addition to bump perturbations in the direction of $N$. In this way, we can model wood-grain patterns, cross-thread patterns in cloth, and streaks in marble or similar materials. Both bump and directional perturbations can be generated using table lookups.
5 OpenGL Texture Functions

An extensive set of texture functions is available in OpenGL. We can specify a pattern for a line, a surface, an interior volume of a spatial region, or as a subpattern that is to be inserted into another texture pattern. And we can apply and manipulate texture patterns in various ways. In addition, texture patterns can be used to simulate environment mapping. The OpenGL texture routines can be used only in RGB or RGBA color mode, although some parameters can be set using a color-table index.

OpenGL Line-Texture Functions

Parameters for a one-dimensional RGBA texture pattern specified in a single-subscript color array are designated, for example, with

```c
glTexImage1D (GL_TEXTURE_1D, 0, GL_RGBA, nTexColors, 0,
            dataFormat, dataType, lineTexArray);
```

We have set the first argument in the `glTexImage1D` function to the OpenGL symbolic constant `GL_TEXTURE_1D` to indicate that we are defining a texture array for a one-dimensional object: a line. If we are not sure that the system will support the texture pattern with the specified parameters, we use the symbolic constant `GL_PROXY_TEXTURE_1D` for the first argument of `glTexImage1D`. This allows us to first query the system before defining the elements of the texture array, and we discuss the query procedures in a later section.

For the second and fifth arguments of this example function, we use the value 0. The first 0 value (second argument) means that this array is not a reduction of some larger texture array. For the fifth argument, the 0 value means that we do not want a border around the texture. If this fifth argument had been assigned the value 1 (the only other possibility), the texture pattern would be displayed with a one-pixel border around it, which is used to merge the pattern with adjacent texture patterns. For the third argument, the value `GL_RGBA` means that each color of the texture pattern is specified with four RGBA values. We could have just used the three RGB color values, but RGBA values are sometimes processed more efficiently because they align with processor memory boundaries. Numerous other color specifications are possible, including a single intensity or luminance value. Parameter `nTexColors`, the fourth argument, is to be assigned a positive integer indicating the number of colors in the linear texture pattern. Because a 0 value is listed for the fifth argument (the border parameter), the number of colors in the texture pattern must be a power of 2. If the fifth argument had been assigned the value 1, then the number of colors in the texture pattern would have to be 2 plus a power of 2. The two border colors are used to provide
color blending with neighboring patterns. We can specify the one-subscript texture pattern with up to 64 + 2 colors, and some OpenGL implementations allow larger texture patterns. Parameters describing the texture colors and the border colors are stored in lineTexArray. In this example, we have no border and each successive group of four elements in the array represents one color component of the texture pattern. Therefore, the number of elements in lineTexArray is $4 \times nTexColors$. As a specific example, if we want to define a texture pattern with 8 colors, the texture array must contain $4 \times 8 = 32$ elements.

Parameters dataFormat and dataType are similar to the arguments in the glDrawPixels and glReadPixels functions. We assign an OpenGL symbolic constant to dataFormat to indicate how the color values are to be specified in the texture array. For instance, we could use the symbolic constant GL_BGR to indicate that the color components are to be given in the order blue, green, red, alpha. To indicate the BGRA or RGBA data type, we can assign the OpenGL constant value GL_UNSIGNED_BYTE to parameter dataType. Other possible possible the data format we choose, include GL_INT and GL_FLOAT.

We can map multiple copies of a texture, or any contiguous subset of the texture colors, to an object in a scene. When a group of texture elements is mapped to one or more pixel areas, the boundaries of the texture elements usually do not align with the positions of the pixel boundaries. A pixel area could be contained within the boundaries of a single RGB (or RGBA) texture element or it could overlap several texture elements. To simplify the calculations in the texture mapping, we use the following functions to give each pixel the color of the nearest texture element:

```c
glTexParameteri (GL_TEXTURE_1D, GL_TEXTURE_MIN_FILTER, GL_NEAREST);
glTexParameteri (GL_TEXTURE_1D, GL_TEXTURE_MAG_FILTER, GL_NEAREST);
```

The first function is used by the texturing routines when a section of the texture pattern must be enlarged to fit a specified coordinate range in a scene, and the second function is used when a texture pattern has to be reduced. (These two texture operations in OpenGL are referred to as magnifying, MAG, and minifying, MIN.) Although assigning the nearest texture color to a pixel can be performed quickly, it can lead to aliasing effects. To calculate the pixel color as a linear combination of overlapping texture colors, we replace the symbolic constant GL_NEAREST with GL_LINEAR. Several other parameter values can be set with the glTexParameteri function, and we take a look at these options later in this chapter.

Specifying OpenGL texture patterns for a scene is somewhat similar to specifying surface-normal vectors, RGB colors, or other attributes. We need to associate the pattern with some object, but, unlike a single color setting, we now have a collection of color values. For a one-dimensional texture space, the color values are referenced with a single $s$ coordinate that varies from 0.0 to 1.0 across the texture space. Thus, the texture pattern is applied to objects in a scene by assigning texture coordinate values to object positions. A particular $s$-coordinate value in one-dimensional texture space is selected with the following command:

```c
glTexCoord1* (sCoord);
```

Allowable suffix codes for this function are b (byte), s (short), i (integer), f (float), and d (double), depending on the data type for the texture coordinate parameter sCoord. In addition, we can use the suffix v if an $s$-coordinate value is given in
an array. As with color and other similar parameters, the s coordinate is a state parameter, which applies to all subsequently defined world-coordinate positions. The default value for the s coordinate is 0.0.

To map a linear texture pattern onto positions within a world-coordinate scene, we assign s coordinates to the endpoints of a line segment. The texture colors can be then be applied to the object in various ways, and the OpenGL default method is to multiply each pixel color value for the object by the corresponding color value in the texture pattern. If the line color is white (1.0, 1.0, 1.0, 1.0), which is the default color for objects in a scene, the line will be displayed only with the texture colors.

In the following example, we create a four-element linear texture pattern with alternating green and red colors. The entire texture pattern, from 0.0 to 1.0, is then assigned to a straight-line segment. Because the line is white, by default, it is displayed in the texture colors.

```c
GLint k;
GLubyte texLine [16]; // 16-element texture array.

/* Define two green elements for the texture pattern. */
/* Each texture color is specified in four array positions. */
for (k = 0; k <= 2; k += 2) {
    texLine [4*k] = 0;
    texLine [4*k+1] = 255;
    texLine [4*k+2] = 0;
    texLine [4*k+3] = 255;
}

/* Define two red elements for the texture pattern. */
for (k = 1; k <= 3; k += 2) {
    texLine [4*k] = 255;
    texLine [4*k+1] = 0;
    texLine [4*k+2] = 0;
    texLine [4*k+3] = 255;
}

glTexParameteri (GL_TEXTURE_1D, GL_TEXTURE_MAG_FILTER, GL_NEAREST);
glTexParameteri (GL_TEXTURE_1D, GL_TEXTURE_MIN_FILTER, GL_NEAREST);
glTexImage1D (GL_TEXTURE_1D, 0, GL_RGBA, 4, 0, GL_RGBA, GL_UNSIGNED_BYTE, texLine);
glEnable (GL_TEXTURE_1D);

/* Assign the full range of texture colors to a line segment. */
 glBegin (GL_LINES);
    glVertex2f (0.0);
    glVertex2f (1.0);
 glEnd ( );

glDisable (GL_TEXTURE_1D);
```

Texturing and Surface Detail Methods
The line segment is displayed with alternate green and red sections along the line path. We can assign any values to the \( s \) coordinates. For example, the middle red and green colors of the texture pattern are mapped onto the line with the following statements:

\[
\begin{align*}
&\text{glBegin (GL_LINES);} \\
&\quad \text{glTexCoord1f (0.25);} \\
&\quad \text{glVertex3fv (wcPt1);} \\
&\quad \text{glTexCoord1f (0.75);} \\
&\quad \text{glVertex3fv (wcPt2);} \\
&\text{glEnd ( );}
\end{align*}
\]

Thus, the first half of the line is red and the second half green. We could also use \( s \) values outside the range from 0.0 to 1.0. For instance, if we assigned the \( s \) value \(-2.0\) to one endpoint of a line and the \( s \) value \( 2.0 \) to the other endpoint, the texture pattern would be mapped onto the line four times. The line would then be displayed with 16 green sections and 16 red sections. For \( s \)-coordinate values outside the unit interval, integer parts are ignored unless we specify that \( s \) values are to be clamped to 0 or 1.0.

A vast number of parameters and options are available with OpenGL texture patterns. However, before we go into these features of the OpenGL texture routines, we will discuss the basic functions needed to generate two-dimensional and three-dimensional texture patterns.

**OpenGL Surface-Texture Functions**

We can set the parameters for a two-dimensional RGBA texture space using functions similar to those for our one-dimensional texture pattern example:

\[
\begin{align*}
&\text{glTexImage2D (GL_TEXTURE_2D, 0, GL_RGBA, texWidth,} \\
&\quad \text{texHeight, 0, dataFormat, dataType, surfTexArray);} \\
&\text{glEnable (GL_TEXTURE_2D);} \\
&\text{glEnd ( );}
\end{align*}
\]

The only difference here is that we must specify both a width (number of columns) and a height (number of rows) for the three-subscript texture array. Both the width and height must be a power of 2, without a border, or 2 plus a power of 2, with a border. We again use RGBA color components, and we state that the pattern has no border and it is not a reduction of a larger texture pattern. Therefore the size of the array stored in \( \text{surfTexArray} \) is \( 4 \times \text{texWidth} \times \text{texHeight} \). For two-dimensional texture patterns, we set the elements in the texture array to the color values in a bottom-to-top order. Starting at the lower-left corner of the color pattern, we set the elements in the first row of the array to the RGBA values corresponding to the bottom row of texture space, and we set the elements in the last row of the array to the RGBA values corresponding to the top of the rectangular texture space (as previously shown in Figure 1).

As with a linear texture pattern, surface pixels in a scene can be assigned the nearest texture color or an interpolated texture color. We select one of these options with the same two \text{glTexParameter} functions that we used for one-dimensional textures. One function specifies the option to use when a texture
pattern is enlarged to fit a coordinate range, and the other function specifies an option that is to be used with pattern reductions. A two-dimensional texture pattern could be stretched in one direction and compressed in the other direction. For example, the following statements instruct the texturing routines to display projected surface positions using the nearest texture color:

```c
glTexParameteri (GL_TEXTURE_2D, GL_TEXTURE_MAG_FILTER, GL_NEAREST);
glTexParameteri (GL_TEXTURE_2D, GL_TEXTURE_MIN_FILTER, GL_NEAREST);
```

To assign an interpolated texture color to surface pixels, we use the symbolic constant `GL_LINEAR` instead of `GL_NEAREST`.

A coordinate position in two-dimensional texture space is selected with

```c
glTexCoord2* (sCoord, tCoord);
```

Texture space is normalized so that the pattern is referenced with coordinate values in the range from 0.0 to 1.0. However, we can use any texture-coordinate values to replicate a pattern across a surface. The texture coordinates can be specified in various formats, and we indicate a data format with a suffix code of `b`, `s`, `i`, `f`, or `d`. We also append the suffix `v` if the texture coordinates are given in an array.

To illustrate the OpenGL functions for a two-dimensional texture space, the following code segment sets up a 32 × 32 pattern and maps it onto a quadrilateral surface. Each texture color is specified with four RGBA components, and the pattern has no border.

```c
GLubyte texArray [32][32][4];
/* Next: assign the texture color components to texArray. */
/* Select nearest-color option. */
glTexParameteri (GL_TEXTURE_2D, GL_TEXTURE_MAG_FILTER, GL_NEAREST);
glTexParameteri (GL_TEXTURE_2D, GL_TEXTURE_MIN_FILTER, GL_NEAREST);
glTexImage2D (GL_TEXTURE_2D, 0, GL_RGBA, 32, 32, 0, GL_RGBA,
             GL_UNSIGNED_BYTE, texArray);

glEnable (GL_TEXTURE_2D);
/* Assign the full range of texture colors to a quadrilateral. */
gBegin (GL_QUADS);
gTexCoord2f (0.0, 0.0); glVertex3fv (vertex1);
gTexCoord2f (1.0, 0.0); glVertex3fv (vertex2);
gTexCoord2f (1.0, 1.0); glVertex3fv (vertex3);
gTexCoord2f (0.0, 1.0); glVertex3fv (vertex4);
gEnd ( );

glDisable (GL_TEXTURE_2D);
```
OpenGL Volume-Texture Functions

Functions for a three-dimensional texture space are simple extensions of those for two-dimensional texture spaces. A four-subscript RGBA texture array with no border, for instance, can be set up with the functions:

```c
glTexImage3D (GL_TEXTURE_3D, 0, GL_RGBA, texWidth, texHeight, texDepth, 0, dataFormat, dataType, volTexArray);
```

```c
glEnable (GL_TEXTURE_3D);
```

The RGBA texture colors are stored in `volTexArray`, which contains \(4 \times \text{texWidth} \times \text{texHeight} \times \text{texDepth}\) elements; and the width, height, and depth of the array must be either a power of 2, or 2 plus a power of 2.

With the following statements, we display pixels using the nearest texture color:

```c
glTexParameteri (GL_TEXTURE_3D, GL_TEXTURE_MAG_FILTER, GL_NEAREST);
```

```c
glTexParameteri (GL_TEXTURE_3D, GL_TEXTURE_MIN_FILTER, GL_NEAREST);
```

For linearly interpolated texture colors, we replace the value `GL_NEAREST` with `GL_LINEAR`.

Three-dimensional texture coordinates are selected with

```c
glTexCoord3* (sCoord, tCoord, rCoord);
```

Each selected position in the texture space is then associated with a spatial coordinate position within a world-coordinate scene.

OpenGL Color Options for Texture Patterns

Elements for a texture space can be specified in many different ways. The third argument in the functions `glTexImage1D`, `glTexImage2D`, and `glTexImage1D` is used to specify the general format and number of color components for each element of a pattern. Nearly 40 symbolic constants are available for this specification. For example, each texture element could be a set of RGBA values, a set of RGB values, a single alpha value, a single red intensity value, a single luminance value, or a luminance value paired with an alpha value. In addition, some constants specify bit size. The OpenGL constant `GL_R3_G3_B2`, for instance, specifies a 1-byte (8-bit) RGB color, with 3 bits allotted to the red component, 3 bits for the green component, and 2 bits for blue.

Parameter `dataFormat` in the texture functions is then used to specify the specific format for the texture elements. We can choose any of 11 symbolic constants for this parameter. This allows us to specify each texture element as an index into a color table, a single alpha value, a single luminance value, a luminance-alpha pair of values, a single intensity value for one of the RGB components, the three RGB components, or the four components of an RGBA specification in the order BGRA. Also, the `dataType` parameter is assigned a value such as `GL_BYTE`, `GL_INT`, `GL_FLOAT`, or a symbolic constant that specifies both data type and bit size. We can choose a value from a set of 20 symbolic constants for the data-type parameter.
OpenGL Texture-Mapping Options

Texture elements can be applied to an object so that the texture values are combined with the current object color components, or the texture values can be used to replace the object color. We select a texture-mapping method with the function

```
glTexEnvi (GL_TEXTURE_ENV, GL_TEXTURE_ENV_MODE,
            applicationMethod);
```

If parameter `applicationMethod` is assigned the value `GL_REPLACE`, then the texture color, luminance, intensity, or alpha value replaces the corresponding object value. For example, a texture pattern of alpha values replaces the object alpha values. Similar replacement operations are used with a texture pattern specified with a single luminance or intensity value. A pattern of green intensity values replaces the green components of the object color.

Assigning the value `GL_MODULATE` to parameter `applicationMethod` results in a “modulation” of the object color values. That is, current object values are multiplied by the texture values. The specific results depend on the format for the elements in the texture pattern so that, for example, alpha values modulate alpha values and intensity values modulate intensity values. The default application method for a texture pattern is `GL_MODULATE`. If an object’s color is white (the default object color), the modulate operation produces the same result as a replace operation, depending on how the elements of the texture pattern have been specified.

We can also use the symbolic constant `GL_DECAL` for the texture-mapping operations, which then employ the RGBA alpha values as transparency coefficients. In this case, the object is treated as if it were transparent with the texture color in the background. If the texture pattern contains only RGB values, with no alpha component, the texture color replaces the object color. Also, in some cases, such as when the texture pattern contains only alpha values, the decal operation is undefined.

When we use `GL_BLEND` as the `applicationMethod` parameter value, the texture routines perform color blending using a color specified with the function

```
glTexEnv* (GL_TEXTURE_ENV, GL_TEXTURE_ENV_COLOR,
              blendingColor);
```

We append the suffix `i` or `f` according to the blending-color data type, and the suffix `v` is also appended if the blending color is given in an array.

OpenGL Texture Wrapping

When coordinate values in texture space are outside the range from 0 to 1.0, we can choose to replicate the patterns described in a texture array using the command

```
glTexParameter* (texSpace, texWrapCoord, GL_REPEAT);
```

Patterns are replicated using only the fractional part of a texture-space coordinate value. Parameter `texSpace` is assigned one of the symbolic values `GL_TEXTURE_1D`, `GL_TEXTURE_2D`, or `GL_TEXTURE_3D`, and parameter `texWrapCoord` designates a texture-space coordinate using either `GL_TEXTURE_WRAP_S`, `GL_TEXTURE_WRAP_T`, or `GL_TEXTURE_WRAP_R`.

To clamp a texture coordinate to the unit interval, we use the symbolic constant `GL_CLAMP` instead of `GL_REPEAT`. If a clamped texture coordinate has a
value greater than 1.0, it is assigned the value 1.0. Similarly, a clamped texture coordinate that has a value less than 0.0 is assigned the value 0.0. We can specify any combination of repeating and clamping for the coordinates in a particular texture space. The default for all coordinates is GL_REPEAT.

Copying OpenGL Texture Patterns from the Frame Buffer

Either an original pattern or a subpattern can be obtained from values stored in the frame buffer. The following function sets up a two-dimensional pattern for the current texture state using a block of RGBA pixel values:

```c
glCopyTexImage2D (GL_TEXTURE_2D, 0, GL_RGBA, x0, y0, texWidth, texHeight, 0);
```

The two 0 values in the argument list again indicate that this pattern is not a reduction and it does not have a border. The frame-buffer position (x0, y0), relative to the lower-left corner of the buffer, references the lower-left corner of a texWidth by texHeight block of pixel colors.

A similar function is available for obtaining a block of pixel colors as a texture subpattern as follows:

```c
glCopyTexSubImage2D (GL_TEXTURE_2D, 0, xTexElement, yTexElement, x0, y0, texSubWidth, texSubHeight);
```

This block of pixel values is placed in the current pattern at texture-element position (xTexElement, yTexElement). Parameters texSubWidth and texSubHeight give the size of the pixel block, whose lower-left corner is at frame-buffer position (x0, y0).

OpenGL Texture-Coordinate Arrays

As with color data, surface-normal vectors, and polygon edge flags, we can specify texture coordinates in lists that can be combined or associated with vertex arrays.

```c
glEnableClientState (GL_TEXTURE_COORD_ARRAY);

glTexCoordPointer (nCoords, dataType, offset, texCoordArray);
```

Parameter nCoords is assigned the value 1, 2, 3, or 4, which designates the dimensionality of the texture pattern. The default value 4 is used to reference texture space in a homogeneous-coordinate form, so that a texture-space position is calculated by dividing the first three coordinate values by the fourth. This form is useful, for example, when the texture pattern is a perspective photograph. Parameter dataType is assigned the constant value GL_SHORT, GL_INT, GL.FLOAT (the default value), or GL_DOUBLE. The byte offset between coordinate positions in array texCoordArray is designated in parameter offset, which has a default value of 0.

Naming OpenGL Texture Patterns

Often it is useful to use several texture patterns in an application, so OpenGL allows multiple, named texture patterns to be created. Then we simply specify which named texture is to be applied at any time. This is a much more efficient method than invoking the glTexImage function each time because each call to glTexImage requires that the pattern be re-created, possibly from color values.
in a data file. To name a texture pattern, we select a positive (unsigned) integer
before the pattern is defined. As an example, the following statements name
and then activate the green and red line pattern from our previous example as
texture 3:

```c
    glBindTexture (GL_TEXTURE_1D, 3);
    glTexImage1D (GL_TEXTURE_1D, 0, GL_RGBA, 4, 0, GL_RGBA,
                 GL_UNSIGNED_BYTE, texLine);
    glBindTexture (GL_TEXTURE_1D, 3);
```

The first `glBindTexture` statement names the pattern, and the second call
to `glBindTexture` designates that pattern as the **current texture state**. If we
have created multiple texture patterns, we could call `glBindTexture` again
with another pattern name to activate that texture for application to some
object in a scene. For a two-dimensional or three-dimensional pattern, we change
the first argument of the `glBindTexture` function to either `GL_TEXTURE_2D`
or `GL_TEXTURE_3D`. When a texture name is first invoked, a texture pattern is
created using the default values for the pattern parameters.

Although we can choose our own name for a pattern, it is generally a better
idea to let OpenGL generate a name for us, so that we don’t have to keep track of
the names that have already been used. For example, we could use the following
code to request one texture name and then use it to create a pattern:

```c
    static GLuint texName;
    glGenTextures (1, &texName);
    glBindTexture (GL_TEXTURE_2D, texName);
```

The first parameter is the number of texture names to be generated, and the
second parameter is an array of `GLuint` into which the names are to be placed.
Because we only wanted one texture name, we had to pass the address of our
single `GLuint` variable as the second parameter so that `glGenTextures` could
place the generated name into it. It is more common to request several texture
names at once, which is why the second parameter can be the address of an array.
For example, the following code obtains a list of 6 unused texture names and uses
one of them to create a pattern:

```c
    static GLuint texNamesArray [6];
    glGenTextures (6, texNamesArray);
    glBindTexture (GL_TEXTURE_2D, texNamesArray [3]);
```

When we are done using a texture pattern, we may want to delete it to release
the space it occupies in OpenGL’s texture memory (for example, to make room for
another texture). We delete one or more existing texture patterns with the function

```c
    glDeleteTextures (nTextures, texNamesArray);
```

The parameters are the same as those for `glGenTextures`: `nTextures` is the
number of pattern names to be deleted, and `texNamesArray` is an array con-
taining those names.

A query command is available in OpenGL to find out if a texture name is in
use for an existing pattern:

```c
    glIsTexture (texName);
```
This function returns the value GL_TRUE if texName is the name of an existing pattern, otherwise the value GL_FALSE is returned. A GL_FALSE value is also returned if texName = 0 or if an error occurs.

OpenGL Texture Subpatterns
Once a texture pattern has been defined, we can create another pattern, called a subpattern, to modify any part, or all, of the original pattern. The texture values in the subpattern replace specified values in the original pattern. This is usually a more efficient process than re-creating a texture with new elements. For example, the following function designates a set of RGBA color values that are to replace a section of a two-dimensional texture that has no border and is not a reduction of a larger pattern:

```c
glTexSubImage2D (GL_TEXTURE_2D, 0, xTexElement,
yTexElement, GL_RGBA, texSubWidth, texSubHeight,
0, dataFormat, dataType, subSurfTexArray);
```

Parameters xTexElement and yTexElement are used to select an integer-coordinate position of a texture element within the original pattern, where position (0, 0) references the texture element at the lower-left corner of the pattern. The subpattern is pasted into the original pattern with its lower-left corner at position (xTexElement, yTexElement). Parameters TexSubWidth and TexSubHeight give the size of the subpattern. The number of color elements in the array subSurfTexArray for a RGBA texture pattern is $4 \times \text{texSubWidth} \times \text{texSubHeight}$. Other parameters are the same as in the glTexImage function, and similar subpatterns can be set up for one-dimensional and three-dimensional textures.

OpenGL Texture Reduction Patterns
For reduced object sizes, we can use OpenGL routines to create a series of texture reduction patterns, referred to as mip maps (see Section 2). One way to create a sequence of reduction patterns is to invoke the glTexImage function repeatedly using higher integer values for the second argument (the "level number") in the function. The original pattern is referenced as reduction-level number 0. A reduction pattern that is half the size of the original pattern is assigned the level number 1, the second one-half size reduction pattern is designated as level number 2, and so on for the other reductions. The copyTexImage function also generates a reduction pattern when we set the level number to 1 or higher.

Alternatively, we can have OpenGL generate reduction patterns automatically. For example, RGBA reduction patterns are obtained for a 16 x 16 surface texture using the following GLU function:

```c
gluBuild2DMipmaps (GL_TEXTURE_2D, GL_RGBA, 16, 16, GL_RGBA,
GL_UNSIGNED_BYTE, surfTexArray);
```

A complete set of four patterns, at the reduced sizes of $8 \times 8$, $4 \times 4$, $2 \times 2$, and $1 \times 1$, is generated by this function. We can also set up selected reductions using

```c
gluBuild2DMipmapLevels (GL_TEXTURE_2D, GL_RGBA, 16, 16,
GL_RGBA, GL_UNSIGNED_BYTE, 0, minLevel, maxLevel,
surfTexArray);
```
This function produces reduction patterns for a range of level numbers specified by parameters minLevel and maxLevel. In each case, the mip maps are constructed for the current texture pattern, specified at level number 0.

We choose a method for determining pixel colors from the reduction patterns using the glTexParameter function and the GL_TEXTURE_MIN_FILTER symbolic constant. For example, the following function designates the mapping procedure for a two-dimensional texture pattern:

\[
\text{glTexParameter} (\text{GL\_TEXTURE\_2D}, \text{GL\_TEXTURE\_MIN\_FILTER}, \\
\quad \text{GL\_NEAREST\_MIPMAP\_NEAREST});
\]

This function specifies that the texture routines should use the reduction pattern that most closely matches the pixel size (MIPMAP\_NEAREST). A pixel is then assigned the color of the nearest texture element (GL\_NEAREST) in that reduction pattern. With the symbolic constant GL\_LINEAR\_MIPMAP\_NEAREST, we specify a linear combination of texture colors from the nearest reduction pattern. With GL\_NEAREST\_MIPMAP\_LINEAR (the default value), we specify an average color calculated from the nearest texture elements in each of the reduction patterns closest to the pixel size. In addition, GL\_LINEAR\_MIPMAP\_LINEAR computes a pixel color using a linear combination of texture colors from a set of closest-size reduction patterns.

### OpenGL Texture Borders

When multiple textures, or multiple copies of a single texture, are applied to an object, aliasing effects may be apparent at the edges of adjacent patterns when pixel colors are computed by linearly interpolating the texture colors. This can be avoided by including a border with each texture pattern, where border colors match the texture edge colors in the adjacent pattern.

We can designate a texture border color in several ways. The color value in an adjacent pattern can be copied to the border in another pattern using the glTexSubImage function, or the border colors can be directly assigned in the texture array specified with the glTexImage function. Another option is to set a border color using the glTexParameter routine. For example, we can assign a border color for a two-dimensional pattern with

\[
\text{glTexParameterfv} (\text{GL\_TEXTURE\_2D}, \text{GL\_TEXTURE\_BORDER\_COLOR}, \\
\quad \text{borderColor});
\]

where parameter borderColor is assigned a four-element set of RGBA color components. The default border color is black (0.0, 0.0, 0.0).

### OpenGL Proxy Textures

In any of the glTexImage functions, we can set the first argument to a symbolic constant, called a texture proxy. The purpose of this constant is to hold the definition of the texture pattern until we find out if there are enough resources to handle this pattern. For a two-dimensional pattern, the proxy constant is GL\_PROXY\_TEXTURE\_2D, and similar constants are available for linear and volumetric texture patterns. Once we have set up the texture proxy, we use glGetTexLevelFunction to determine whether specific parameter values can be accommodated.
As an example of using a texture proxy, the following statements query the system to determine whether the height specified for a two-dimensional pattern can be used:

```c
GLint texHeight;

glTexImage2D (GL_PROXY_TEXTURE_2D, 0, GL_RGBA12, 16, 16, 0,
GL_RGBA, GL_UNSIGNED_BYTE, NULL);
gGetTexLevelParameteriv (GL_PROXY_TEXTURE_2D, 0, GL_RGBA12,
GL_TEXTURE_HEIGHT, &texHeight);
```

If the system cannot accommodate the requested pattern height (16, in this case), a value of 0 is returned in parameter `texHeight`. Otherwise, the value returned is the value requested. Other pattern parameters can be queried similarly using symbolic constants such as `GL_TEXTURE_WIDTH`, `GL_TEXTURE_DEPTH`, `GL_TEXTURE_BORDER`, and `GL_TEXTURE_BLUE_SIZE`. In each case, a returned value of 0 indicates that the requested parameter value in the `glTexImage` function cannot be accommodated. For floating-point data values, we replace the suffix `i` with the code `f`.

Although we might obtain an affirmative answer for a proposed texture, we still might not be able to store the pattern in memory. This can occur when another pattern is occupying the available memory.

**Automatic Texturing of Quadric Surfaces**

Routines are available in OpenGL for automatically generating texture coordinates in certain applications. This feature is particularly useful when it may be difficult to determine surface coordinates for an object directly, and a GLU function is available for applying these routines to quadric surfaces.

To map a texture pattern to a quadric surface, we first set up the parameters for the texture space. Then we invoke the following function and define the quadric object:

```c
gluQuadricTexture (quadSurfObj, GL_TRUE)
```

Parameter `quadSurfObj` in this function is the name of the quadric object. If we want to deactivate the texturing of the quadric surface, we change the symbolic constant `GL_TRUE` to `GL_FALSE`.

**Homogeneous Texture Coordinates**

A four-dimensional texture-space position is specified with

```c
gTexCoord4* (sCoord, tCoord, rCoord, hCoord);
```

Texture coordinates are transformed using a $4 \times 4$ matrix in the same way that scene coordinates are transformed: Each coordinate is divided by the homogeneous parameter. Thus, the values for the texture coordinates $s$, $t$, and $r$ in this function are divided by the homogeneous parameter $h_{\text{tex}}$ to produce an actual texture-space position.

Homogeneous coordinates in texture space are useful when multiple perspective effects are combined in one display. For example, a perspective view of an object may include a texture pattern produced with a different perspective-projection transformation. The texture pattern can then be modified using homogeneous texture coordinates to adjust the texture perspective. Many other effects are possible using homogeneous texture coordinates to manipulate a texture mapping.
Additional OpenGL Texture Options

Functions are available in OpenGL for performing many other texture manipulations and applications. If we obtain a texture pattern (from a photograph or other source) that is not a power of 2, OpenGL provides a function to modify the size of the pattern. In some implementations of OpenGL, multitexturing routines are available for pasting multiple texture patterns onto an object. Environment-mapping can be simulated in OpenGL by creating a texture map in the shape of a spherical surface, and texture coordinates for spherical environment patterns, as well as other texture applications, can be generated automatically.

6 Summary

Surface detail can be added to objects using polygon facets, texture mapping, bump mapping, or frame mapping. Small polygon facets can be overlaid on larger surfaces to provide various kinds of designs. Alternatively, texture patterns can be defined in one-dimensional, two-dimensional, and three-dimensional spaces, which can be used to add texture to a line, a surface, or a volume of space. Procedural texture mapping uses functions to calculate variations in object lighting effects. Bump mapping is a means for modeling surface irregularities by applying a bump function to perturb surface normal vectors. Frame mapping is an extension of bump mapping that can be used to model characteristics of anisotropic materials by allowing for horizontal surface variations, as well as vertical variations.

The core library of OpenGL contains a number of functions for applying texture array patterns to objects. Texture patterns can be obtained from a number of different sources. Table 1 provides a summary of these OpenGL texture-mapping functions.

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glTexImage1D</td>
<td>Specifies parameters for setting up a one-dimensional texture space. (Activates texturing with glEnable.)</td>
</tr>
<tr>
<td>glTexImage2D</td>
<td>Specifies parameters for setting up a two-dimensional texture space.</td>
</tr>
<tr>
<td>glTexImage3D</td>
<td>Specifies parameters for setting up a three-dimensional texture space.</td>
</tr>
<tr>
<td>glTexParameter</td>
<td>Specifies parameters for the texture-mapping routines.</td>
</tr>
<tr>
<td>glTexCoord</td>
<td>Specifies a value for a texture coordinate in one-dimensional, two-dimensional, three-dimensional, or four-dimensional texture space.</td>
</tr>
<tr>
<td>glTexEnv</td>
<td>Specifies texture-environment parameters, such as a blending color for texture mapping.</td>
</tr>
</tbody>
</table>

(Continued)
Summary of OpenGL Texture-Mapping Functions (Continued)

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glCopyTexImage</td>
<td>Copies a block of frame-buffer pixel colors for use as a texture pattern.</td>
</tr>
<tr>
<td>glCopyTexSubImage</td>
<td>Copies a block of frame-buffer pixel colors for use as a texture subimage.</td>
</tr>
<tr>
<td>glTexCoordPointer</td>
<td>Specifies texture coordinates in a list that is associated with a vertex list.</td>
</tr>
<tr>
<td>glBindTexture</td>
<td>Assigns a name to a texture pattern; also used to activate a named pattern.</td>
</tr>
<tr>
<td>glDeleteTextures</td>
<td>Eliminates a list of named textures.</td>
</tr>
<tr>
<td>glGenTextures</td>
<td>Generates texture names automatically.</td>
</tr>
<tr>
<td>glIsTexture</td>
<td>Queries command to determine whether a named texture exists.</td>
</tr>
<tr>
<td>glTexSubImage</td>
<td>Creates a texture subpattern.</td>
</tr>
<tr>
<td>gluBuild*Mipmaps</td>
<td>Generates texture reduction patterns automatically for a one-dimensional, two-dimensional, or three-dimensional texture space.</td>
</tr>
<tr>
<td>gluBuild*MipmapLevels</td>
<td>Generates texture reduction patterns automatically for a specified level for a one-dimensional, two-dimensional, or three-dimensional texture space.</td>
</tr>
<tr>
<td>glGetTexLevelParameter</td>
<td>Queries the system to determine whether a texture parameter value can be accommodated.</td>
</tr>
<tr>
<td>gluQuadricTexture</td>
<td>Activates or deactivates texturing for a quadric surface.</td>
</tr>
</tbody>
</table>

REFERENCES

Texture-mapping methods and applications are discussed in Williams (1983), Segal, et al. (1992), and Demers (2002).

Additional programming examples using OpenGL texture functions are given in Woo, et al. (1999). Programming examples for the OpenGL and texture functions are also available at Nate Robins’s tutorial website: http://www.xmission.com/~nate/opengl.html. Finally, a complete listing of OpenGL rendering functions is provided in Shreiner (2000).

EXERCISES

1 Write a program to map a given texture pattern onto each face of a specified cube.

2 Modify the program in the preceding exercise so that the pattern is mapped to each face of a tetrahedron.

3 Modify the program in the preceding exercise so that the pattern is mapped to a specified section of a spherical surface.

4 Write a program to map a given one-dimensional texture pattern onto a specified face of a cube as a diagonal stripe.

5 Modify the program in the preceding example so that the one-dimensional texture is mapped to the surface of a sphere, given two points on the spherical surface.
Texturing and Surface Detail Methods

6 Given a spherical surface, write a bump-mapping procedure to simulate the bumpy surface of an orange.

7 Write a bump-mapping routine to produce surface-normal variations for any specified bump function.

8 Write a complete OpenGL program to display a set of diagonal lines using various one-dimensional texture patterns.

9 Write a program using a two-dimensional OpenGL texture pattern to display a red-and-black checkerboard on a gray background.

10 Modify the program in the preceding exercise so that the checkerboard has blue and yellow squares and the background is black.

11 Write a program using a two-dimensional OpenGL texture pattern to display a red rectangle with a set of evenly spaced horizontal blue stripes. Set the background color to white.

12 Modify the program in the preceding exercise to map the texture pattern onto the surface of a sphere.

13 Modify the program in the preceding exercise to map the texture pattern onto the surface of the GLUT teapot.

IN MORE DEPTH

1 Examine the objects in your scene that you have rendered thus far with simple shading models and choose those that would more appropriately be rendered with more complex surfaces. Obtain from another source, or design yourself, texture patterns for each of these objects that most appropriately fit their desired textural properties. Discuss the different possible ways that you could map each of these textures onto their respective objects based on the shapes of the objects.

2 Use the OpenGL surface-mapping functions to map the texture patterns that you produced in the previous exercise onto the appropriate objects in your scene. Adjust the blending parameters between the mapped textures and the object color values as appropriate to obtain the desired look of the objects in your scene.
Texturing and Surface-Detail Methods

Color Plate 18
Modeling and rendering stages in the simulation of a square plate struck by a rigid projectile: (a) A wire-frame model of the scene; (b) A final rendered version. (Image created by M. Wicke, D. Ritchie, B. Klingner, S. Burke, J. Shewchuk, and J. O’Brien, University of California, Berkeley.)

Color Plate 19
Scenes illustrating computer-graphics generation of surface detail for various objects: (a) cactus plants with added spines and flowers (Courtesy of Deborah R. Fowler, Przemyslaw Prusinkiewicz, and Johannes Battjes, University of Calgary. © 1992), (b) seashells with various patterns and fluted surfaces (Courtesy of Deborah R. Fowler, Hans Meinhardt, and Przemyslaw Prusinkiewicz, University of Calgary. © 1992), (c) a table of fruit (Courtesy of SOFTIMAGE, Inc.), and (d) surface patterns on chess pieces and a chessboard produced with texture-mapping methods (Courtesy of SOFTIMAGE, Inc.).

**Color Plate 20**
A scene with object characteristics modeled using solid-texture methods. (Courtesy of Peter Shirley, Computer Science Department, University of Utah.)

**Color Plate 21**
Rendering the characteristics of rough surfaces using bump mapping. (Courtesy of (a) Peter Shirley, Computer Science Department, University of Utah, and (b) SOFTIMAGE, Inc.)
Our discussions of color up to this point have concentrated on methods involving red, green, and blue (RGB) components, which we use for generating displays on video monitors. Several other color descriptions are useful as well in computer-graphics applications. Some methods are used to describe color output on printers and plotters, some are used for transmitting and storing color information, and others are used to provide a more intuitive color-parameter interface to a program.
1 Properties of Light

Light exhibits many different characteristics, and we describe the properties of light in different ways in different contexts. Physically, we can characterize light as radiant energy, but we also need other concepts to describe our perception of light.

The Electromagnetic Spectrum

In physical terms, color is electromagnetic radiation within a narrow frequency band. Some of the other frequency groups in the electromagnetic spectrum are referred to as radio waves, microwaves, infrared waves, and X-rays. Figure 1 shows the approximate frequency ranges for these various aspects of electromagnetic radiation.

Each frequency value within the visible region of the electromagnetic spectrum corresponds to a distinct spectral color. At the low-frequency end (approximately \(3.8 \times 10^{14}\) hertz) are the red colors, and at the high-frequency end (approximately \(7.9 \times 10^{14}\) hertz) are the violet colors. Actually, the human eye is sensitive to some frequencies into the infrared and ultraviolet bands. Spectral colors range from shades of red through orange and yellow, at the low-frequency end, to shades of green, blue, and violet at the high end.

In the wave model of electromagnetic radiation, light can be described as oscillating transverse electric and magnetic fields propagating through space. The electric and magnetic fields are oscillating in directions that are perpendicular to each other and to the direction of propagation. For each spectral color, the rate of oscillation of the field magnitude is given by the frequency \(f\). Figure 2
illust\textquotesingle{}lates the time-varying oscillations for the magnitude of the electric field within one plane. The time between any two consecutive positions on the wave that have the same amplitude is called the \textit{period} ($T$) of the wave, which is the inverse of the frequency (i.e., $T = 1/f$). And the distance that the wave has traveled from the beginning of one oscillation to the beginning of the next oscillation is called the \textit{wavelength} ($\lambda$). For one spectral color (a monochromatic wave), the wavelength and frequency are inversely proportional to each other, with the proportionality constant as the speed of light ($c$):

$$c = \lambda f$$ (1)

Frequency for each spectral color is a constant for all materials, but the speed of light and the wavelength are material dependent. In a vacuum, the speed of light is very nearly $c = 3 \times 10^{10}$ cm/sec. Light wavelengths are very small, so length units for designating spectral colors are usually given in angstroms ($1\,\text{Å} = 10^{-8}$ cm) or in nanometers ($1\,\text{nm} = 10^{-7}$ cm). An equivalent term for nanometer is milli-micron. Light at the low-frequency end of the spectrum (red) has a wavelength of approximately 780 nanometers (nm), and the wavelength at the other end of the spectrum (violet) is about 380 nm. Because wavelength units are somewhat more convenient to deal with than frequency units, spectral colors are typically specified in terms of the wavelength values in a vacuum.

A light source such as the sun or a standard household light bulb emits all frequencies within the visible range to produce white light. When white light is incident upon an opaque object, some frequencies are reflected and some are absorbed. The combination of frequencies present in the reflected light determines what we perceive as the color of the object. If low frequencies are predominant in the reflected light, the object is described as red. In this case, we say that the perceived light has a \textbf{dominant frequency} (or \textbf{dominant wavelength}) at the red end of the spectrum. The dominant frequency is also called the \textbf{hue}, or simply the \textbf{color}, of the light.

\textbf{Psychological Characteristics of Color}

Other properties besides frequency are needed to characterize our perception of light. When we view a source of light, our eyes respond to the color (or dominant frequency) and two other basic sensations. One of these we call the \textbf{brightness}, which corresponds to the total light energy and can be quantified as the luminance of the light. The third perceived characteristic is called the \textbf{purity}, or the \textbf{saturation}, of the light. Purity describes how close a light appears to be to a pure spectral color, such as red. Pastels and pale colors have low purity (low saturation) and they appear to be nearly white. Another term, \textit{chromaticity}, is used to refer collectively to the two properties describing color characteristics: purity and dominant frequency (hue).

Radiation emitted by a white light source has an energy distribution that can be represented over the visible frequencies as in Figure 3. Each frequency component within the range from red to violet contributes more or less equally to the total energy, and the color of the source is described as white. When a dominant frequency is present, the energy distribution for the source takes a form such as that in Figure 4. We would describe this light as a red color (the dominant frequency), with a relatively high value for the purity. The energy density of the dominant light component is labeled as $E_D$ in this figure, and the contributions from the other frequencies produce white light of energy density $E_W$. We can calculate the brightness of the source as the area under the curve, which gives the total energy density emitted. Purity (saturation) depends on the difference between $E_D$ and...
Energy distribution for a white light source.

\( E_W \). The larger the energy \( E_D \) of the dominant frequency compared to the white-light component \( E_W \), the higher the purity of the light. We have a purity of 100 percent when \( E_W = 0 \) and a purity of 0 percent when \( E_W = E_D \).

## 2 Color Models

Any method for explaining the properties or behavior of color within some particular context is called a **color model**. No single model can explain all aspects of color, so we make use of different models to help describe different color characteristics.

### Primary Colors

When we combine the light from two or more sources with different dominant frequencies, we can vary the amount (intensity) of light from each source to generate a range of additional colors. This represents one method for forming a color model. The hues that we choose for the sources are called the **primary colors**, and the **color gamut** for the model is the set of all colors that we can produce from the primary colors. Two primaries that produce white are referred to as **complementary colors**. Examples of complementary color pairs are red and cyan, green and magenta, and blue and yellow.

No finite set of real primary colors can be combined to produce all possible visible colors. Nevertheless, three primaries are sufficient for most purposes, and colors not in the color gamut for a specified set of primaries can still be described using extended methods. Given a set of three primary colors, we can characterize any fourth color using color-mixing processes. Thus, a mixture of one or two of the primaries with the fourth color can be used to match some combination of the remaining primaries. In this extended sense, a set of three primary colors can be considered to describe all colors. Figure 5 shows a set of **color-matching functions** for three primaries and the amount of each needed to produce any spectral color. The curves plotted in Figure 5 were obtained by averaging the judgments of a large number of observers. Colors in the vicinity of 500 nm can be matched only by “subtracting” an amount of red light from a combination of blue and green lights. This means that a color around 500 nm is described only by combining that color with an amount of red light to produce the blue-green combination specified in the diagram. Thus, an RGB color monitor cannot display colors in the neighborhood of 500 nm.
Intuitive Color Concepts

An artist creates a color painting by mixing color pigments with white and black pigments to form the various shades, tints, and tones in the scene. Starting with the pigment for a "pure color" ("pure hue"), the artist adds a black pigment to produce different shades of that color. The more black pigment, the darker the shade. Similarly, different tints of the color are obtained by adding a white pigment to the original color, making it lighter as more white is added. Tones of the color are produced by adding both black and white pigments.

To many, these color concepts are more intuitive than describing a color as a set of three numbers that give the relative proportions of the primary colors. It is generally much easier to think of creating a pastel red color by adding white to pure red and producing a dark blue color by adding black to pure blue. Therefore, graphics packages providing color palettes to a user often employ two or more color models. One model provides an intuitive color interface for the user, and the others describe the color components for the output devices.

3 Standard Primaries and the Chromaticity Diagram

Because no finite set of light sources can be combined to display all possible colors, three standard primaries were defined in 1931 by the International Commission on Illumination, referred to as the CIE (Commission Internationale de l’Eclairage). The three standard primaries are imaginary colors. They are defined mathematically with positive color-matching functions (Figure 6) that
specify the amount of each primary needed to describe any spectral color. This provides an international standard definition for all colors, and the CIE primaries eliminate negative-value color-matching and other problems associated with selecting a set of real primaries.

The XYZ Color Model

The set of CIE primaries is generally referred to as the XYZ color model, where parameters \( X \), \( Y \), and \( Z \) represent the amount of each CIE primary needed to produce a selected color. Thus, a color is described with the XYZ model in the same way that we described a color using the RGB model.

In the three-dimensional XYZ color space, we represent any color \( C(\lambda) \) as

\[
C(\lambda) = (X, Y, Z)
\]  

where \( X \), \( Y \), and \( Z \) are calculated from the color-matching functions (Figure 6):

\[
X = k \int_{\text{visible} \lambda} f_X(\lambda) I(\lambda) \, d\lambda \\
Y = k \int_{\text{visible} \lambda} f_Y(\lambda) I(\lambda) \, d\lambda \\
Z = k \int_{\text{visible} \lambda} f_Z(\lambda) I(\lambda) \, d\lambda
\]  

Parameter \( k \) in these calculations has the value 683 lumens/watt, where lumen is a unit of measure for light radiation per unit solid angle from a “standard” point light source (once called a candle). The function \( I(\lambda) \) represents the spectral radiance, which is the selected light intensity in a particular direction, and the color-matching function \( f_Y \) is chosen so that parameter \( Y \) is the luminance for that color. Luminance values are normally adjusted to the range from 0 to 100.0, where 100.0 represents the luminance of white light.

Any color can be represented in the XYZ color space as an additive combination of the primaries using unit vectors \( X, Y, Z \). Thus, we can write Equation 2 as

\[
C(\lambda) = X X + YX + Z X
\]  

Normalized XYZ Values

In discussing color properties, it is convenient to normalize the amounts in Equation 3 against the sum \( X + Y + Z \), which represents the total light energy. Normalized amounts are thus calculated as

\[
x = \frac{X}{X + Y + Z}, \quad y = \frac{Y}{X + Y + Z}, \quad z = \frac{Z}{X + Y + Z}
\]  

Because \( x + y + z = 1 \), any color can be represented with just the \( x \) and \( y \) amounts. Also, we have normalized against total energy, so parameters \( x \) and \( y \) depend only on hue and purity and are called the chromaticity values. However, the \( x \) and \( y \) values alone do not allow us to describe all properties of the color completely, and we cannot obtain the amounts \( X \), \( Y \), and \( Z \). Therefore, a complete description of a color is typically given with three values: \( x \), \( y \), and the luminance \( Y \). The remaining CIE amounts are then calculated as

\[
X = \frac{x}{y} Y, \quad Z = \frac{z}{y} Y
\]  

where \( z = 1 - x - y \). Using chromaticity coordinates \((x, y)\), we can represent all colors on a two-dimensional diagram.
The CIE Chromaticity Diagram

When we plot the normalized amounts $x$ and $y$ for colors in the visible spectrum, we obtain the tongue-shaped curve shown in Figure 7. This curve is called the **CIE chromaticity diagram**. Points along the curve are the spectral colors (pure colors). The line joining the red and violet spectral points, referred to as the *purple line*, is not part of the spectrum. Interior points represent all possible visible color combinations. Point $C$ in the diagram corresponds to the white-light position. Actually, this point is plotted for a white light source known as **illuminant C**, which is used as a standard approximation for average daylight.

Luminance values are not available in the chromaticity diagram because of normalization. Colors with different luminance but with the same chromaticity map to the same point. The chromaticity diagram is useful for:

- Comparing color gamuts for different sets of primaries.
- Identifying complementary colors.
- Determining purity and dominant wavelength for a given color.

**Color Gamuts**

We identify color gamuts on the chromaticity diagram as straight-line segments or polygon regions. All colors along the straight line joining points $C_1$ and $C_2$ in Figure 8 can be obtained by mixing appropriate amounts of the colors $C_1$ and $C_2$. If a greater proportion of $C_1$ is used, the resultant color is closer to $C_1$ than to $C_2$. The color gamut for three points, such as $C_3$, $C_4$, and $C_5$ in Figure 8, is a triangle with vertices at the three color positions. These three primaries can generate only the colors inside or on the bounding edges of the triangle. Thus, the chromaticity diagram helps us to understand why no set of three primaries can be additively combined to generate all colors, because no triangle within the diagram can encompass all colors. Color gamuts for video monitors and hard-copy devices are compared conveniently on the chromaticity diagram.

**Complementary Colors**

Because the color gamut for two points is a straight line, complementary colors must be represented on the chromaticity diagram as two points on opposite sides of $C$ and collinear with $C$, as in Figure 9. The distances of the two colors $C_1$ and $C_2$ to $C$ determine the amounts of each needed to produce white light.
Dominant Wavelength
To determine the dominant wavelength of a color, we draw a straight line from $C$ through that color point to a spectral color on the chromaticity curve. The spectral color $C_s$ in Figure 10 is the dominant wavelength for color $C_1$ in this diagram. Thus, color $C_1$ can be represented as a combination of white light $C$ and the spectral color $C_s$. This method for determining dominant wavelength will not work for color points that are between $C$ and the purple line. Drawing a line from $C$ through point $C_2$ in Figure 10 takes us to point $C_p$ on the purple line, which is not in the visible spectrum. In this case, we take the compliment of $C_p$ on the spectral curve, which is the point $C_{sp}$, as the dominant wavelength. Colors such as $C_2$ in this diagram have spectral distributions with subtractive dominant wavelengths. We can describe such colors by subtracting the spectral dominant wavelength from white light.

Purity
For a color point such as $C_1$ in Figure 10, we determine the purity as the relative distance of $C_1$ from $C$ along the straight line joining $C$ to $C_s$. If $d_{C_1}$ denotes the distance from $C$ to $C_1$ and $d_{C_s}$ is the distance from $C$ to $C_s$, we can represent purity as the ratio $d_{C_1}/d_{C_s}$. Color $C_1$ in this figure is about 25 percent pure, because it is situated at about one-fourth the total distance from $C$ to $C_s$. At position $C_s$, the color point would be 100 percent pure.

4 The RGB Color Model
According to the tristimulus theory of vision, our eyes perceive color through the stimulation of three visual pigments in the cones of the retina. One of the pigments is most sensitive to light with a wavelength of about 630 nm (red), another has its peak sensitivity at about 530 nm (green), and the third pigment is most receptive to light with a wavelength of about 450 nm (blue). By comparing intensities in a light source, we perceive the color of the light. This theory of vision is the basis for displaying color output on a video monitor using the three primaries red, green, and blue, which is referred to as the RGB color model.

We can represent this model using the unit cube defined on $R$, $G$, and $B$ axes, as shown in Figure 11. The origin represents black and the diagonally opposite
vertex, with coordinates (1, 1, 1), is white. Vertices of the cube on the axes represent the primary colors, and the remaining vertices are the complementary color points for each of the primary colors.

As with the XYZ color system, the RGB color scheme is an additive model. Each color point within the unit cube can be represented as a weighted vector sum of the primary colors, using unit vectors $R$, $G$, and $B$:

$$C(\lambda) = (R, G, B) = R R + G G + B B$$

where parameters $R$, $G$, and $B$ are assigned values in the range from 0 to 1.0. For example, the magenta vertex is obtained by adding maximum red and blue values to produce the triple (1, 0, 1), and white at (1, 1, 1) is the sum of the maximum values for red, green, and blue. Shades of gray are represented along the main diagonal of the cube from the origin (black) to the white vertex. Points along this diagonal have equal contributions from each primary color, and a gray shade halfway between black and white is represented as (0.5, 0.5, 0.5). The color graduations along the front and top planes of the RGB cube are illustrated in Color Plate 22.

Chromaticity coordinates for the National Television System Committee (NTSC) standard RGB phosphors are listed in Table 1. Also listed are the RGB chromaticity coordinates within the CIE color model and the approximate values used for phosphors in color monitors. Figure 12 shows the approximate color gamut for the NTSC standard RGB primaries.

<table>
<thead>
<tr>
<th>RGB (x, y) Chromaticity Coordinates</th>
<th>NTSC Standard</th>
<th>CIE Model</th>
<th>Approx. Color Monitor Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>R (0.670, 0.330)</td>
<td>(0.735, 0.265)</td>
<td>(0.628, 0.346)</td>
<td></td>
</tr>
<tr>
<td>G (0.210, 0.710)</td>
<td>(0.274, 0.717)</td>
<td>(0.268, 0.588)</td>
<td></td>
</tr>
<tr>
<td>B (0.140, 0.080)</td>
<td>(0.167, 0.009)</td>
<td>(0.150, 0.070)</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 12**
The RGB color gamut for NTSC chromaticity coordinates. Illuminant C is at position (0.310, 0.316), with a luminance value of $Y = 100.0$. 

---

**TABLE 1**
RGB (x, y) Chromaticity Coordinates

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</tr>
</tbody>
</table>
5 The YIQ and Related Color Models

Although an RGB graphics monitor requires separate signals for the red, green, and blue components of an image, a television monitor uses a composite signal. NTSC color encoding for forming the composite video signal is called the YIQ color model.

The YIQ Parameters

In the YIQ color model, parameter \( Y \) is the same as the \( Y \) component in the CIE \( XYZ \) color space. Luminance (brightness) information is conveyed by the \( Y \) parameter, while chromaticity information (hue and purity) is incorporated into the \( I \) and \( Q \) parameters. A combination of red, green, and blue is chosen for the \( Y \) parameter to yield the standard luminosity curve. Because \( Y \) contains the luminance information, black-and-white television monitors use only the \( Y \) signal. Parameter \( I \) contains orange-cyan color information that provides the flesh-tone shading, and parameter \( Q \) carries green-magenta color information.

The NTSC composite color signal is designed to provide information in a form that can be received by black-and-white television monitors, which obtain grayscale information for a picture within a 6-MHz bandwidth. Thus, the \( YIQ \) information is also encoded within a 6-MHz bandwidth, but the luminance and chromaticity values are encoded on separate analog signals. In this way, the luminance signal is unchanged for black-and-white monitors, and the color information is simply added within the same bandwidth. Luminance information, the \( Y \) value, is conveyed as an amplitude modulation on a carrier signal with a bandwidth of about 4.2 MHz. Chromaticity information, the \( I \) and \( Q \) values, is combined on a second carrier signal that has a bandwidth of about 1.8 MHz. The parameter names \( I \) and \( Q \) refer to the modulation methods used to encode the color information on this carrier. An amplitude-modulation encoding (the “in-phase” signal) transmits the \( I \) value, using about 1.3 MHz of the bandwidth. And a phase-modulation encoding (the “quadrature” signal), using about 0.5 MHz, carries the \( Q \) value.

Luminance values are encoded at a higher precision in the NTSC signal (4.2 MHz bandwidth) than the chromaticity values (1.8 MHz bandwidth), because we can detect small brightness changes more easily compared to small color changes. However, the lower precision for the chromaticity encoding does result in some degradation of the color quality for an NTSC picture.

We can calculate the luminance value for an RGB color. One method for producing chromaticity values is to subtract the luminance from the red and blue components of the color. Thus,

\[
Y = 0.299 \, R + 0.587 \, G + 0.114 \, B \\
I = R - Y \\
Q = B - Y 
\]  

Transformations Between RGB and YIQ Color Spaces

An RGB color is converted to a set of YIQ values using an NTSC encoder that implements the calculations in Equation 9 and modulates the carrier signals. The conversion from RGB space to YIQ space is accomplished using the following transformation matrix:

\[
\begin{bmatrix}
Y \\
I \\
Q
\end{bmatrix} = 
\begin{bmatrix}
0.299 & 0.587 & 0.114 \\
0.701 & -0.587 & -0.114 \\
-0.299 & -0.587 & 0.886
\end{bmatrix}
\begin{bmatrix}
R \\
G \\
B
\end{bmatrix} 
\]  

(9)
Conversely, an NTSC video signal is converted to RGB color values using an NTSC decoder, which first separates the video signal into the YIQ components, and then converts the YIQ values to RGB values. The conversion from YIQ space to RGB space is accomplished with the inverse of transformation 9:

\[
\begin{bmatrix}
R \\
G \\
B
\end{bmatrix} =
\begin{bmatrix}
1.000 & 1.000 & 0.000 \\
1.000 & 0.000 & 1.000 \\
1.000 & 0.000 & 0.000
\end{bmatrix}
\begin{bmatrix}
Y \\
I \\
Q
\end{bmatrix}
\] (10)

The YUV and \(Y_C, C_b\) Systems

Because of the lower bandwidth assigned to the chromaticity information in the NTSC composite analog video signal, the color quality of an NTSC picture is somewhat impaired. Therefore, variations of the YIQ encoding have been developed to improve the color quality of video transmissions. One such encoding is the YUV set of color parameters, which provides the composite color information for video transmissions by systems such as Phase Alternation Line (PAL) Broadcasting, used in most of Europe, as well as Africa, Australia, and Eurasia. Another variation of YIQ is the digital encoding called \(Y_C, C_b\). This color representation is used for digital video transformations, and it is incorporated into various graphics file formats, such as the JPEG system.

6 The CMY and CMYK Color Models

A video monitor displays color patterns by combining light that is emitted from the screen phosphors, which is an additive process. However, hard-copy devices, such as printers and plotters, produce a color picture by coating a paper with color pigments. We see the color patterns on the paper by reflected light, which is a subtractive process.

The CMY Parameters

A subtractive color model can be formed with the three primary colors cyan, magenta, and yellow. As we have noted, cyan can be described as a combination of green and blue. Therefore, when white light is reflected from cyan-colored ink, the reflected light contains only the green and blue components, and the red component is absorbed, or subtracted, by the ink. Similarly, magenta ink subtracts the green component from incident light, and yellow subtracts the blue component. A unit cube representation for the CMY model is illustrated in Figure 13.

In the CMY model, the spatial position \((1, 1, 1)\) represents black, because all components of the incident light are subtracted. The origin represents white light. Equal amounts of each of the primary colors produce shades of gray along the main diagonal of the cube. A combination of cyan and magenta ink produces blue light, because the red and green components of the incident light are absorbed. Similarly, a combination of cyan and yellow ink produces green light, and a combination of magenta and yellow ink yields red light.

The CMY printing process often uses a collection of four ink dots, which are arranged in a close pattern somewhat as an RGB monitor uses three phosphor dots. Thus, in practice, the CMY color model is referred to as the CMYK model, where \(K\) is the black color parameter. One ink dot is used for each of the primary colors (cyan, magenta, and yellow), and one ink dot is black. A black dot is included because reflected light from the cyan, magenta, and yellow inks typically produce only shades of gray. Some plotters produce different color combinations...
by spraying the ink for the three primary colors over each other and allowing them to mix before they dry. For black-and-white or grayscale printing, only the black ink is used.

**Transformations Between CMY and RGB Color Spaces**

We can express the conversion from an RGB representation to a CMY representation using the following matrix transformation:

\[
\begin{bmatrix}
C \\
M \\
Y
\end{bmatrix} = \begin{bmatrix} 1 \\ 1 \\ 1 \end{bmatrix} - \begin{bmatrix} R \\ G \\ B \end{bmatrix}
\] (11)

where the white point in RGB space is represented as the unit column vector. And we convert from a CMY color representation to an RGB representation using the matrix transformation

\[
\begin{bmatrix}
R \\
G \\
B
\end{bmatrix} = \begin{bmatrix} 1 \\ 1 \\ 1 \end{bmatrix} - \begin{bmatrix} C \\ M \\ Y \end{bmatrix}
\] (12)

In this transformation, the unit column vector represents the black point in the CMY color space.

For the conversion from RGB to the CMYK color space, we first set \( K = \max(R, G, B) \). Then \( K \) is subtracted from each of \( C, M, \) and \( Y \) in Equation 11. Similarly, for the transformation from CMYK to RGB, we first set \( K = \min(R, G, B) \). Then \( K \) is subtracted from each of \( R, G, \) and \( B \) in Equation 12. In practice, these transformation equations are often modified to improve the printing quality for a particular system.

### 7 The HSV Color Model

Interfaces for selecting colors often use a color model based on intuitive concepts, rather than a set of primary colors. We can give a color specification in an intuitive model by selecting a spectral color and the amounts of white and black that are to be added to that color to obtain different shades, tints, and tones (Section 2).

**The HSV Parameters**

Color parameters in this model are called **hue** \((H)\), **saturation** \((S)\), and **value** \((V)\). We derive this three-dimensional color space by relating the HSV parameters to the directions in the RGB cube. If we imagine viewing the cube along the diagonal from the white vertex to the origin (black), we see an outline of the cube that has the hexagon shape shown in Figure 14. The boundary of the hexagon represents the various hues, and it is used as the top of the HSV hexcone (Figure 15). In HSV space, saturation \( S \) is measured along a horizontal axis, and the value parameter \( V \) is measured along a vertical axis through the center of the hexcone.

Hue is represented as an angle about the vertical axis, ranging from 0° at red through 360°. Vertices of the hexagon are separated by 60° intervals. Yellow is at 60°, green at 120°, and cyan (opposite the red point) is at \( H = 180° \). Complementary colors are 180° apart.

Saturation parameter \( S \) is used to designate the purity of a color. A pure color (spectral color) has the value \( S = 1.0 \), and decreasing \( S \) values tend toward the grayscale line \((S = 0)\) at the center of the hexcone.
Value $V$ varies from 0 at the apex of the hexcone to 1.0 at the top plane. The apex of the hexcone is the black point. At the top plane, colors have their maximum intensity. When $V = 1.0$ and $S = 1.0$, we have the pure hues. Parameter values for the white point are $V = 1.0$ and $S = 0$.

For most users, this is a more convenient model for selecting colors. Starting with a selection for a pure hue, which specifies the hue angle $H$ and sets $V = S = 1.0$, we describe the color we want in terms of adding either white or black to the pure hue. Adding black decreases the setting for $V$ while $S$ is held constant. To get a dark blue, for instance, $V$ could be set to 0.4 with $S = 1.0$ and $H = 240^\circ$. Similarly, when white is to be added to the selected hue, parameter $S$ is decreased while keeping $V$ constant. A light blue could be designated with $S = 0.3$ while $V = 1.0$ and $H = 240^\circ$. By adding some black and some white, we decrease both $V$ and $S$. An interface for this model typically presents the HSV parameter choices in a color palette containing sliders and a color wheel.

**Selecting Shades, Tints, and Tones**

Color regions for selecting shades, tints, and tones are represented in the cross-sectional plane of the HSV hexcone shown in Figure 16. Adding black to a spectral color decreases $V$ along the side of the hexcone toward the black point. Thus, various shades are represented with the values $S = 1.0$ and $0.0 \leq V \leq 1.0$. Adding white to spectral colors produces the tints across the top plane of the hexcone, where parameter values are $V = 1.0$ and $0 \leq S \leq 1.0$. Various tones are obtained by adding both black and white to spectral colors, which generates color points within the triangular cross-sectional area of the hexcone.

The human eye can distinguish about 128 different hues and about 130 different tints (saturation levels). For each of these, a number of shades (value settings) can be detected, depending on the hue selected. About 23 shades are discernible with yellow colors, and about 16 different shades can be seen at the blue end of the spectrum. This means that we can distinguish about $128 \times 130 \times 23 = 382,720$ different colors. For most graphics applications, 128 hues, 8 saturation levels, and
16 value settings are sufficient. With this range of parameters in the HSV color model, 16,384 colors are available to a user. These color values can be stored in 14 bits per pixel, or we could use color-lookup tables and fewer bits per pixel.

Transformations Between HSV and RGB Color Spaces

To determine the operations required for the transformations between the HSV and RGB spaces, we first consider how the HSV hexcone can be constructed from the RGB cube. The diagonal of the RGB cube from black (the origin) to white corresponds to the $V$ axis of the hexcone. Also, each subcube of the RGB cube corresponds to a hexagonal cross-sectional area of the hexcone. At any cross section, all sides of the hexagon and all radial lines from the $V$ axis to any vertex have the value $V$. Thus, for any set of RGB values, $V$ is equal to the value of the maximum RGB component. The HSV point corresponding to this set of RGB values lies on the hexagonal cross section at value $V$. Parameter $S$ is then determined as the relative distance of this point from the $V$ axis. Parameter $H$ is determined by calculating the relative position of the point within each sextant of the hexagon. An algorithm for mapping any set of RGB values into the corresponding HSV values is given in the following procedure:

```cpp
class rgbSpace {public: float r, g, b;};
class hsvSpace {public: float h, s, v;};

const float noHue = -1.0;
inline float min(float a, float b) {return (a < b)? a : b;}
inline float max(float a, float b) {return (a > b)? a : b;}

void rgbToHsv (rgbSpace& rgb, hsvSpace& hsv)
{
    /* RGB and HSV values are in the range from 0 to 1.0 */
    float minRGB = min (r, min (g, b)), maxRGB = max (r, max (g, b));
    float deltaRGB = maxRGB - minRGB;

    v = maxRGB;
    if (maxRGB != 0.0)
        s = deltaRGB / maxRGB;
    else
        s = 0.0;
    ```
if (s <= 0.0)
    h = noHue;
else {
    if (r == maxRGB)
        h = (g - b) / deltaRGB;
    else
        if (g == maxRGB)
            h = 2.0 + (b - r) / deltaRGB;
        else
            if (b == maxRGB)
                h = 4.0 + (r - g) / deltaRGB;
            h *= 60.0;
    if (h < 0.0)
        h += 360.0;
    h /= 360.0;
}

We obtain the transformation from HSV space to RGB space by determining
the inverse of the operations in the preceding procedure. These inverse operations
are carried out for each sextant of the hexcone, and the resulting transformation
equations are summarized in the following algorithm:

class rgbSpace {public: float r, g, b;};
class hsvSpace {public: float h, s, v;};

void hsvToRgb (hsvSpace& hsv, rgbSpace& rgb)
{
    /* HSV and RGB values are in the range from 0 to 1.0 */
    int k
    float aa, bb, cc, f;

    if (s <= 0.0)
        r = g = b = v; // Have gray scale if s = 0.
    else {
        if (h == 1.0)
            h = 0.0;
        h *= 6.0;
        k = floor (h);
        f = h - k;
        aa = v * (1.0 - s);
        bb = v * (1.0 - (s * f));
        cc = v * (1.0 - (s * (1.0 - f)));
        switch (k)
        {
        case 0: r = v; g = cc; b = aa; break;
        case 1: r = bb; g = v; b = aa; break;
        case 2: r = aa; g = v; b = cc; break;
        case 3: r = aa; g = bb; b = v; break;
        case 4: r = cc; g = aa; b = v; break;
        case 5: r = v; g = aa; b = bb; break;
        }
    }
}
Another model based on intuitive color parameters is the HLS system used by the Tektronix Corporation. This color space has the double-cone representation shown in Figure 17. The three parameters in this color model are called hue (H), lightness (L), and saturation (S).

Hue has the same meaning as in the HSV model. It specifies an angle about the vertical axis that locates a hue (spectral color). In this model, \( H = 0 \) corresponds to blue. The remaining colors are specified around the perimeter of the cone in the same order as in the HSV model. Magenta is located at \( H = 60^\circ \), red is at \( H = 120^\circ \), and cyan is at \( H = 300^\circ \). Again, complementary colors are 180\(^\circ\) apart on the double cone.

The vertical axis in this model is called lightness, L. At \( L = 0 \), we have black, and at \( L = 1.0 \), we have white. Grayscale values are along the \( L \) axis, and the pure colors lie on the \( L = 0.5 \) plane.

Saturation parameter S again specifies the purity of a color. This parameter varies from 0 to 1.0, and pure colors are those for which \( S = 1.0 \) and \( L = 0.5 \). As \( S \) decreases, more white is added to a color. The grayscale line is at \( S = 0 \).

To specify a color, we begin by selecting hue angle H. Then a particular shade, tint, or tone for that hue is obtained by adjusting parameters \( L \) and \( S \). We obtain a lighter color by increasing \( L \), and we obtain a darker color by decreasing \( L \). When \( S \) is decreased, the spatial color point moves toward the grayscale line.

**Figure 17**  
The HLS double cone.
9 Color Selection and Applications

A graphics package can provide color capabilities in a way that aids us in making color selections. For example, an interface can contain sliders and color wheels instead of requiring that all color specifications be provided as numerical values for the RGB components. In addition, some aids can be provided for choosing harmonious color combinations and for basic color selection guidelines.

One method for obtaining a set of coordinating colors is to generate the color combinations from a small subspace of a color model. If colors are selected at regular intervals along any straight line within the RGB or CMY cube, for example, we can expect to obtain a set of well-matched colors. Randomly selected hues can be expected to produce harsh and clashing color combinations. Another consideration in color displays is the fact that we perceive colors at different depths. This occurs because our eyes focus on colors according to their frequency. Blues, in particular, tend to recede. Displaying a blue pattern next to a red pattern can cause eye fatigue, because we continually need to refocus when our attention is switched from one area to the other. This problem can be reduced by separating these colors or by using colors from one-half or less of the color hexagon in the HSV model. With this technique, a display contains either blues and greens or reds and yellows.

As a general rule, the use of a smaller number of colors produces a better-looking display than one with a large number of colors. Also, tints and shades tend to blend better than the pure hues. For a background, gray or the complement of one of the foreground colors is usually best.

10 Summary

Light can be described as electromagnetic radiation with a certain energy distribution propagating through space, and the color components of light correspond to frequencies within a narrow band of the electromagnetic spectrum. However, light exhibits other properties, and we characterize the different aspects of light using a variety of parameters. With the light theories for wave-particle duality, we can explain the physical features of visible radiation. And we quantify our perceptions of a light source using terms such as dominant frequency (hue), luminance (brightness), and purity (saturation). Hue and purity are referred to collectively as the chromaticity properties of a color.

We also use color models to explain the effects of combining light sources. One method for defining a color model is to specify a set of two or more primary colors that are combined to produce various other colors. However, no finite set of primary colors is capable of producing all colors or describing all features of color. The set of colors that can be generated by a set of primaries is called a color gamut. Two colors that combine to produce white light are called complementary colors.

In 1931, the International Commission on Illumination (CIE) adopted a set of three hypothetical color-matching functions as a standard. This set of colors is referred to as the XYZ model, where X, Y, and Z represent the amounts of each color needed to match any color in the electromagnetic spectrum. The color-matching functions are structured so that all functions are positive and the Y amount for any color represents the luminance value. Normalized X and Y values, called x and y, are used to plot positions for all spectral colors on the CIE chromaticity diagram. We can use the chromaticity diagram to compare color gamuts for different...
color models, to identify complementary colors, and to determinant dominant frequency and purity for a specified color.

Other color models based on a set of three primaries are the RGB, YIQ, and CMY models. We use the RGB model to describe colors that are displayed on a video monitor. The YIQ model is used to describe the composite video signal for television broadcasting. And the CMY model is used to describe color on hard-copy devices.

User interfaces often provide intuitive color models, such as the HSV and HLS models, for selecting color values. With these models, we specify a color as a mixture of a selected hue and certain amounts of white and black. Adding black produces color shades, adding white produces tints, and adding both black and white produces tones.

Color selection is an important factor in the design of effective displays. To avoid clashing color combinations, we can choose adjacent colors in a display that do not differ greatly in dominant frequency. Also, we can select color combinations from a small subspace of a color model. As a general rule, a small number of color combinations formed with tints and shades, rather than pure hues, results in a more harmonious color display.

REFERENCES


EXERCISES
1 Derive the expressions for converting RGB color parameters to HSV values.
2 Derive the expressions for converting HSV color values to RGB values.
3 Design an interactive procedure that allows selection of HSV color parameters from a displayed menu; then, the HSV values are to be converted to RGB values for storage in a frame buffer.
4 Write a program to select colors using a set of three sliders to select values for the HSV color parameters.
5 Modify the program in the preceding exercise to display the numeric values for the RGB components of a selected color.
6 Modify the program in the preceding exercise to display the RGB color components and the combined color in small display windows.
7 Derive expressions for converting RGB color values to HLS color parameters.
8 Derive expressions for converting HLS color values to RGB values.
9 Design an interactive procedure that allows selection of HLS color parameters from a displayed menu; then, the HLS values are to be converted to RGB values for storage in a frame buffer.
10 Write a program that will produce a set of colors that are linearly interpolated between any two specified positions in RGB space.
11 Write an interactive routine for selecting color values from within a specified subspace of RGB space.
12 Write a program that will produce a set of colors that are linearly interpolated between any two specified positions in HSV space.
13 Write an interactive routine for selecting color values from within a specified subspace of HSV space.
14 Write a program that will produce a set of colors that are linearly interpolated between any two specified positions in HLS space.
15 Write an interactive routine for selecting color values from within a specified subspace of HLS space.
16 Write a program to display two adjacent RGB color rectangles. Fill one rectangle with a set of randomly selected RGB color points, and fill
the other rectangle with a set of color points
that are selected from a small RGB subspace.
Experiment with different random selections and
different subspaces to compare the two color
patterns.

17 Display the two color rectangles in the preceding exercise using color selections from either the HSV or the HLS color space.

18 Write a program that will produce a randomly selected color from within a color gamut specified by three positions in RGB space.

19 Write a program that will produce a randomly selected color from within a color gamut specified by three positions in HSV space.

20 Write a program that will produce a randomly selected color from within a color gamut specified by three positions in HLS space.

IN MORE DEPTH

1 Write a routine that takes in a pixel position of a scene in your application and a color space identifier and returns a vector representing the color value of that pixel in the selected color space. The routine should produce correct output for the RGB, CMY, HSV, and HLS color spaces.

2 Use the routine developed in the previous exercise to write another routine that outputs to a file a bitmap of a scene in your application in a specified color space (either RGB, CMY, HSV, or HLS). That is, the routine should take in a color space identifier, call the routine in the previous exercise on each pixel to obtain the color of that pixel in the specified color space, and write the color value vector of each pixel out to a file. Each color value vector should appear on a separate line, and pixels should be processed in row-major order.
Color Models and Color Applications Color Plates

Color Plate 22
Two views of the RGB color cube. View (a) is along the gray-scale diagonal from white to black, and view (b) is along the gray-scale diagonal from black to white.
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Although we can construct programs and provide input data using the methods and program commands discussed in the previous chapters, it is often useful to be able to specify graphical input interactively. During the execution of a program, for example, we might want to change the position of the camera or the location of an object in a scene by pointing to a screen position, or we might want to change animation parameters using menu selections. In design applications, control-point coordinates for spline constructions are chosen interactively, and pictures are often constructed using interactive painting or drawing methods. There are several kinds of data that are used by a graphics program, and a variety of interactive input methods have been devised for processing these data values. In addition, interfaces for systems now involve extensive interactive graphics, including display windows, icons, menus, and a mouse or other cursor-control devices.
1 Graphical Input Data

Graphics programs use several kinds of input data, such as coordinate positions, attribute values, character-string specifications, geometric-transformation values, viewing conditions, and illumination parameters. Many graphics packages, including the International Standards Organization (ISO) and American National Standards Institute (ANSI) standards, provide an extensive set of input functions for processing such data. But input procedures require interaction with display-window managers and specific hardware devices. Therefore, some graphics systems, particularly those that provide mainly device-independent functions, often include relatively few interactive procedures for dealing with input data.

A standard organization for input procedures in a graphics package is to classify the functions according to the type of data that is to be processed by each function. This scheme allows any physical device, such as a keyboard or a mouse, to input any data class, although most input devices can handle some data types better than others.

2 Logical Classification of Input Devices

When input functions are classified according to data type, any device that is used to provide the specified data is referred to as a logical input device for that data type. The standard logical input-data classifications are

- **LOCATOR**: A device for specifying one coordinate position.
- **STROKE**: A device for specifying a set of coordinate positions.
- **STRING**: A device for specifying text input.
- **VALUATOR**: A device for specifying a scalar value.
- **CHOICE**: A device for selecting a menu option.
- **PICK**: A device for selecting a component of a picture.

Locator Devices

Interactive selection of a coordinate point is usually accomplished by positioning the screen cursor at some location in a displayed scene, although other methods, such as menu options, could be used in certain applications. We can use a mouse, touchpad, joystick, trackball, spaceball, thumbwheel, dial, hand cursor, or digitizer stylus for screen-cursor positioning. In addition, various buttons, keys, or switches can be used to indicate processing options for the selected location.

Keyboards are used for locator input in several ways. A general-purpose keyboard usually has four cursor-control keys that move the screen cursor up, down, left, and right. With an additional four keys, we can move the cursor diagonally as well. Rapid cursor movement is accomplished by holding down the selected cursor key. Sometimes a keyboard includes a touchpad, joystick, trackball, or other device for positioning the screen cursor. For some applications, it may also be convenient to use a keyboard to type in numerical values or other codes to indicate coordinate positions.

Other devices, such as a light pen, have also been used for interactive input of coordinate positions. But light pens record screen positions by detecting light from the screen phosphors, and this requires special implementation procedures.
Stroke Devices
This class of logical devices is used to input a sequence of coordinate positions, and the physical devices used for generating locator input are also used as stroke devices. Continuous movement of a mouse, trackball, joystick, or hand cursor is translated into a series of input coordinate values. The graphics tablet is one of the more common stroke devices. Button activation can be used to place the tablet into “continuous” mode. As the cursor is moved across the tablet surface, a stream of coordinate values is generated. This procedure is used in paintbrush systems to generate drawings using various brush strokes. Engineering systems also use this process to trace and digitize layouts.

String Devices
The primary physical device used for string input is the keyboard. Character strings in computer-graphics applications are typically used for picture or graph labeling.

Other physical devices can be used for generating character patterns for special applications. Individual characters can be sketched on the screen using a stroke or locator-type device. A pattern recognition program then interprets the characters using a stored dictionary of predefined patterns.

Valuator Devices
We can employ valuator input in a graphics program to set scalar values for geometric transformations, viewing parameters, and illumination parameters. In some applications, scalar input is also used for setting physical parameters such as temperature, voltage, or stress-strain factors.

A typical physical device used to provide valuator input is a panel of control dials. Dial settings are calibrated to produce numerical values within some predefined range. Rotary potentiometers convert dial rotation into a corresponding voltage, which is then translated into a number within a defined scalar range, such as −10.5 to 25.5. Instead of dials, slide potentiometers are sometimes used to convert linear movements into scalar values.

Any keyboard with a set of numeric keys can be used as a valuator device. Although dials and slide potentiometers are more efficient for fast input.

Joysticks, trackballs, tablets, and other interactive devices can be adapted for valuator input by interpreting pressure or movement of the device relative to a scalar range. For one direction of movement, say left to right, increasing scalar values can be input. Movement in the opposite direction decreases the scalar input value. Selected values are usually echoed on the screen for verification.

Another technique for providing valuator input is to display graphical representations of sliders, buttons, rotating scales, and menus on the video monitor. Cursor positioning, using a mouse, joystick, spaceball, or other device, can be used to select a value on one of thesevaluators. As a feedback mechanism for the user, selected values are usually displayed in text or color fields elsewhere within the graphical display belonging to the application.

Choice Devices
Menus are typically used in graphics programs to select processing options, parameter values, and object shapes that are to be used in constructing a picture. Commonly used choice devices for selecting a menu option are cursor-positioning devices such as a mouse, trackball, keyboard, touch panel, or button box.

Keyboard function keys or separate button boxes are often used to enter menu selections. Each button or function key is programmed to select a particular
operation or value, although preset buttons or keys are sometimes included on an input device.

For screen selection of listed menu options, we use a cursor-positioning device. When a screen-cursor position \((x, y)\) is selected, it is compared to the coordinate extents of each listed menu item. A menu item with vertical and horizontal boundaries at the coordinate values \(x_{\text{min}}, x_{\text{max}}, y_{\text{min}},\) and \(y_{\text{max}}\) is selected if the input coordinates satisfy the inequalities

\[
\begin{align*}
  x_{\text{min}} &\leq x \leq x_{\text{max}}, \\
  y_{\text{min}} &\leq y \leq y_{\text{max}}
\end{align*}
\]  

For larger menus with relatively few options displayed, a touch panel is commonly used. A selected screen position is compared to the coordinate extents of the individual menu options to determine what process is to be performed.

Alternate methods for choice input include keyboard and voice entry. A standard keyboard can be used to type in commands or menu options. For this method of choice input, some abbreviated format is useful. Menu listings can be numbered or given short identifying names. A similar encoding scheme can be used with voice input systems. Voice input is particularly useful when the number of options is small (20 or fewer).

Pick Devices

We use a pick device to select a part of a scene that is to be transformed or edited in some way. Several different methods can be used to select a component of a displayed scene, and any input mechanism used for this purpose is classified as a pick device. Most often, pick operations are performed by positioning the screen cursor. Using a mouse, joystick, or keyboard, for example, we can perform picking by positioning the screen cursor and pressing a button or key to record the pixel coordinates. This screen position can then be used to select an entire object, a facet of a tessellated surface, a polygon edge, or a vertex. Other pick methods include highlighting schemes, selecting objects by name, or a combination of methods.

Using the cursor-positioning approach, a pick procedure could map a selected screen position to a world-coordinate location using the inverse viewing and geometric transformations that were specified for the scene. Then, the world-coordinate position can be compared to the coordinate extents of objects. If the pick position is within the coordinate extents of a single object, the pick object has been identified. The object name, coordinates, or other information about the object can then be used to apply the desired transformation or editing operations. But if the pick position is within the coordinate extents of two or more objects, further testing is necessary. Depending on the type of object to be selected and the complexity of a scene, several levels of search may be required to identify the pick object. For example, if we are attempting to pick a sphere whose coordinate extents overlap the coordinate extents of some other three-dimensional object, the pick position could be compared to the coordinate extents of the individual surface facets of the two objects. If this test fails, the coordinate extents of individual line segments can be tested.

When coordinate-extent tests do not uniquely identify a pick object, the distances from the pick position to individual line segments could be computed. Figure 1 illustrates a pick position that is within the coordinate extents of two line segments. For a two-dimensional line segment with pixel endpoint coordinates \((x_1, y_1)\) and \((x_2, y_2)\), the perpendicular distance squared from a pick position \((x, y)\) to the line is calculated as

\[
d^2 = \frac{[(\Delta x)(y - y_1) - (\Delta y)(x - x_1)]^2}{\Delta x^2 + \Delta y^2}
\]
where $\Delta x = x_2 - x_1$ and $\Delta y = y_2 - y_1$. Other methods, such as comparing distances to endpoint positions, have been proposed to simplify the line-picking operations.

Pick procedures can be simplified if coordinate-extent testing is not carried out for the surface facets and line segments of an object. When the pick position is within the coordinate extents of two or more objects, the pick procedures can simply return a list of all candidate pick objects.

Another picking technique is to associate a **pick window** with a selected cursor position. The pick window is centered on the cursor position, as shown in Figure 2, and clipping procedures are used to determine which objects intersect the pick window. For line picking, we can set the pick-window dimensions $w$ and $h$ to very small values, so that only one line segment intersects the pick window. Some graphics packages implement three-dimensional picking by reconstructing a scene using the viewing and projection transformations with the pick window as the clipping window. Nothing is displayed from this reconstruction, but clipping procedures are applied to determine which objects are within the pick view volume. A list of information for each object in the pick view volume can then be returned for processing. This list can contain information such as object name and depth range, where the depth range could be used to select the nearest object in the pick view volume.

Highlighting can also be used to facilitate picking. One way to do this is to successively highlight those objects whose coordinate extents overlap a pick position (or pick window). As each object is highlighted, a user could issue a “reject” or “accept” action using keyboard keys. The sequence stops when the user accepts a highlighted object as the pick object. Picking could also be accomplished simply by successively highlighting all objects in the scene without selecting a cursor position. The highlighting sequence can be initiated with a button or function key, and a second button can be used to stop the process when the desired object is highlighted. If very many objects are to be searched in this way, additional buttons can be used to speed up the highlighting process. One button initiates a rapid successive highlighting of structures. A second button is activated to stop the process, and a third button is used to back up slowly through the highlighting process. Finally, a stop button could be pressed to complete the pick procedure.

If picture components can be selected by name, keyboard input can be used to pick an object. This is a straightforward, but less interactive, pick-selection method. Some graphics packages allow picture components to be named at various levels down to the individual primitives. Descriptive names can be used to help a user in the pick process, but this approach has drawbacks. It is generally slower than interactive picking on the screen, and a user will probably need prompts to remember the various structure names.

### 3 Input Functions for Graphical Data

Graphics packages that use the logical classification for input devices provide several functions for selecting devices and data classes. These functions allow a user to specify the following options:

- The input interaction mode for the graphics program and the input devices. Either the program or the devices can initiate data entry, or both can operate simultaneously.

- Selection of a physical device that is to provide input within a particular logical classification (for example, a tablet used as a stroke device).

- Selection of the input time and device for a particular set of data values.
**Input Modes**

Some input functions in an interactive graphics system are used to specify how the program and input devices should interact. A program could request input at a particular time in the processing (request mode), or an input device could independently provide updated input (sample mode), or the device could independently store all collected data (event mode).

In **request mode**, the application program initiates data entry. When input values are requested, processing is suspended until the required values are received. This input mode corresponds to the typical input operation in a general programming language. The program and the input devices operate alternately. Devices are put into a wait state until an input request is made; then the program waits until the data are delivered.

In **sample mode**, the application program and input devices operate independently. Input devices may be operating at the same time that the program is processing other data. New values obtained from the input devices replace previously input data values. When the program requires new data, it samples the current values that have been stored from the device input.

In **event mode**, the input devices initiate data input to the application program. The program and the input devices again operate concurrently, but now the input devices deliver data to an input queue, also called an **event queue**. All input data is saved. When the program requires new data, it goes to the data queue.

Typically, any number of devices can be operating at the same time in sample and event modes. Some can be operating in sample mode, while others are operating in event mode. But only one device at a time can deliver input in request mode.

Other functions in the input library are used to specify physical devices for the logical data classes. The input procedures in an interactive package can involve complicated processing for some kinds of input. For instance, to obtain a world-coordinate position, the input procedures must process an input screen location back through the viewing and other transformations to the original world-coordinate description of a scene. This processing also involves information from the display-window routines.

**Echo Feedback**

Requests can usually be made in an interactive input program for an echo of input data and associated parameters. When an echo of the input data is requested, it is displayed within a specified screen area. Echo feedback can include, for example, the size of the pick window, the minimum pick distance, the type and size of a cursor, the type of highlighting to be employed during pick operations, the range (minimum and maximum) for valuator input, and the resolution (scale) for valuator input.

**Callback Functions**

For device-independent graphics packages, a limited set of input functions can be provided in an auxiliary library. Input procedures can then be handled as callback functions that interact with the system software. These functions specify what actions are to be taken by a program when an input event occurs. Typical input events are moving a mouse, pressing a mouse button, or pressing a key on the keyboard.
Interactive Picture-Construction Techniques

A variety of interactive methods are often incorporated into a graphics package as aids in the construction of pictures. Routines can be provided for positioning objects, applying constraints, adjusting the sizes of objects, and designing shapes and patterns.

Basic Positioning Methods

We can interactively choose a coordinate position with a pointing device that records a screen location. How the position is used depends on the selected processing option. The coordinate location could be an endpoint position for a new line segment, or it could be used to position some object—for instance, the selected screen location could reference a new position for the center of a sphere; or the location could be used to specify the position for a text string, which could begin at that location or it could be centered on that location. As an additional positioning aid, numeric values for selected positions can be echoed on the screen. With the echoed coordinate values as a guide, a user could make small interactive adjustments in the coordinate values using dials, arrow keys, or other devices.

Dragging

Another interactive positioning technique is to select an object and drag it to a new location. Using a mouse, for instance, we position the cursor at the object position, press a mouse button, move the cursor to a new position, and release the button. The object is then displayed at the new cursor location. Usually, the object is displayed at intermediate positions as the screen cursor moves.

Constraints

Any procedure for altering input coordinate values to obtain a particular orientation or alignment of an object is called a constraint. For example, an input line segment can be constrained to be horizontal or vertical, as illustrated in Figures 3 and 4. To implement this type of constraint, we compare the input coordinate values at the two endpoints. If the difference in the \( y \) values of the two endpoints is smaller than the difference in the \( x \) values, a horizontal line is displayed. Otherwise, a vertical line is drawn. The horizontal-vertical constraint is useful, for instance, in forming network layouts, and it eliminates the need for precise positioning of endpoint coordinates.

![Figure 3](image-url)
Other kinds of constraints can be applied to input coordinates to produce a variety of alignments. Lines could be constrained to have a particular slant, such as $45^\circ$, and input coordinates could be constrained to lie along predefined paths, such as circular arcs.

**Grids**

Another kind of constraint is a rectangular grid displayed in some part of the screen area. With an activated grid constraint, input coordinates are rounded to the nearest grid intersection. Figure 5 illustrates line drawing using a grid. Each of the cursor positions in this example is shifted to the nearest grid intersection point, and a line is drawn between these two grid positions. Grids facilitate object constructions, because a new line can be joined easily to a previously drawn line by selecting any position near the endpoint grid intersection of one end of the displayed line. Spacing between grid lines is often an option, and partial grids or grids with different spacing could be used in different screen areas.

**Rubber-Band Methods**

Line segments and other basic shapes can be constructed and positioned using rubber-band methods that allow the sizes of objects to be interactively stretched or contracted. Figure 6 demonstrates a rubber-band method for interactively specifying a line segment. First, a fixed screen position is selected for one endpoint of the line. Then, as the cursor moves around, the line is displayed from the start position to the current position of the cursor. The second endpoint of the line is input when a button or key is pressed. Using a mouse, we construct a rubber-band line while pressing a mouse key. When the mouse key is released, the line display is completed.
We can use similar rubber-band methods to construct rectangles, circles, and other objects. Figure 7 demonstrates rubber-band construction of a rectangle, and Figure 8 shows a rubber-band circle construction. We can implement rubber-band constructions in various ways. For example, the shape and size of a rectangle can be adjusted by independently moving only the top edge of the rectangle, or the bottom edge, or one of the side edges.

Gravity Field

In the construction of figures, we sometimes need to connect lines at positions between endpoints that are not at grid intersections. Because exact positioning of the screen cursor at the connecting point can be difficult, a graphics package can include a procedure that converts any input position near a line segment into a position on the line using a gravity field area around the line. Any selected position within the gravity field of a line is moved (“gravitated”) to the nearest position on the line. A gravity field area around a line is illustrated with the shaded region shown in Figure 9.

Gravity fields around the line endpoints are enlarged to make it easier for a designer to connect lines at their endpoints. Selected positions in one of the circular areas of the gravity field are attracted to the endpoint in that area. The size of gravity fields is chosen large enough to aid positioning, but small enough to reduce chances of overlap with other lines. If many lines are displayed, gravity areas can overlap, and it may be difficult to specify points correctly. Normally, the boundary for the gravity field is not displayed.

Interactive Painting and Drawing Methods

Options for sketching, drawing, and painting come in a variety of forms. Straight lines, polygons, and circles can be generated with methods discussed in the
previous sections. Curve-drawing options can be provided using standard curve shapes, such as circular arcs and splines, or with freehand sketching procedures. Splines are interactively constructed by specifying a set of control points or a freehand sketch that gives the general shape of the curve. Then the system fits the set of points with a polynomial curve. In freehand drawing, curves are generated by following the path of a stylus on a graphics tablet or the path of the screen cursor on a video monitor. Once a curve is displayed, the designer can alter the curve shape by adjusting the positions of selected points along the curve path.

Line widths, line styles, and other attribute options are also commonly found in painting and drawing packages. Various brush styles, brush patterns, color combinations, object shapes, and surface texture patterns are also available on many systems, particularly those designed as artists’ workstations. Some paint systems vary the line width and brush strokes according to the pressure of the artist’s hand on the stylus. Color Plate 23 shows a window and menu system used with a painting package that allows an artist to select variations of a specified object shape, different surface textures, and a variety of lighting conditions for a scene.

5 Virtual-Reality Environments

A typical virtual-reality environment is illustrated in Color Plate 24. Interactive input is accomplished in this environment with a data glove, which is capable of grasping and moving objects displayed in a virtual scene. The computer-generated scene is displayed through a head-mounted viewing system as a stereographic projection. Tracking devices compute the position and orientation of the headset and data glove relative to the object positions in the scene. With this system, a user can move through the scene and rearrange object positions with the data glove.

Another method for generating virtual scenes is to display stereographic projections on a raster monitor, with the two stereographic views displayed on alternate refresh cycles. The scene is then viewed through stereographic glasses. Interactive object manipulations can again be accomplished with a data glove and a tracking device to monitor the glove position and orientation relative to the position of objects in the scene.

6 OpenGL Interactive Input-Device Functions

Interactive device input in an OpenGL program is handled with routines in the OpenGL Utility Toolkit (GLUT), because these routines need to interface with a window system. In GLUT, we have functions to accept input from standard devices, such as a mouse or a keyboard, as well as from tablets, space balls, button boxes, and dials. For each device, we specify a procedure (the call back function) that is to be invoked when an input event from that device occurs. These GLUT commands are placed in the main procedure along with the other GLUT statements. In addition, a combination of functions from the basic library and the GLU library can be used with the GLUT mouse function for pick input.
GLUT Mouse Functions

We use the following function to specify ("register") a procedure that is to be called when the mouse pointer is in a display window and a mouse button is pressed or released:

\[ \text{glutMouseFunc (mouseFcn)}; \]

This mouse callback procedure, which we named \text{mouseFcn}, has four arguments:

\[ \text{void mouseFcn (GLint button, GLint action, GLint xMouse, GLint yMouse)} \]

Parameter \text{button} is assigned a GLUT symbolic constant that denotes one of the three mouse buttons, and parameter \text{action} is assigned a symbolic constant that specifies which button action we want to use to trigger the mouse activation event. Allowable values for \text{button} are \text{GLUT_LEFT_BUTTON}, \text{GLUT_MIDDLE_BUTTON}, and \text{GLUT_RIGHT_BUTTON}. (If we have only a two-button mouse, then we use just the left-button and right-button designations; with a one-button mouse, we can assign parameter \text{button} only the value \text{GLUT_LEFT_BUTTON}.) Parameter \text{action} can be assigned either \text{GLUT_DOWN} or \text{GLUT_UP}, depending on whether we want to initiate an action when we press a mouse button or when we release it. When procedure \text{mouseFcn} is invoked, the display-window location of the mouse cursor is returned as the coordinate position \((xMouse, yMouse)\). This location is relative to the top-left corner of the display window, so that \text{xMouse} is the pixel distance from the left edge of the display window and \text{yMouse} is the pixel distance down from the top of the display window.

By activating a mouse button while the screen cursor is within the display window, we can select a position for displaying a primitive such as a single point, a line segment, or a fill area. We could also use the mouse as a pick device by comparing the returned screen position with the coordinate extents of displayed objects in a scene. However, OpenGL does provide routines for using the mouse as a pick device, and we discuss these routines in a later section.

As a simple example of the use of the \text{glutMouseFunc} routine, the following program plots a red point, with a point size equal to 3, at the position of the mouse cursor in the display window, each time that we press the left mouse button. Because the coordinate origin for the OpenGL primitive functions is the lower-left corner of the display window, we need to flip the returned \text{yMouse} value in the procedure \text{mousePtPlot}.

```c
#include <GL/glut.h>

GLsizei winWidth = 400, winHeight = 300; // Initial display-window size.

void init (void)
{
    glClearColor (0.0, 0.0, 1.0, 1.0) // Set display-window color to blue.
    glMatrixMode (GL_PROJECTION);
    gluOrtho2D (0.0, 200.0, 0.0, 150.0);
}
```
void displayFcn (void) 
{
    glClear (GL_COLOR_BUFFER_BIT); // Clear display window.
    glColor3f (1.0, 0.0, 0.0); // Set point color to red.
    glPointSize (3.0); // Set point size to 3.0.
}

void winReshapeFcn (GLint newWidth, GLint newHeight) 
{
    /* Reset viewport and projection parameters */
    glViewport (0, 0, newWidth, newHeight);
    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ( );
    gluOrtho2D (0.0, GLdouble (newWidth), 0.0, GLdouble (newHeight));

    /* Reset display-window size parameters. */
    winWidth = newWidth;
    winHeight = newHeight;
}

void plotPoint (GLint x, GLint y) 
{
    glBegin (GL_POINTS);
    glVertex2i (x, y);
    glEnd ( );
}

void mousePtPlot (GLint button, GLint action, GLint xMouse, GLint yMouse) 
{
    if (button == GLUT_LEFT_BUTTON && action == GLUT_DOWN)
    plotPoint (xMouse, winHeight - yMouse);

    glFlush ( );
}

void main (int argc, char** argv) 
{
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Mouse Plot Points");

    init ( );
    glutDisplayFunc (displayFcn);
    glutReshapeFunc (winReshapeFcn);
    glutMouseFunc (mousePtPlot);

    glutMainLoop ( );
}
The next program example uses mouse input to select an endpoint position for a straight-line segment. Selected line segments are connected to demonstrate interactive construction of a polyline. Initially, two display-window locations must be selected with the left mouse button to generate the first line segment. Each subsequent position that we select adds another segment to the polyline.

```c
#include <GL/glut.h>

GLsizei winWidth = 400, winHeight = 300; // Initial display-window size.
GLint endPtCtr = 0; // Initialize line endpoint counter.

class scrPt {
public:
    GLint x, y;
};

void init (void)
{
    glClearColor (0.0, 0.0, 1.0, 1.0) // Set display-window color to blue.
    glMatrixMode (GL_PROJECTION);
    gluOrtho2D (0.0, 200.0, 0.0, 150.0);
}

void displayFcn (void)
{
    glClear (GL_COLOR_BUFFER_BIT);
}

void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    /* Reset viewport and projection parameters */
    glViewport (0, 0, newWidth, newHeight);
    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ( );
    gluOrtho2D (0.0, GLdouble (newWidth), 0.0, GLdouble (newHeight));

    /* Reset display-window size parameters. */
    winWidth = newWidth;
    winHeight = newHeight;
}

void drawLineSegment (scrPt endPt1, scrPt endPt2)
{
    glBegin (GL_LINES);
    glVertex2i (endPt1.x, endPt1.y);
    glVertex2i (endPt2.x, endPt2.y);
    glEnd ( );
}
void polyline (GLint button, GLint action, GLint xMouse, GLint yMouse) {
    static scrPt endPt1, endPt2;
    if (ptCtr == 0) {
        if (button == GLUT_LEFT_BUTTON && action == GLUT_DOWN) {
            endPt1.x = xMouse;
            endPt1.y = winHeight - yMouse;
            ptCtr = 1;
        } else
            if (button == GLUT_RIGHT_BUTTON) // Quit the program.
                exit (0);
    } else
        if (button == GLUT_LEFT_BUTTON && action == GLUT_DOWN) {
            endPt2.x = xMouse;
            endPt2.y = winHeight - yMouse;
            drawLineSegment (endPt1, endPt2);
            endPt1 = endPt2;
        } else
            if (button == GLUT_RIGHT_BUTTON) // Quit the program.
                exit (0);
    glFlush ( );
}

void main (int argc, char** argv) {
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Draw Interactive Polyline");
    init ( );
    glutDisplayFunc (displayFcn);
    glutReshapeFunc (winReshapeFcn);
    glutMouseFunc (polyline);
    glutMainLoop ( );
}

Another GLUT mouse routine that we can use is

glutMotionFunc (fcnDoSomething);

This routine invokes fcnDoSomething when the mouse is moved within the
display window with one or more buttons activated. The function that is invoked
in this case has two arguments:

    void fcnDoSomething (GLint xMouse, GLint yMouse)
where \((x_{\text{Mouse}}, y_{\text{Mouse}})\) is the mouse location in the display window relative to the top-left corner, when the mouse is moved with a button pressed.

Similarly, we can perform some action when we move the mouse within the display window without pressing a button:

\[
\text{glutPassiveMotionFunc (fcnDoSomethingElse)};
\]

Again, the mouse location is returned to \(\text{fcnDoSomethingElse}\) as coordinate position \((x_{\text{Mouse}}, y_{\text{Mouse}})\), relative to the top-left corner of the display window.

### GLUT Keyboard Functions

With keyboard input, we use the following function to specify a procedure that is to be invoked when a key is pressed:

\[
\text{glutKeyboardFunc (keyFcn)};
\]

The specified procedure has three arguments:

\[
\text{void keyFcn (GLubyte key, GLint x_{\text{Mouse}}, GLint y_{\text{Mouse}})}
\]

Parameter \(\text{key}\) is assigned a character value or the corresponding ASCII code. The display-window mouse location is returned as position \((x_{\text{Mouse}}, y_{\text{Mouse}})\) relative to the top-left corner of the display window. When a designated key is pressed, we can use the mouse location to initiate some action, independently of whether any mouse buttons are pressed.

In the following code, we present a simple curve-drawing procedure using keyboard input. A freehand curve is generated by moving the mouse within the display window while holding down the \(\text{“c”}\) key. This displays a sequence of red dots at each recorded mouse position. By slowly moving the mouse, we can obtain a solid curved line. Mouse buttons have no effect in this example.

```c
#include <GL/glut.h>

GLsizei winWidth = 400, winHeight = 300; // Initial display-window size.

void init (void)
{
    glClearColor (0.0, 0.0, 1.0, 1.0); // Set display-window color to blue.
    glMatrixMode (GL_PROJECTION);
    gluOrtho2D (0.0, 200.0, 0.0, 150.0);
}

void displayFcn (void)
{
    glClear (GL_COLOR_BUFFER_BIT); // Clear display window.
    glColor3f (1.0, 0.0, 0.0); // Set point color to red.
    glPointSize (3.0); // Set point size to 3.0.
    glutMainLoop();
}
```

void winReshapeFcn (GLint newWidth, GLint newHeight)
{
   /* Reset viewport and projection parameters */
   glViewport (0, 0, newWidth, newHeight);
   glMatrixMode (GL_PROJECTION);
   glLoadIdentity ( );
   gluOrtho2D (0.0, GLdouble (newWidth), 0.0, GLdouble (newHeight));

   /* Reset display-window size parameters. */
   winWidth = newWidth;
   winHeight = newHeight;
}

void plotPoint (GLint x, GLint y)
{
   glBegin (GL_POINTS);
   glVertex2i (x, y);
   glEnd ( );
}

/* Move cursor while pressing c key enables freehand curve drawing. */
void curveDrawing (GLubyte curvePlotKey, GLint xMouse, GLint yMouse)
{
   GLint x = xMouse;
   GLint y = winHeight - yMouse;
   switch (curvePlotKey)
   {
      case 'c':
         plotPoint (x, y);
         break;
      default:
         break;
   }
   glFlush ( );
}

void main (int argc, char** argv)
{
   glutInit (&argc, argv);
   glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
   glutInitWindowPosition (100, 100);
   glutInitWindowSize (winWidth, winHeight);
   glutCreateWindow ("Keyboard Curve-Drawing Example");

   init ( );
   glutDisplayFunc (displayFcn);
   glutReshapeFunc (winReshapeFcn);
   glutKeyboardFunc (curveDrawing);

   glutMainLoop ( );
}
For function keys, arrow keys, and other special-purpose keys, we can use the command

```c
glutSpecialFunc (specialKeyFcn);
```

The specified procedure has the same three arguments:

```c
void specialKeyFcn (GLint specialKey, GLint xMouse,
                    GLint yMouse)
```

but now parameter `specialKey` is assigned an integer-valued GLUT symbolic constant. To select a function key, we use one of the constants `GLUT_KEY_F1` through `GLUT_KEY_F12`. For the arrow keys, we use constants such as `GLUT_KEY_UP` and `GLUT_KEY_RIGHT`. Other keys can be designated using `GLUT_KEY_PAGE_DOWN`, `GLUT_KEY_HOME`, and similar constants for the page up, end, and insert keys. The backspace, delete, and escape keys can be designated with the `glutKeyboardFunc` routine using their ASCII codes, which are 8, 127, and 27, respectively.

An interactive program using the mouse, keyboard, and function keys is demonstrated in the following code. Mouse input is used to select a location for the lower-left corner of a red square. Keyboard input is used to scale the size of the square, and a new square is obtained with each click of the left mouse button.

```c
#include <GL/glut.h>
#include <stdlib.h>

GLsizei winWidth = 400, winHeight = 300; // Initial display-window size.
GLint edgeLength = 10; // Initial edge length for square.

void init (void)
{
    glClearColor (0.0, 0.0, 1.0, 1.0) // Set display-window color to blue.
    glMatrixMode (GL_PROJECTION);
    gluOrtho2D (0.0, 200.0, 0.0, 150.0);
}

void displayFcn (void)
{
    glClear (GL_COLOR_BUFFER_BIT); // Clear display window.
    glColor3f (1.0, 0.0, 0.0); // Set fill color to red.
}

void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    /* Reset viewport and projection parameters */
    glViewport (0, 0, newWidth, newHeight);
    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ();
    gluOrtho2D (0.0, GLdouble (newWidth), 0.0, GLdouble (newHeight));
}
/ * Reset display-window size parameters. */
winWidth = newWidth;
winHeight = newHeight;
}
/* Display a red square with a selected edge-length size. */
void fillSquare (GLint button, GLint action, GLint xMouse, GLint yMouse)
{
    GLint x1, y1, x2, y2;
    /* Use left mouse button to select a position for the */
    /* lower-left corner of the square. */
    if (button == GLUT_LEFT_BUTTON && action == GLUT_DOWN)
    {
        x1 = xMouse;
y1 = winHeight - yMouse;
x2 = x1 + edgeLength;
y2 = y1 + edgeLength;
glRecti (x1, y1, x2, y2);
    }
    else
    if (button == GLUT_RIGHT_BUTTON) // Use right mouse button to quit.
        exit (0);
glFlush ( );
}
/* Use keys 2, 3, and 4 to enlarge the square. */
void enlargeSquare (GLubyte sizeFactor, GLint xMouse, GLint yMouse)
{
    switch (sizeFactor)
    {
    case '2':
        edgeLength *= 2;
        break;
    case '3':
        edgeLength *= 3;
        break;
    case '4':
        edgeLength *= 4;
        break;
    default:
        break;
    }
}
/* Use function keys F2 and F4 for reduction factors 1/2 and 1/4. */
void reduceSquare (GLint reductionKey, GLint xMouse, GLint yMouse)
{
    switch (reductionKey)
    {
    case GLUT_KEY_F2:
        edgeLength /= 2;
        break;
GLUT Tablet Functions

Usually, tablet activation occurs only when the mouse cursor is in the display window. A button event for tablet input is then recorded with

```c
    glutTabletButtonFunc (tabletFcn);
```

and the arguments for the invoked function are similar to those for a mouse:

```c
    void tabletFcn (GLint tabletButton, GLint action, 
                   GLint xTablet, GLint yTablet)
```

We designate a tablet button with an integer identifier such as 1, 2, 3, and so on, and the button action is again specified with either GLUT_UP or GLUT_DOWN. The returned values xTablet and yTablet are the tablet coordinates. We can determine the number of available tablet buttons with the command

```c
    glutDeviceGet (GLUT_NUM_TABLET_BUTTONS);
```

Motion of a tablet stylus or cursor is processed with the following function:

```c
    glutTabletMotionFunc (tabletMotionFcn);
```

where the invoked function has the form

```c
    void tabletMotionFcn (GLint xTablet, GLint yTablet)
```

The returned values xTablet and yTablet give the coordinates on the tablet surface.
GLUT Spaceball Functions

We use the following function to specify an operation when a spaceball button is activated for a selected display window:

```c
glutSpaceballButtonFunc (spaceballFcn);
```

The callback function has two parameters:

```c
void spaceballFcn (GLint spaceballButton, GLint action)
```

Spaceball buttons are identified with the same integer values as a tablet, and parameter action is assigned either the value GLUT_UP or the value GLUT_DOWN. We can determine the number of available spaceball buttons with a call to `glutDeviceGet` using the argument GLUT_NUM_SPACEBALL_BUTTONS.

Translational motion of a spaceball, when the mouse is in the display window, is recorded with the function call

```c
glutSpaceballMotionFunc (spaceballTranlFcn);
```

The three-dimensional translation distances are passed to the invoked function as, for example:

```c
void spaceballTranslFcn (GLint tx, GLint ty, GLint tz)
```

These translation distances are normalized within the range from $-1000$ to $1000$.

Similarly, a spaceball rotation is recorded with

```c
glutSpaceballRotateFunc (spaceballRotFcn);
```

The three-dimensional rotation angles are then available to the callback function, as follows:

```c
void spaceballRotFcn (GLint thetaX, GLint thetaY, GLint thetaZ)
```

GLUT Button-Box Function

Input from a button box is obtained with the following statement:

```c
glutButtonBoxFunc (buttonBoxFcn);
```

Button activation is then passed to the invoked function:

```c
void buttonBoxFcn (GLint button, GLint action);
```

The buttons are identified with integer values, and the button action is specified as GLUT_UP or GLUT_DOWN.

GLUT Dials Function

A dial rotation can be recorded with the following routine:

```c
glutDialsFunc (dialsFcn);
```

In this case, we use the callback function to identify the dial and obtain the angular amount of rotation:

```c
void dialsFcn (GLint dial, GLint degreeValue);
```

Dials are designated with integer values, and the dial rotation is returned as an integer degree value.
OpenGL Picking Operations

In an OpenGL program, we can interactively select objects by pointing to screen positions. However, the picking operations in OpenGL are not straightforward. Basically, we perform picking using a designated pick window to form a revised view volume. We assign integer identifiers to objects in a scene, and the identifiers for those objects that intersect the revised view volume are stored in a pick-buffer array. Thus, to use the OpenGL pick features, we need to incorporate the following procedures into a program:

1. Create and display a scene.
2. Pick a screen position and, within the mouse callback function, do the following:
   - Set up a pick buffer.
   - Activate the picking operations (selection mode).
   - Initialize an ID name stack for object identifiers.
   - Save the current viewing and geometric-transformation matrix.
   - Specify a pick window for the mouse input.
   - Assign identifiers to objects and reprocess the scene using the revised view volume. (Pick information is then stored in the pick buffer.)
   - Restore the original viewing and geometric-transformation matrix.
   - Determine the number of objects that have been picked, and return to the normal rendering mode.
3. Process the pick information.

We can also use a modification of these procedures to select objects without interactive input from a mouse. This is accomplished by specifying the vertices for the revised view volume, instead of designating a pick window.

A pick-buffer array is set up with the command

```c
glSelectBuffer (pickBuffSize, pickBuffer);
```

Parameter `pickBuffer` designates an integer array with `pickBuffSize` elements. The `glSelectBuffer` function must be invoked before the OpenGL picking operations (selection mode) are activated. An integer information record is stored in pick-buffer array for each object that is selected with a single pick input. Several records of information can be stored in the pick buffer, depending on the size and location of the pick window. Each record in the pick buffer contains the following information:

1. The stack position of the object, which is the number of identifiers in the name stack, up to and including the position of the picked object.
2. The minimum depth of the picked object.
3. The maximum depth of the picked object.
4. The list of the identifiers in the name stack from the first (bottom) identifier to the identifier for the picked object.

The integer depth values stored in the pick buffer are the original values in the range from 0 to 1.0, multiplied by $2^{32} - 1$.

The OpenGL picking operations are activated with

```c
glRenderMode (GL_SELECT);
```
This places us in selection mode, which means that a scene is processed through the viewing pipeline but not stored in the frame buffer. A record of information for each object that would have been displayed in the normal rendering mode is placed in the pick buffer. In addition, this command returns the number of picked objects, which is equal to the number of information records in the pick buffer. To return to the normal rendering mode (the default), we invoke the `glRenderMode` routine using the argument `GL_RENDER`. A third option is the argument `GL_FEEDBACK`, which stores object coordinates and other information in a feedback buffer without displaying the objects. Feedback mode is used to obtain information about primitive types, attributes, and other parameters associated with the objects in a scene.

We use the following statement to activate the integer-ID name stack for the picking operations:

```
glInitNames ( );
```

The ID stack is initially empty, and this stack can be used only in selection mode. To place an unsigned integer value on the stack, we can invoke the following function:

```
glPushName (ID);
```

This places the value for parameter ID on the top of the stack and pushes the previous top name down to the next position in the stack. We can also simply replace the top of the stack using

```
glLoadName (ID);
```

but we cannot use this command to place a value on an empty stack. To eliminate the top of the ID stack, we issue the command

```
glPopName ( );
```

A pick window within a selected viewport is defined using the following GLU function:

```
gluPickMatrix (xPick, yPick, widthPick, heightPick, vpArray);
```

Parameters `xPick` and `yPick` give the double-precision, screen-coordinate location for the center of the pick window relative to the lower-left corner of the viewport. When these coordinates are given with mouse input, the mouse coordinates are relative to the upper-left corner, and thus we need to invert the input `yMouse` value. The double-precision values for the width and height of the pick window are specified with parameters `widthPick` and `heightPick`. Parameter `vpArray` designates an integer array containing the coordinate position and size parameters for the current viewport. We can obtain the viewport parameters using the `glGetIntegerv` function. This pick window is then used as a clipping window to construct a revised view volume for the viewing transformations. Information for objects that intersect this revised view volume is placed in the pick buffer.

We illustrate the OpenGL picking operations in the following program, which displays three color rectangles with the colors red, blue, and green. For this picking example, we use a $5 \times 5$ pick window, and the center of the pick window is given
with mouse input. Therefore, we need to invert the input \texttt{yMouse} value using
the viewport height, which is the fourth element of the array \texttt{vpArray}. The red
rectangle is assigned ID = 30, the blue rectangle is assigned ID = 10, and the green
rectangle is assigned ID = 20. Depending on the input mouse position, we can
pick no rectangles, one rectangle, two of the rectangles, or all three rectangles at
one time. The rectangle identifiers are entered into the ID stack in the color order:
red, blue, green. Therefore, when we process a picked rectangle, we could use
either its identifier or its stack position number. For example, if the stack position
number, which is the first item in the pick record, is 2, then we have picked the blue
rectangle and there are two rectangle identifiers listed at the end of the record.
Alternatively, we could use the last entry in the record, which is the identifier
for the picked object. In this example program, we simply list the contents of the
pick buffer. The rectangles are defined in the \textit{xy} plane, so all depth values are 0.
A sample output is given in Example 1 for a mouse input position that is near
the boundary between the red and green rectangles. No mechanism is provided
for terminating the program, so any number of mouse inputs can be processed.

```c
#include <GL/glut.h>
#include <stdio.h>

const GLint pickBuffSize = 32;

/* Set initial display-window size. */
GLsizei winWidth = 400, winHeight = 400;

void init (void)
{
    /* Set display-window color to white. */
    glClearColor (1.0, 1.0, 1.0, 1.0);
}

/* Define 3 rectangles and associated IDs. */
void rects (GLenum mode)
{
    if (mode == GL_SELECT)
        glPushName (30);   // Red rectangle.
        glColor3f (1.0, 0.0, 0.0);
        glRecti (40, 130, 150, 260);
    
    if (mode == GL_SELECT)
        glPushName (10);   // Blue rectangle.
        glColor3f (0.0, 0.0, 1.0);
        glRecti (150, 130, 260, 260);
    
    if (mode == GL_SELECT)
        glPushName (20);   // Green rectangle.
        glColor3f (0.0, 1.0, 0.0);
        glRecti (40, 40, 260, 130);
}

/* Print the contents of the pick buffer for each mouse selection. */
void processPicks (GLint nPicks, GLuint pickBuffer [ ])
```
GLint j, k;
GLuint objID, *ptr;

printf (" Number of objects picked = %d\n", nPicks);
printf ("\n");
ptr = pickBuffer;

/* Output all items in each pick record. */
for (j = 0; j < nPicks; j++) {
    objID = *ptr;
    printf (" Stack position = %d\n", objID);
    ptr++;
    printf (" Min depth = %g,\n", float (*ptr/0x7fffffff));
    ptr++;
    printf (" Max depth = %g\n", float (*ptr/0x7fffffff));
    ptr++;
    printf (" Stack IDs are: \n");
    for (k = 0; k < objID; k++) {
        printf (" %d ",*ptr);
        ptr++;
    }
    printf ("\n\n");
}

void pickRects (GLint button, GLint action, GLint xMouse, GLint yMouse)
{
    GLuint pickBuffer [pickBuffSize];
    GLint nPicks, vpArray [4];
    if (button != GLUT_LEFT_BUTTON || action != GLUT_DOWN)
        return;
    glSelectBuffer (pickBuffSize, pickBuffer); // Designate pick buffer.
    glRenderMode (GL_SELECT); // Activate picking operations.
    glInitNames ( ); // Initialize the object-ID stack.
    /* Save current viewing matrix. */
    glMatrixMode (GL_PROJECTION);
    glPushMatrix ( );
    glLoadIdentity ( );
    /* Obtain the parameters for the current viewport. Set up
     * a 5 x 5 pick window, and invert the input yMouse value
     * using the height of the viewport, which is the fourth
     * element of vpArray.
     */
    gl.GetIntegerv (GL_VIEWPORT, vpArray);
    gluPickMatrix (GLdouble (xMouse), GLdouble (vpArray [3] - yMouse),
    5.0, 5.0, vpArray);
gluOrtho2D (0.0, 300.0, 0.0, 300.0);
rects (GL_SELECT); // Process the rectangles in selection mode.

/* Restore original viewing matrix. */
glMatrixMode (GL_PROJECTION);
glPopMatrix ();
glFlush ();

/* Determine the number of picked objects and return to the normal rendering mode. */
nPicks = glRenderMode (GL_RENDER);
processPicks (nPicks, pickBuffer); // Process picked objects.
glutPostRedisplay ();
}

void displayFcn (void)
{
    glClear (GL_COLOR_BUFFER_BIT);
    rects (GL_RENDER); // Display the rectangles.
glFlush ();
}

void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    /* Reset viewport and projection parameters. */
    glViewport (0, 0, newWidth, newHeight);
    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ();
    gluOrtho2D (0.0, 300.0, 0.0, 300.0);
    glMatrixMode (GL_MODELVIEW);

    /* Reset display-window size parameters. */
    winWidth = newWidth;
    winHeight = newHeight;
}

void main (int argc, char** argv)
{
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowPosition (100, 100);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Example Pick Program");

    init ( );
    glutDisplayFunc (displayFcn);
    glutReshapeFunc (winReshapeFcn);
    glutMouseFunc (pickRects);

    glutMainLoop ( );
}
EXAMPLE 1 Sample Output from Procedure `pickrects`.

```
Number of objects picked = 2
  Stack position = 1
  Min depth = 0, Max depth = 0
  Stack IDs are:
    30

  Stack position = 3
  Min depth = 0, Max depth = 0
  Stack IDs are:
    30 10 20
```

7 OpenGL Menu Functions

In addition to the input-device routines, GLUT contains various functions for adding simple pop-up menus to programs. With these functions, we can set up and access a variety of menus and associated submenus. The GLUT menu commands are placed in procedure `main` along with the other GLUT functions.

Creating a GLUT Menu

A pop-up menu is created with the statement

```c
    glutCreateMenu (menuFcn);
```

where parameter `menuFcn` is the name of a procedure that is to be invoked when a menu entry is selected. This procedure has one argument, which is the integer value corresponding to the position of a selected option.

```c
    void menuFcn (GLint menuItemNumber)
```

The integer value passed to parameter `menuItemNumber` is then used by `menuFcn` to perform an operation. When a menu is created, it is associated with the current display window.

Once we have designated the menu function that is to be invoked when a menu item is selected, we must specify the options that are to be listed in the menu. We do this with a series of statements that list the name and position for each option. These statements have the general form

```c
    glutAddMenuEntry (charString, menuItemNumber);
```

Parameter `charString` specifies text that is to be displayed in the menu, and parameter `menuItemNumber` gives the location for that entry in the menu. For example, the following statements create a menu with two options:

```c
    glutCreateMenu (menuFcn);
    glutAddMenuEntry ("First Menu Item", 1);
    glutAddMenuEntry ("Second Menu Item", 2);
```

Next, we must specify a mouse button that is to be used to select a menu option. This is accomplished with

```c
    glutAttachMenu (button);
```
where parameter button is assigned one of the three GLUT symbolic constants referencing the left, middle, or right mouse button.

To illustrate the creation and use of a GLUT menu, the following program provides two options for displaying the interior fill of a triangle. Initially, the triangle is defined with two white vertices, one red vertex, and a fill color determined by an interpolation of the vertex colors. We use the glShadeModel function to select a polygon fill that is either a solid color or an interpolation (Gouraud rendering) of the vertex colors. A menu is created in this program that allows us to choose between these two options using the right mouse button, when the mouse cursor is inside the display window. This pop-up menu is displayed with the upper-left corner at the position of the mouse cursor. A menu option is highlighted when we move the mouse cursor over that option. The highlighted option is then selected by releasing the right button. If the option “Solid-Color Fill” is selected, the triangle is filled with the color specified for the last vertex (which is red). At the end of the menu-display procedure, fillOption, we include a glutPostRedisplay command to indicate that the triangle should be redrawn when the menu is displayed.

```c
#include <GL/glut.h>

GLsizei winWidth = 400, winHeight = 400; // Initial display-window size.
GLfloat red = 1.0, green = 1.0, blue = 1.0; // Initial triangle color: white.
GLenum fillMode = GL_SMOOTH; // Initial polygon fill: color interpolation.

void init (void)
{
  glClearColor (0.6, 0.6, 0.6, 1.0); // Set display-window color to gray.
  glMatrixMode (GL_PROJECTION);
  gluOrtho2D (0.0, 300.0, 0.0, 300.0);
}

void fillOption (GLint selectedOption)
{
  switch (selectedOption) {
  case 1: fillMode = GL_FLAT; break; // Flat surface rendering.
  case 2: fillMode = GL_SMOOTH; break; // Gouraud rendering.
  }
  glutPostRedisplay ( );
}

void displayTriangle (void)
{
  glClear (GL_COLOR_BUFFER_BIT);
  glShadeModel (fillMode); // Set fill method for triangle.
  glColor3f (red, green, blue); // Set color for first two vertices.
  glBegin (GL_TRIANGLES);
  glVertex2i (280, 20);
  glVertex2i (160, 280);
```
Creating and Managing Multiple GLUT Menus

When a menu is created, it is associated with the current display window. We can create multiple menus for a single display window, and we can create different menus for different windows. As each menu is created, it is assigned an integer identifier, starting with the value 1 for the first menu created. The integer identifier for a menu is returned by the glutCreateMenu routine, and we can record this value with a statement such as

```c
menuID = glutCreateMenu (menuFcn);
```
A newly created menu becomes the **current menu** for the current display window. To activate a menu for the current display window, we use the statement

```c
glutSetMenu (menuID);
```

This menu then becomes the current menu, which will pop up in the display window when the mouse button that has been attached to that menu is pressed.

We eliminate a menu with the command

```c
glutDestroyMenu (menuID);
```

If the designated menu is the current menu for a display window, then that window has no menu assigned as the current menu, even though other menus may exist.

The following function is used to obtain the identifier for the current menu in the current display window:

```c
currentMenuID = glutGetMenu ( );
```

A value of 0 is returned if no menus exist for this display window or if the previous current menu was eliminated with the `glutDestroyMenu` function.

### Creating GLUT Submenus

A submenu can be associated with a menu by first creating the submenu using `glutCreateMenu`, along with a list of suboptions, and then listing the submenu as an additional option in the main menu. We can add the submenu to the option list in a main menu (or other submenu) using a sequence of statements such as

```c
submenuID = glutCreateMenu (submenuFcn);
    glutAddMenuEntry ("First Submenu Item", 1);
    ...
    ...
    glutCreateMenu (menuFcn);
    glutAddMenuEntry ("First Menu Item", 1);
    ...
    ...
    glutAddSubMenu ("Submenu Option", submenuID);
```

The `glutAddSubMenu` function can also be used to add the submenu to the current menu.

In the following program, we illustrate the creation of a submenu. This program, which is a modification of the previous menu program, displays a submenu that provides three color choices (blue, green, and white) for the first two vertices of the triangle. The main menu is now listed with three options, and the third option is displayed with an arrow symbol to indicate that a pop-up submenu will be displayed when that option is highlighted. A `glutPostRedisplay` function is included at the end of both the main-menu function and the submenu function.
#include <GL/glut.h>

GLsizei winWidth = 400, winHeight = 400; // Initial display-window size.
GLfloat red = 1.0, green = 1.0, blue = 1.0; // Initial color values.
GLenum renderingMode = GL_SMOOTH; // Initial fill method.

void init (void)
{
    glClearColor (0.6, 0.6, 0.6, 1.0); // Set display-window color to gray.
    glMatrixMode (GL_PROJECTION);
    gluOrtho2D (0.0, 300.0, 0.0, 300.0);
}

void mainMenu (GLint renderingOption)
{
    switch (renderingOption) {
    case 1: renderingMode = GL_FLAT; break;
    case 2: renderingMode = GL_SMOOTH; break;
    }
    glutPostRedisplay ( );
}

/* Set color values according to the submenu option selected. */
void colorSubMenu (GLint colorOption)
{
    switch (colorOption) {
    case 1:
        red = 0.0; green = 0.0; blue = 1.0;
        break;
    case 2:
        red = 0.0; green = 1.0; blue = 0.0;
        break;
    case 3:
        red = 1.0; green = 1.0; blue = 1.0;
    }
    glutPostRedisplay ( );
}

void displayTriangle (void)
{
    glClear (GL_COLOR_BUFFER_BIT);
    glShadeModel (renderingMode); // Set fill method for triangle.
    glColor3f (red, green, blue); // Set color for first two vertices.
    glBegin (GL_TRIANGLES);
    glVertex2i (280, 20);
    glVertex2i (160, 280);
    glColor3f (1.0, 0.0, 0.0); // Set color of last vertex to red.
    glVertex2i (20, 100);
    glEnd ( );
    glFlush ( );
}
void reshapeFcn (GLint newWidth, GLint newHeight)
{
    glViewport (0, 0, newWidth, newHeight);

    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ();
    gluOrtho2D (0.0, GLfloat (newWidth), 0.0, GLfloat (newHeight));

    displayTriangle ( );
    glFlush ( );
}

void main (int argc, char **argv)
{
    GLint subMenu; // Identifier for submenu.

    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowPosition (200, 200);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Submenu Example");

    init ( );
    glutDisplayFunc (displayTriangle);

    subMenu = glutCreateMenu (colorSubMenu);
    glutAddMenuEntry ("Blue", 1);
    glutAddMenuEntry ("Green", 2);
    glutAddMenuEntry ("White", 3);

    glutCreateMenu (mainMenu); // Create main pop-up menu.
    glutAddMenuEntry ("Solid-Color Fill", 1);
    glutAddMenuEntry ("Color-Interpolation Fill", 2);
    glutAddSubMenu ("Color", subMenu);

    /* Select menu option using right mouse button. */
    glutAttachMenu (GLUT_RIGHT_BUTTON);

    glutReshapeFunc (reshapeFcn);
    glutMainLoop ( );
}

Modifying GLUT Menus

If we want to change the mouse button that is used to select a menu option, we first cancel the current button attachment and then attach the new button. A button attachment is cancelled for the current menu with

    glutDetachMenu (mouseButton);

where parameter mouseButton is assigned the GLUT constant identifying the button (left, middle, or right) that was previously attached to the menu. Once we
have detached the menu from the button, we can use `glutAttachMenu` to attach it to a different button.

Options within an existing menu can also be changed. For example, we can delete an option in the current menu with the function

```c
    glutRemoveMenuItem (itemNumber);
```

where parameter `itemNumber` is assigned the integer value of the menu option that is to be deleted.

Other GLUT routines allow us to modify the names or status of items within an existing menu. For example, we can use these routines to change the displayed name of a menu option, to change the item number of the option, or to change an option into a submenu.

---

### 8 Designing a Graphical User Interface

A common feature of modern applications software is a graphical user interface (GUI) composed of display windows, icons, menus, and other features to aid a user in applying the software to a particular problem. Specialized interactive dialogues are designed so that programming options are selected using familiar terms within a particular field, such as architectural and engineering design, drafting, business graphics, geology, economics, chemistry, or physics. Other considerations for a user interface (whether graphical or not) are the accommodation of various skill levels, consistency, error handling, and feedback.

#### The User Dialogue

For any application, the user’s model serves as the basis for the design of the dialogue by describing what the system is designed to accomplish and what operations are available. It states the type of objects that can be displayed and how the objects can be manipulated. For example, if the system is to be used as a tool for architectural design, the model describes how the package can be used to construct and display views of buildings by positioning walls, doors, windows, and other building components. A facility-layout package might include a set of furniture items along with the operations for positioning and removing different objects in a specified floor plan. A circuit-design program provides electrical or logic symbols and the positioning operations for adding or deleting elements within a layout.

All information in the user dialogue is presented in the language of the application. In an architectural design package, this means that all interactions are described only in architectural terms, without reference to particular data structures, computer-graphics terms, or other concepts that may be unfamiliar to an architect.

#### Windows and Icons

Typical GUIs provide visual representations both for the objects that are to be manipulated in an application and for the actions to be performed on the application objects.

In addition to the standard display-window operations, such as opening, closing, positioning, and resizing, other operations are needed for working with the sliders, buttons, icons, and menus. Some systems are capable of supporting multiple window managers so that different window styles can be accommodated,
each with its own window manager, which could be structured for a particular application.

Icons representing objects such as walls, doors, windows, and circuit elements are often referred to as **application icons**. The icons representing actions, such as rotate, magnify, scale, clip, or paste, are called **control icons**, or **command icons**.

**Accommodating Multiple Skill Levels**

Usually, interactive GUIs provide several methods for selecting actions. For example, an option could be specified by pointing to an icon, accessing a pull-down or pop-up menu, or by typing a keyboard command. This allows a package to accommodate users that have different skill levels.

A less experienced user may find an interface with a large, comprehensive set of operations to be difficult to use, so a smaller interface with fewer but more easily understood operations and detailed prompting may be preferable. A simplified set of menus and options is easy to learn and remember, and the user can concentrate on the application instead of on the details of the interface. Simple point-and-click operations are often easiest for an inexperienced user of an applications package. Therefore, interfaces typically provide a means for masking the complexity of a package, so that beginners can use the system without being overwhelmed with too much detail.

Experienced users, on the other hand, typically want speed. This means fewer prompts and more input from the keyboard or with multiple mouse-button clicks. Actions are selected with function keys or with simultaneous combinations of keyboard keys, because experienced users will remember these shortcuts for commonly used actions.

An interface may be designed to provide different sets of options to users with different experience levels. This may be selectable by the user through an application preference setting, or suggested by the application itself as the user gains experience with it. Similarly, help facilities can be designed on several levels so that beginners can carry on a detailed dialogue, while more experienced users can reduce or eliminate prompts and messages. Help facilities can also include one or more tutorial applications, which provide users with an introduction to the capabilities and use of the system.

**Consistency**

An important design consideration in an interface is consistency. An icon shape should always have a single meaning, rather than serving to represent different actions or objects depending on the context. Some other examples of consistency are always placing menus in the same relative positions so that a user does not have to hunt for a particular option, always using the same combination of keyboard keys for an action, and always using the same color encoding so that a color does not have different meanings in different situations.

**Minimizing Memorization**

Operations in an interface should also be structured so that they are easy to understand and to remember. Obscure, complicated, inconsistent, and abbreviated command formats lead to confusion and reduction in the effective application of the software. One key or button used for all delete operations, for example, is easier to remember than a number of different keys for different kinds of delete procedures.
Icons and window systems can also be organized to minimize memorization. Different kinds of information can be separated into different windows so that a user can identify and select items easily. Icons should be designed as easily recognizable shapes that are related to application objects and actions. To select a particular action, a user should be able to select an icon that resembles that action.

Backup and Error Handling
A mechanism for undoing a sequence of operations is another common feature of an interface, which allows a user to explore the capabilities of a system, knowing that the effects of a mistake can be corrected. Typically, systems can now undo several operations, thus allowing a user to reset the system to some specified action. For those actions that cannot be reversed, such as closing an application without saving changes, the system asks for a verification of the requested operation.

In addition, good diagnostics and error messages help a user to determine the cause of an error. Interfaces can attempt to minimize errors by anticipating certain actions that could lead to an error; and users can be warned if they are requesting ambiguous or incorrect actions, such as attempting to apply a procedure to multiple application objects.

Feedback
Responding to user actions is another important feature of an interface, particularly for an inexperienced user. As each action is entered, some response should be given. Otherwise, a user might begin to wonder what the system is doing and whether the input should be reentered.

Feedback can be given in many forms, such as highlighting an object, displaying an icon or message, and displaying a selected menu option in a different color. When the processing of a requested action is lengthy, the display of a flashing message, clock, hourglass, or other progress indicator is important. It may also be possible for the system to display partial results as they are completed, so that the final display is built up a piece at a time. The system might also allow a user to input other commands or data while one instruction is being processed.

Standard symbol designs are used for typical kinds of feedback. A cross, a frowning face, or a thumbs-down symbol is often used to indicate an error, and some kind of time symbol or a blinking “at work” sign is used to indicate that an action is being processed. This type of feedback can be very effective with a more experienced user, but the beginner may need more detailed feedback that not only clearly indicates what the system is doing but also what the user should input next.

Clarity is another important feature of feedback. A response should be easily understood, but not so overpowering that the user’s concentration is interrupted. With function keys, feedback can be given as an audible click or by lighting up the key that has been pressed. Audio feedback has the advantage that it does not use up screen space, and it does not divert the user’s attention from the work area.

A fixed message area can be used so that a user always know where to look for messages, but it may be advantageous in some cases to place feedback messages in the work area near the cursor. Feedback can also be displayed in different colors to distinguish it from other displayed objects.

Echo feedback is often useful, particularly for keyboard input, so that errors can be detected quickly. Button and dial input can be echoed in the same way. Scalar values that are selected with dials or from displayed scales are usually
echoed on the screen so that a user can check the input values for accuracy. Selection of coordinate points can be echoed with a cursor or other symbol that appears at the selected position. For more precise echoing of selected positions, the coordinate values could also be displayed on the screen.

9 Summary

Input to graphics programs can come from many different hardware devices, with more than one device providing the same general class of input data. Graphics input functions are often designed to be independent of hardware by adopting a logical classification for input devices. A device is then specified according to the type of graphics input. The six logical devices used in the ISO and ANSI standards are locator, stroke, string, valuator, choice, and pick. Locator devices input a single coordinate position. Stroke devices input a stream of coordinates. String devices input text. Valuator devices enter a scalar value. Choice devices are used for menu selections, and pick devices allow us to select scene components. Device-independent graphics packages offer a limited set of input functions that are defined in an auxiliary library.

Three modes are commonly used for input functions. Request mode places input under the control of the application program. Sample mode allows the input devices and program to operate concurrently. Event mode allows input devices to initiate data entry and control processing of data. Once we have chosen a mode for a logical device class and the particular physical device to be used to enter this class of data, input functions are used to enter data values into the program. An application program can make simultaneous use of several physical input devices operating in different modes.

Interactive picture-construction methods are commonly used in a variety of applications, including design and painting packages. These methods provide users with the capability to specify object positions, constrain objects to predefined orientations or alignments, and interactively draw or paint objects into a scene. Grids, gravity fields, and rubber-band methods are used to aid in positioning and other picture-construction operations.

Graphical user interfaces (GUIs) are now standard features of applications software. A dialogue for the software is designed from the user’s model, which describes the purpose and functions of the applications package. All elements of the dialogue are presented in the language of the application.

Window systems provide a typical interface with procedures for manipulating display windows, menus, and icons. General window systems can be designed to support multiple window managers.

Considerations in user-dialogue design are ease of use, clarity, and flexibility. Specifically, GUIs are designed to maintain consistency in user interaction and to provide for different user skill levels. In addition, interfaces are designed to minimize user memorization, to provide sufficient feedback, and to provide adequate backup and error handling capabilities.

In the Utility Toolkit, GLUT, input functions are available for interactive devices, such as a mouse, tablet, spaceball, button box, and dial box. In addition, GLUT provides a function for accepting a combination of input values from a mouse and a keyboard. Picking operations can be performed using functions from the GLU library and the basic OpenGL library. We can also display pop-up menus and submenus using a set of functions in the GLUT library. A summary of the OpenGL input and menu functions is given in Tables 1 and 2.
<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glutMouseFunc</td>
<td>Specifies a mouse callback function that is to be invoked when a mouse button is pressed.</td>
</tr>
<tr>
<td>glutMotionFunc</td>
<td>Specifies a mouse callback function that is to be invoked when the mouse cursor is moved while a button is pressed.</td>
</tr>
<tr>
<td>glutPassiveMotionFunc</td>
<td>Specifies a mouse callback function that is to be invoked when the mouse cursor is moved without pressing a button.</td>
</tr>
<tr>
<td>glutKeyboardFunc</td>
<td>Specifies a keyboard callback function that is to be invoked when a standard key is pressed.</td>
</tr>
<tr>
<td>glutSpecialFunc</td>
<td>Specifies a keyboard callback function that is to be invoked when a special-purpose key (e.g., a function key) is pressed.</td>
</tr>
<tr>
<td>glutTabletButtonFunc</td>
<td>Specifies a tablet callback function that is to be invoked when a tablet button is pressed while the mouse cursor is in a display window.</td>
</tr>
<tr>
<td>glutTabletMotionFunc</td>
<td>Specifies a tablet callback function that is to be invoked when a tablet stylus or cursor is moved while the mouse cursor is in a display window.</td>
</tr>
<tr>
<td>glutSpaceballButtonFunc</td>
<td>Specifies a spaceball callback function that is to be invoked when a spaceball button is pressed while the mouse cursor is in a display window, or using another display-window activation method.</td>
</tr>
<tr>
<td>glutSpaceballMotionFunc</td>
<td>Specifies a spaceball callback function that is to be invoked when a spaceball translational motion occurs for an activated display window.</td>
</tr>
<tr>
<td>glutSpaceballRotateFunc</td>
<td>Specifies a spaceball callback function that is to be invoked when a spaceball rotational motion occurs for an activated display window.</td>
</tr>
<tr>
<td>glutButtonBoxFunc</td>
<td>Specifies a button-box callback function that is to be invoked when a button is pressed.</td>
</tr>
<tr>
<td>glutDialsFunc</td>
<td>Specifies a dial callback function that is to be invoked when a dial is rotated.</td>
</tr>
</tbody>
</table>
**TABLE 1**

(Continued)

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>glSelectBuffer</code></td>
<td>Specifies size and name for the pick buffer.</td>
</tr>
<tr>
<td><code>glRenderMode</code></td>
<td>Activates pick operations using the argument <code>GL_SELECT</code>. This function is</td>
</tr>
<tr>
<td></td>
<td>also used to activate the normal rendering mode or the feedback mode.</td>
</tr>
<tr>
<td><code>glInitNames</code></td>
<td>Activates the object-ID name stack.</td>
</tr>
<tr>
<td><code>glPushName</code></td>
<td>Pushes an object identifier onto the ID stack.</td>
</tr>
<tr>
<td><code>glLoadName</code></td>
<td>Replaces the top identifier on the ID stack with a specified value.</td>
</tr>
<tr>
<td><code>glPopName</code></td>
<td>Eliminates the top item on the ID stack.</td>
</tr>
<tr>
<td><code>gluPickMatrix</code></td>
<td>Defines a pick window and forms a revised view volume for the picking operations.</td>
</tr>
</tbody>
</table>

**TABLE 2**

Summary of OpenGL Menu Functions

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>glutCreateMenu</code></td>
<td>Creates a pop-up menu and specifies a procedure that is to be invoked when a menu item is selected; an integer identifier is assigned to the created menu.</td>
</tr>
<tr>
<td><code>glutAddMenuEntry</code></td>
<td>Specifies an option that is to be listed in a pop-up menu.</td>
</tr>
<tr>
<td><code>glutAttachMenu</code></td>
<td>Specifies the mouse button that is to used for selecting menu options.</td>
</tr>
<tr>
<td><code>glutSetMenu</code></td>
<td>Specifies the current menu for the current display window.</td>
</tr>
<tr>
<td><code>glutDestroyMenu</code></td>
<td>Specifies an identifier for a menu that is to be eliminated.</td>
</tr>
<tr>
<td><code>glutGetMenu</code></td>
<td>Returns the identifier for the current menu attached to the current window.</td>
</tr>
<tr>
<td><code>glutAddSubMenu</code></td>
<td>Specifies a submenu that is to be included in a menu listing, where the indicated submenu has been set up using the <code>glutCreateMenu</code> routine.</td>
</tr>
<tr>
<td><code>glutDetachMenu</code></td>
<td>Cancels a specified mouse-button attachment for the current menu.</td>
</tr>
<tr>
<td><code>glutRemoveMenuItem</code></td>
<td>Deletes a specified option in the current menu.</td>
</tr>
</tbody>
</table>
REFERENCES
The evolution of the concept of logical (or virtual) input
devices is discussed in Wallace (1976) and in Rosenthal,
et al. (1982). Implementations for various input proce-
dures are given in Glassner (1990), Arvo (1991), Kirk
programming examples using mouse and keyboard input
can be found in Woo, et al. (1999). A complete
listing of the functions in the OpenGL basic library and
the GLU library is given in Shreiner (2000). The GLUT
input and menu functions are listed in detail in Kilgard
(1996).

Guidelines for user-interface design are presented in
and Cunningham (1989), Digital (1989), OSF/MOTIF
(1989), and Laurel (1990).

EXERCISES
1 Design an algorithm that allows objects to be
positioned on the screen using a locator device.
An object menu of geometric shapes is to be pre-
ented to a user who is to select an object and
a placement position. The program should allow
any number of objects to be positioned until a
“terminate” signal is given.
2 Extend the algorithm of the previous exercise so
that selected objects can be scaled and rotated
before positioning. The transformation choices
and transformation parameters are to be pre-
nented to the user as menu options.
3 Set up a procedure for interactively sketching pic-
tures using a stroke device.
4 Discuss the methods that could be employed in
a pattern-recognition procedure to match input
characters against a stored library of shapes.
5 Write a routine that displays a linear scale and a
slider on the screen and allows numeric values to
be selected by positioning the slide along the scale
line. The selected numeric value is to be echoed
in a box displayed near the linear scale.
6 Write a program that makes use of the slider
developed in the previous exercise to allow the
user to scale an object displayed in a display
window between some minimum and maximum
value.
7 Write a routine that displays a circular scale and a
pointer or a slider that can be moved around the
circle to select angles (in degrees). The angular
value selected is to be echoed in a box displayed
near the circular scale.
8 Write a program that makes use of the circular
slider developed in the previous exercise to allow
the user to rotate an object around its center.

9 Write a drawing program that allows users to cre-
ate a picture as a set of straight-line segments
drawn between specified endpoints. The coordi-
nates of the individual line segments are to be
selected with a locator device.
10 Write a drawing package that allows pictures to
be created with straight-line segments drawn
between specified endpoints. Set up a gravity
field around each line in a picture, as an aid in
connecting new lines to existing lines.
11 Modify the drawing package in the previous exer-
cise so that lines can be constrained horizontally
or vertically.
12 Develop a drawing package that can display
an optional grid pattern so that selected screen
positions are rounded to grid intersections. The
package is to provide line-drawing capabilities,
with line endpoints selected using a locator
device.
13 Write a routine that allows a designer to create a
picture by sketching straight lines using a rubber-
band method.
14 Design a drawing package that allows straight
lines, rectangles, and circles to be constructed
using rubber-band methods.
15 Write a procedure that allows a user to pick com-
ponents of a two-dimensional scene. The coordi-
nate extents for each object are to be stored and
used to identify the picked object, given an input
cursor position.
16 Develop a procedure that allows a user to design a
picture from a menu of displayed basic shapes by
dragging each selected shape into position with a
pick device.
17 Design an implementation of the input functions
for request mode.
18 Design an implementation of the sample mode
input functions.
19 Design an implementation of the input functions
for event mode.
20 Design a procedure for implementing input func-
tions for request, sample, and event mode.
21 Expand the OpenGL point-plotting program in
Section 6 to include a menu that allows a user
to select point size and point color.
22 Expand the OpenGL polyline program in Section
6 to include a menu that allows a user to choose
the line attributes: size, color, and width.
23 Modify the program in the preceding exercise
to allow a texture pattern to be chosen for the
polyline.
24 Write an interactive OpenGL program to display a circle of radius 150 that can be at any input position within a display window. The input position is to be the center of the circle. Include a menu of color options for displaying the circle in a solid color.

25 Modify the program in the preceding exercise so that the input position is rejected if the entire circle cannot be displayed within the display window.

26 Modify the program in the preceding exercise to include a menu of texture options for the circle. Include a solid color texture, a gradient texture, and a minimum of two other texture patterns.

27 Set up an interactive OpenGL program for displaying an input character string at any position within a display window. The input position is the starting position for the text.

28 Write an interactive OpenGL program for positioning a single two-dimensional object at any position within a display window. The object is to be selected from a menu of basic shapes, including (minimally) a square, circle, and triangle.

29 Modify the program in the preceding exercise to allow any arrangement of the two-dimensional objects to be displayed, with each object selected from the menu until a quit option is chosen from the menu.

30 Modify the program in the preceding exercise to allow objects to be scaled or rotated. Geometric transformation operations are to be listed in a menu.

31 Write an interactive OpenGL program for positioning a single three-dimensional object within a display window. The object is to be selected from a menu list of GLUT wire-frame solids, such as a sphere, solid, or cylinder, and it is to be centered on an input position.

32 Modify the program in the preceding exercise to allow the objects to be displayed in either a wireframe or solid form. For solid-object displays, include a point light source at the viewing position, and use default parameters for the illumination and surface shading.

33 Write a program to implement the OpenGL picking operations for a three-dimensional scene containing several objects. For each pick selection, create a small pick window and remove the most distant object within that pick window from the scene, replacing it with a new object at a random location in the scene.

34 Write an interactive OpenGL program to display a two-dimensional cubic Bézier curve. The four control-point positions are to be selected with mouse input.

35 Modify the program in the preceding exercise to display a Bézier curve with a selected degree of three, four, or five.

36 Write an interactive OpenGL program to display a two-dimensional cubic B-spline. The spline parameters are to be given as input, and the control points are to be selected with a mouse.

37 Write an interactive OpenGL program to display a cubic Bézier surface patch. The x and y coordinates for the control points can be selected with a mouse, and the z coordinate can be given as a height above a ground plane.

38 Select some graphics application with which you are familiar and set up a user model that will serve as the basis for the design of a user interface for graphics applications in that area.

39 List possible help facilities that can be provided in a user interface and discuss which types of help would be appropriate for different levels of users.

40 Summarize the methods for handling backup and errors. Which methods are suitable for a beginner? Which methods are better for an experienced user?

41 List the possible formats for presenting menus to a user, and explain under what circumstances each might be appropriate.

42 Discuss alternatives for feedback in terms of the various levels of users.

43 List the functions that must be performed by a window manager in handling screen layouts with multiple, overlapping windows.

44 Set up a design for a window-manager package.

45 Design a user interface for a painting program.

46 Design a user interface for a two-level hierarchical modeling package.

**In More Depth**

1 In this chapter’s exercises, you will add the necessary input devices that will make your program interactive. Think about the types of tasks that the user should be able to perform in your application and decide how best to implement the interaction necessary to achieve that behavior. What types of responses will occur in the graphical view of the application in response to each type of input? There is usually more than one way to obtain a certain type of input, and more than one way to present input options to the user. Discuss these alternatives and write an input specification employing those that you think would be most intuitive for the user. Add menus and other graphical user interface (GUI) components to make your application more user-friendly.
2 Implement the input specification you designed in the previous exercise using the OpenGL interactive input-device functions defined in the GLUT libraries. Use mouse and keyboard input to allow the user to interact with the application, adding GUI components where necessary or appropriate. Write routines to update the current scene of your application in response to each type of input. Extensively test your application as a user and experiment with modifications to the input specification to make controlling the application as intuitive as possible.
Interactive Input Methods and Graphical User Interfaces

Color Plates

Color Plate 23
A screen layout showing one type of interface for an artist's painting package. (Courtesy of Thomson Digital Image.)

Color Plate 24
Using a head-tracking stereo display, called the BOOM (Fake Space Labs, Inc.), and a Dataglove (VPL, Inc.), a researcher interactively manipulates exploratory probes in the unsteady flow around a Harrier jet airplane. Software developed by Steve Bryson; data from Harrier. (Courtesy of Sam Usetton, NASA Ames Research Center.)
Illumination models in computer graphics are often approximations of the physical laws that describe surface-lighting effects. To reduce computations, most packages use empirical models based on simplified photometric calculations. Surface rendering is performed through calculating the interaction of an object’s surface with the light striking it. This type of illumination model is known as local illumination, and considers only the properties of that object and the light that directly strikes it.

To produce more realistic lighting effects, we must also consider the contribution of light that is reflected from other objects onto the surface of the object being shaded. This type of illumination, called global illumination, can be more accurate, but that accuracy comes at the expense of additional computation.

Some global illumination methods, such as ray-tracing, attempt to determine surface shading by following light rays from the eye-point back into the scene through the pixels of the image plane.
More accurate models, such as the radiosity algorithm, compute light intensities by considering the propagation of radiant energy or by following the movement of photons from between the light sources and the various surfaces in a scene. Also, it is possible to combine these techniques with other shading methods such as texture mapping to simulate the effect of a surrounding environment on the surface of an object.

1 Ray-Tracing Methods

Ray casting is used in constructive solid geometry for locating surface intersections along a ray from a pixel position. Ray casting is also a means for identifying visible surfaces in a scene. Ray tracing is the generalization of the basic ray-casting procedure. Instead of merely looking for the visible surface from each pixel position, we continue to bounce the pixel ray around in the scene, as illustrated in Figure 1, to collect the various intensity contributions. This provides a simple and powerful rendering technique for obtaining global reflection and transmission effects. In addition, the basic ray-tracing algorithm detects visible surfaces, identifies shadow areas, provides for the rendering of transparency effects, generates perspective-projection views, and accommodates illumination effects from multiple light sources. Many extensions to the basic algorithm have been developed to produce photo-realistic displays. Ray-traced pictures of scenes can be highly realistic, particularly when the scene contains shiny objects, but ray-tracing algorithms involve considerable computation time. An example of the global reflection and transmission effects possible with ray tracing is demonstrated in Color Plate 25.

Basic Ray-Tracing Algorithm

The coordinate system for a ray-tracing algorithm is typically set up as shown in Figure 2, with the projection reference point on the z axis and the pixel positions on the xy plane. We then describe the geometry of a scene in this coordinate system and generate the pixel rays. For a perspective-projection view of the scene, each ray

---

**Figure 1**

Multiple reflection and transmission paths for a ray from the projection reference point through a pixel position and on into a scene containing several objects.
originates at the projection reference point (center of projection), passes through a pixel center, and continues into the scene to form the various branches of the ray along reflection and transmission paths. Contributions to the pixel intensity are then accumulated at the intersected surfaces. This rendering approach is based on the principles of geometric optics. Light rays from the surfaces in a scene emanate in all directions, and some pass through the pixel positions on the projection plane. Because there are an infinite number of ray emanations, we determine the intensity contributions for a particular pixel by tracing a light path backward from the pixel position into the scene. In the basic ray-tracing algorithm, one reverse light ray is generated for each pixel, which is approximately equivalent to viewing the scene through a pinhole camera.

As each pixel ray is generated, the list of surfaces in the scene is processed to determine whether there are any ray–surface intersections. If the ray does intersect a surface, we calculate the distance from the pixel to the surface intersection point. After all surfaces have been tested for a ray intersection, the smallest calculated intersection distance identifies the visible surface for that pixel. We then reflect the ray off the visible surface along a specular-reflection path (where the angle of reflection equals the angle of incidence). For a transparent surface, we also send a ray through the surface in the refraction direction. The reflection and refraction rays are referred to as secondary rays.

We then repeat the ray-processing procedures for the secondary rays. Surfaces are tested for intersections, and the nearest intersected surface, if any, along a secondary ray path is used to recursively produce the next generation of reflection and refraction paths. As the rays from a pixel ricochet through the scene, each successively intersected surface is added to a binary ray-tracing tree, as shown in Figure 3. We use left branches in the tree to represent reflection paths and right branches to represent transmission paths. The maximum depth of the ray-tracing trees can be set as a user option, or can be determined by the amount of storage available. We terminate a path in the binary tree for a pixel if any of the following conditions is satisfied:

- The ray intersects no surfaces.
- The ray intersects a light source that is not a reflecting surface.
- The tree has been generated to its maximum allowable depth.

At each surface intersection, we invoke the basic illumination model to determine the surface intensity contribution. This intensity value is stored at the surface-node position in the pixel tree. A ray that intersects a nonreflecting light source can be assigned the intensity of the source, although light sources in the
basic ray-tracing algorithm are usually point sources at positions beyond the coordinate limits of the scene. Figure 4 shows a surface intersected by a ray and the unit vectors used for the reflected light intensity calculations. Unit vector $u$ is in the direction of the ray path, $N$ is the unit surface normal, $R$ is the unit reflection vector, $L$ is the unit vector indicating the direction to a point light source, and $H$ is the unit vector halfway between $L$ and $V$. For the ray-tracing calculations, the viewing direction is $V = -u$. The path along the direction of $L$ is referred to as the shadow ray. If any object intersects the shadow ray between the surface and the point light source, the surface position is in shadow with respect to that source. Ambient light at the surface is calculated as $k_a I_a$; diffuse reflection due to the source is proportional to $k_d (N \cdot L)$; and the specular reflection component is proportional to $k_s (H \cdot N)^n$. The specular-reflection direction for the secondary ray path $R$ depends upon the surface normal and the incoming ray direction, as follows:

$$R = u - (2u \cdot N)N$$

For a transparent surface, we also need to obtain intensity contributions from light transmitted (refracted) through the material. We can locate the source of this contribution by tracing a secondary ray along the transmission direction $T$, as shown in Figure 5. The unit transmission vector $T$ can be obtained from vectors $u$ and $N$ as

$$T = \frac{n_r}{n_t} u - \left( \frac{n_r}{n_t} \cos \theta_t - \frac{n_t}{n_r} \cos \theta_i \right) N$$
Parameters $\eta_i$ and $\eta_r$ are the indices of refraction in the incident material and the refracting material, respectively. Angle of refraction $\theta_r$ can be calculated from Snell’s law:

$$\cos \theta_r = \sqrt{1 - \left(\frac{\eta_i}{\eta_r}\right)^2 \left(1 - \cos^2 \theta_i\right)}$$

(3)

After the binary tree has been completed for a pixel, the intensity contributions are accumulated, starting at the bottom (terminal nodes) of the tree. The surface intensity from each node in the tree is attenuated by the distance from the parent surface (the next node up the tree) and added to the intensity of the parent surface. The intensity assigned to the pixel is the sum of the attenuated intensities at the root node of the ray tree. If the primary ray for a pixel does not intersect an object in the scene, the ray-tracing tree is empty and the pixel is assigned the background intensity.

Ray-tracing is a highly view-dependent method—that is, because the rays that are traced into the scene are emitted from the projection reference point, if the viewing position changes, the rays must be retraced through the scene. Similarly, if the positions of any objects in the scene change, all or some of the rays must be retraced because of the resulting changes in the reflection pattern of rays. These can be significant issues in applications where near-real-time results are desired unless there is significant hardware support for the required computation.

**Ray–Surface Intersection Calculations**

A ray can be described with an initial position $P_0$ and a unit direction vector $u$, as illustrated in Figure 6. The coordinates for any point $P$ along the ray at a distance $s$ from $P_0$ are then computed from the following ray equation:

$$P = P_0 + su$$

(4)
Initially, vector $P_0$ can be set to the position $P_{pix}$ of the pixel on the projection plane, or it could be chosen to be the projection reference point. Unit vector $u$ is initially obtained from the position of the pixel through which the ray passes and the projection reference point:

$$u = \frac{P_{pix} - P_{prp}}{|P_{pix} - P_{prp}|} \quad (5)$$

Although it is not necessary for $u$ to be a unit vector, this will simplify some calculations.

To locate the ray-intersection position on a surface, we use the surface equation to solve for position $P$, as represented by Equation 4. This gives us a value for parameter $s$, which is the distance from $P_0$ to the surface intersection point along the ray path.

At each intersected surface, vectors $P_0$ and $u$ are updated for the secondary rays at the ray-surface intersection point. For the secondary rays, the reflection direction for $u$ is $R$ and the transmission direction is $T$. When a secondary ray-surface intersection is detected, we simultaneously solve the ray equation and the surface equation to obtain the intersection coordinates. We then update the binary tree and generate the next set of reflection and refraction rays.

Efficient ray-surface intersection algorithms have been devised for most commonly occurring shapes, including various spline surfaces. The general procedure is to combine the ray equation with the equations describing a surface and solve for parameter $s$. In many cases, numerical root-finding methods and incremental calculations are used to locate intersection points over a surface. For complex objects, it is often convenient to transform the ray equation into the local coordinate system in which an object is defined. And intersection calculations for a complex object can be simplified in many cases by transforming the object into a more congenial shape. For example, we can ray-trace an ellipsoid by transforming the ray and surface equations into a sphere-intersection problem.

**Ray–Sphere Intersections**

The simplest objects to ray-trace are spheres. If we have a sphere of radius $r$ and center position $P_c$ (Figure 7), then any point $P$ on the surface satisfies the sphere equation:

$$|P - P_c|^2 - r^2 = 0 \quad (6)$$

Substituting Equation 4 for $P$ in the preceding equation, we have

$$|P_0 + su - P_c|^2 - r^2 = 0 \quad (7)$$

![Figure 7](image)

*Figure 7*  
A ray intersecting a sphere with radius $r$ and center position $P_c$. 
If we represent $P_c - P_0$ as $\Delta P$ and expand the dot product, we obtain the quadratic equation

$$s^2 - 2(u \cdot \Delta P)s + (|\Delta P|^2 - r^2) = 0$$

whose solution is

$$s = u \cdot \Delta P \pm \sqrt{(u \cdot \Delta P)^2 - |\Delta P|^2 + r^2}$$

If the discriminant is negative, either the ray does not intersect the sphere or the sphere is behind $P_0$. In either case, we can eliminate the sphere from further consideration, because we assume that the scene is in front of the projection plane. When the discriminant is not negative, the surface intersection coordinates are obtained from Equation 4 using the smaller of the two values from Equation 9. Color Plate 26 shows a ray-traced scene containing a snowflake pattern formed with shiny spheres, which illustrates the global surface reflections possible with ray-tracing.

Some optimizations are possible in the ray–sphere intersection calculations to reduce processing time. In addition, Equation 9 is susceptible to round-off errors when a small sphere far from the initial ray position is processed. That is, if

$$r^2 \ll |\Delta P|^2$$

we could lose the $r^2$ term in the precision error of $|\Delta P|^2$. We can avoid this in most cases by rearranging the calculation for distance $s$ as

$$s = u \cdot \Delta P \pm \sqrt{r^2 - |\Delta P - (u \cdot \Delta P)u|^2}$$

**Ray–Polyhedron Intersections**

Intersection calculations for polyhedra are more complicated than the sphere-intersection procedures. Therefore, it is often more efficient to process a polyhedron by performing an initial intersection test on a bounding volume. For example, Figure 8 shows a polyhedron inside a sphere. If a ray does not intersect the bounding sphere, we eliminate the polyhedron from further testing. Otherwise, we next identify the front faces of the polyhedron as those polygons that satisfy the inequality

$$u \cdot N < 0$$

where $N$ is the surface normal for the polygon. For each face of the polyhedron that satisfies condition 11, we solve the plane equation

$$N \cdot P = -D$$

for surface position $P$ that also satisfies Equation 4. Here, $N = (A, B, C)$ and $D$ is the fourth plane parameter. Position $P$ is both on the plane and on the ray path if

$$N \cdot (P_0 + su) = -D$$

**FIGURE 8**
A polyhedron enclosed by a bounding sphere.
and the distance from the initial ray position to the plane is

\[ s = \frac{D + \mathbf{N} \cdot \mathbf{P}_0}{\mathbf{N} \cdot \mathbf{u}} \]  

(14)

This gives us a position on the infinite plane that contains the polygon face, but this position may not be inside the polygon boundaries (Figure 9). So we need to perform an inside-outside test to determine whether the ray intersected this face of the polyhedron. We perform this test for each face satisfying inequality 11. The smallest distance \( s \) to an intersected polygon identifies the intersection position on the polyhedron surface. If no intersection positions from Equation 14 are inside points, the ray does not intersect the polyhedron.

**Reducing Object-Intersection Calculations**

Ray–surface intersection calculations can account for as much as 95 percent of the processing time in a ray tracer. For a scene with many objects, most of the processing time for each ray is spent checking objects that are not visible along the ray path. Therefore, several methods have been developed for reducing the processing time spent on these intersection calculations.

One method for reducing the intersection calculations is to enclose groups of adjacent objects within a bounding volume, such as a sphere or a box (Figure 10). We can then test for ray intersections with the bounding volume. If the ray does not intersect the surface of the bounding object, we eliminate the enclosed surfaces from further intersection tests. This approach can be extended to include a hierarchy of bounding volumes. That is, we enclose several bounding volumes within a larger volume and carry out the intersection tests hierarchically. First, we test the outer bounding volume; then, if necessary, we test the smaller inner bounding volumes; and so on.

**Space-Subdivision Methods**

Another way to reduce intersection calculations is to use **space-subdivision** procedures. We can enclose an entire scene within a cube, then we successively sub-divide the cube until each subregion (cell) contains no more than a preset maximum number of surfaces. For example, we could require that each cell contain no more than one surface. If parallel and vector processing capabilities are available, the maximum number of surfaces per cell can be determined by the size of the vector registers and the number of processors. Space subdivisions of the
cube can be stored in an octree or in a binary-partition tree. In addition, we can perform a uniform subdivision by dividing the cube into eight equal size octants at each step, or we can perform an adaptive subdivision by subdividing only those regions of the cube that contain objects.

We then trace rays through the individual cells of the cube, performing intersection tests only within those cells containing surfaces. The first surface intersected is the visible surface for that ray. There is a trade-off, however, between the cell size and the number of surfaces per cell. As we reduce the maximum number of allowable surfaces per cell, we reduce the amount of processing needed for the surface-intersection tests, but this also reduces cell size so that more calculations are needed to determine the ray path through the cells.

Figure 11 illustrates the intersection of a pixel ray with the front face of a cube surrounding a scene. The intersection position on the front face of the cube identifies the initial cell that is to be traversed by this ray. We then process the ray through the cells of the cube by determining the coordinates for the entry and exit positions (Figure 12). At each nonempty cell, we test for surface intersections. This processing continues until the ray intersects an object surface or exits the bounding cube.

Given a unit ray direction vector $u$ and a ray entry position $P_{\text{in}}$ for a cell, we identify the potential exit faces of a cell as those that satisfy the inequality

$$u \cdot N_k > 0$$

where $N_k$ represents the unit surface normal vector for face $k$ of the cell. If the unit normal vectors for the cell faces in Figure 12 are aligned with the Cartesian-coordinate axes, then

$$N_k = \begin{cases} 
(\pm 1, 0, 0) \\
(0, \pm 1, 0) \\
(0, 0, \pm 1)
\end{cases}$$

and we can determine the three candidate exit planes merely by checking the sign of each component of $u$. The exit position on each candidate plane is obtained from the ray equation as follows:

$$P_{\text{out},k} = P_{\text{in}} + s_k u$$

where $s_k$ is the distance along the ray from $P_{\text{in}}$ to $P_{\text{out},k}$. Substituting the ray equation into the plane equation for each cell face, we have

$$N_k \cdot P_{\text{out},k} = -D_k$$
and the ray distance to each candidate exit face is computed as

$$s_k = \frac{-D_k - N_k \cdot P_{in}}{N_k \cdot u}$$  \hspace{1cm} (19)

The smallest value computed for $s_k$ identifies the exit face for the cell. With the cell faces aligned parallel to the Cartesian-coordinate planes, normal vectors $N_k$ are the unit axis vectors, and we can simplify the calculations in Equation 19. For example, if a candidate exit plane has the normal vector $(1, 0, 0)$, then for that plane, we have

$$s_k = \frac{x_k - x_0}{u_x}$$  \hspace{1cm} (20)

where $u = (u_x, u_y, u_z)$, $x_k = -D_k$ is the coordinate position of the candidate exit plane, and $x_0$ is the coordinate position of the cell entry face.

Various modifications can be made to the cell traversal procedures to speed up the processing. One possibility is to take a trial exit plane $k$ as the one perpendicular to the direction of the largest component of $u$. This trial exit plane is then divided into sectors, as shown in the example of Figure 13. The sector on the trial plane containing $P_{out,k}$ determines the true exit plane. For example, if the intersection point $P_{out,k}$ is in sector 0 for the example plane of Figure 13, the trial plane is the true exit plane, and we are done. If the intersection point is in sector 1, the true exit plane is the top plane, and we need simply to calculate the exit point on the top boundary of the cell. Similarly, sector 3 identifies the bottom plane as the true exit plane; and sectors 4 and 2 identify the true exit plane as either the left or right cell plane, respectively. When the trial exit point falls in sector 5, 6, 7, or 8, we must carry out two additional intersection calculations to identify the true exit plane. Implementation of these methods on parallel vector machines provides further improvements in performance.

A light-buffer technique, which is a form of spatial partitioning, was used to render the scene in Color Plate 27. Here, a cube is centered on each point light source, and each side of the cube is partitioned using a grid of squares. A sorted list of objects that are visible to the light through each square is then maintained by the ray tracer to speed up processing of shadow rays. As a means for determining surface illumination effects, a square for each shadow ray is computed and the shadow ray is then processed against the list of objects for that square.

Intersection tests in ray-tracing programs can also be reduced with directional subdivision procedures, by considering sectors that contain a bundle of rays. Within each sector, we can sort surfaces in depth order, as in Figure 14. Each ray then needs to test only the objects within the sector that contains that ray.

**Simulating Camera Focusing Effects**

To model camera effects in a scene, we specify the focal length and other parameters for a convex lens (or camera aperture) that is to be positioned in front of the projection plane. Lens parameters are then set so that some objects in a scene can be in focus while other objects are out of focus. The lens focal length is the distance from the center of the lens to the focal point $F$, which is the convergence position for a set of parallel rays passing through the lens, as illustrated in Figure 15. A typical value for the focal length of a 35 mm camera is $f = 50$ mm. Camera apertures are usually described with a parameter $n$, called the f-number or f-stop, which is the ratio of the focal length to the aperture diameter:

$$n = \frac{f}{2r}$$  \hspace{1cm} (21)
Therefore, we could use either the radius $r$ or the $f$-number $n$, along with the focal length $f$, to specify the camera parameters. For a more accurate focusing model we could use the film size (width and height) and focal length to simulate the camera effects.

Ray-tracing algorithms typically determine focusing effects using the thin-lens equation from geometric optics:

$$\frac{1}{d} + \frac{1}{d_i} = \frac{1}{f} \tag{22}$$

Parameter $d$ is the distance from the lens center to an object position, and $d_i$ is the distance from the lens center to the image plane where that object is in focus. The object point and its image are on opposite sides of the lens along a line through the lens center, and $d > f$ (Figure 16). Therefore, to focus on a particular object at a distance $d$ from the lens, we position the pixel plane at a distance $d_i$ behind the lens.

For a scene position at some distance $d' \neq d$, the projected point will be out of focus on the image plane. If $d' > d$, the point is in focus at a position in front of the image plane; and if $d' < d$, the point is in focus at a position in back of the image plane. The projection of a point at position $d'$ on the image plane is approximately
Global Illumination

Figure 16
Thin-lens parameters. An object at a distance \(d\) from the lens is in focus on the image plane at a distance \(d_i\) from the lens.

A small circle, called the circle of confusion, and the diameter of this circle can be computed as

\[
2r_c = \frac{|d' - d| f}{nf}
\]  

(23)

We can choose the camera parameters to minimize the size of the circle of confusion for a range of distances, called the depth of field for the camera. In addition, multiple rays are traced for each pixel to sample positions throughout the lens area, and we discuss these distributed-ray tracing methods in a later section of this chapter.

Antialiased Ray Tracing

Two basic antialiasing techniques employed in ray-tracing algorithms are supersampling and adaptive sampling. In supersampling and adaptive sampling, the pixel is treated as a finite square area instead of a single point. Supersampling typically uses multiple, evenly spaced rays (samples) over each pixel area. Adaptive sampling uses unevenly spaced rays in some regions of the pixel area. For example, more rays can be used near object edges to obtain a better estimate of the pixel intensities. (Another method for sampling is to distribute the rays randomly over the pixel area. We discuss this approach in the next section.) When multiple rays per pixel are used, the intensities of the pixel rays are averaged to produce the overall pixel intensity.

Figure 17 illustrates a simple supersampling procedure. Here, one ray is generated through each corner of the pixel. If the intensities computed for the four rays are not approximately equal, or if some small object lies between the four rays, we divide the pixel area into subpixels and repeat the process. For example, the pixel in Figure 18 is divided into nine subpixels using 16 rays, one at each subpixel corner. Adaptive sampling is then used to subdivide further those subpixels that subtend a small object or do not have nearly equal-intensity rays. This subdivision process can be continued until the subpixel rays have approximately equal intensities, or until an upper bound, say 256, has been reached for the maximum number of rays per pixel.
Instead of passing rays through pixel corners, we can generate rays through subpixel centers, as in Figure 19. With this approach, we can weight the rays according to one of the sampling schemes you already know.

Another method for antialiasing displayed scenes is to treat a pixel ray as a cone, as shown in Figure 20. Only one ray is generated per pixel, but the ray now has a finite cross-section. To determine the percent of pixel-area coverage with objects, we calculate the intersection of the pixel cone with the object surface. For a sphere, this requires finding the intersection of two circles. For a polyhedron, we must find the intersection of a circle with a polygon.

Distributed Ray Tracing

Distributed ray tracing (also referred to as distribution ray tracing) is a stochastic sampling method that randomly distributes rays according to the various parameters in an illumination model. Illumination parameters include pixel area, reflection and refraction directions, camera lens area, and time. Aliasing effects are thus replaced with low-level noise, which improves picture quality and allows more accurate modeling of surface gloss and translucency, finite camera apertures, finite light sources, and motion-blur displays of moving objects. Distributed ray
tracing essentially provides a Monte Carlo evaluation of the multiple integrals that occur in an accurate physical description of surface lighting.

Pixel sampling is accomplished by randomly distributing a number of rays over the pixel area. Choosing ray positions completely at random, however, can result in a clustering of rays in a small region of the pixel area, and leaving large parts of the pixel unsampled. A better approximation of the light distribution over a pixel area is obtained by using a technique called jittering on a regular subpixel grid. This is usually done by initially dividing the pixel area (a unit square) into the 16 subareas shown in Figure 21 and generating a random jitter position in each subarea. The random ray positions are obtained by jittering the center coordinates of each subarea by small amounts, $\delta_x$ and $\delta_y$, where $-0.5 < \delta_x, \delta_y < 0.5$. We then choose the jitter position as $(x + \delta_x, y + \delta_y)$, where $(x, y)$ is the center position of the pixel.

Integer codes 1 through 16 are assigned randomly to each of the 16 rays, and a table lookup is used to obtain values for the other parameters, such as reflection angle and time. Each subpixel ray is processed through the scene to determine the intensity contribution for that ray. The 16 ray intensities are then averaged to produce the overall pixel intensity. If the subpixel intensities vary too much, we can subdivide the pixel further.

To model camera-lens effects, we process pixel rays through a lens that is positioned in front of the pixel plane. As we noted earlier, a camera is simulated using a focal length and other parameters so that selected objects will be in focus. Then we distribute subpixel rays over the aperture area. Assuming that we have 16 rays per pixel, we can subdivide the aperture area into 16 zones. Each subpixel is then assigned a center position in one of the zones, and the following procedure can be used to determine the distribution sampling for the pixel. A jittered position is calculated from each zone center, and a ray is projected into the scene from this jittered zone position through the focal point of the lens. We locate the focal point for a ray along the line from the center of the subpixel through the lens center, shown as point $F$ in Figure 22. With the pixel plane at a distance $d_i$ from the lens (Figure 16), positions along the ray near the object plane (the focusing plane), at a distance $d_i$ in front of the lens, are in focus. Other positions along the ray are blurred. To improve the display for out-of-focus objects, we increase the number of subpixel rays.

Reflection and transmission paths are also distributed throughout a spatial region. To simulate surface gloss, rays reflected from a surface position are
distributing subpixel rays about the reflection direction \( \mathbf{R} \) and the transmission direction \( \mathbf{T} \). The maximum spread about \( \mathbf{R} \) is divided into 16 angular zones, and each ray is reflected in a jittered position from the zone center corresponding to its integer code. We can use the Phong model, \( \cos^n \phi \), to determine the maximum distribution for the reflection angles. If the material is transparent, refracted rays can be distributed about the transmission direction \( \mathbf{T} \) in a similar manner to model translucency.

Extended light sources are handled by distributing a number of shadow rays over the area of the light source, as demonstrated in Figure 24. The light source is divided into zones, and shadow rays are assigned jitter directions to the various zones. In addition, zones can be weighted according to the intensity of the light source within that zone and the size of the projected area of the zone onto the object surface. More shadow rays are then sent to zones with heavier weights. If some shadow rays intersect opaque objects between the surface and the light source, a penumbra (partly illuminated region) is generated at that surface point. If all shadow rays are blocked, however, the surface point is within an umbra region (completely dark) for that light source. Figure 25 illustrates the regions for the umbra and penumbra on a surface partially shielded from a light source.

We create motion blur by distributing rays over time. A total frame time and the frame-time subdivisions are determined according to the motion dynamics required for the scene. Time intervals are labeled with integer codes, and each ray is assigned to a jittered time within the interval corresponding to the ray code. Objects are then moved to their positions at that time, and the ray is traced through the scene. Additional rays are used for highly blurred objects. To reduce calculations, we can use bounding boxes or spheres for initial ray-intersection tests. That is, we move the bounding object according to the motion requirements and test for intersection. If the ray does not intersect the bounding object, we need not process the individual surfaces within the bounding volume.

Color Plate 28 illustrates focusing, refraction, and antialiasing effects with distributed ray tracing.
Radiosity Lighting Model

Although the basic illumination model produces reasonable results for many applications, there are a variety of lighting effects that are not accurately described by the simple approximations in this model. We can model lighting effects more precisely by considering the physical laws governing the radiant-energy transfers within an illuminated scene. This method for computing pixel color values is generally referred to as the **radiosity model**.

Radiant-Energy Terms

In the quantum model of light, the energy of the radiation is carried by the individual photons. For monochromatic light radiation, the energy of each photon is calculated as

\[ E_{\text{photon}, f} = hf \]  

(24)

where the frequency \( f \), measured in hertz (cycles per second), characterizes the color of the light. A blue light has a high frequency within the visible band of the electromagnetic spectrum, and a red light has a low frequency. The frequency also gives the oscillation rate for the amplitude of the electric and magnetic components of the radiation. Parameter \( h \) is Planck's constant, which has the value \( 6.6262 \times 10^{-34} \) joules \( \cdot \) sec, independent of the light frequency.

The total energy for monochromatic light radiation is

\[ E_f = \sum_{\text{all photons}} hf \]  

(25)

The radiant energy at a particular light frequency is also referred to as a **spectral radiance**. However, any actual light radiations, even those from a “monochromatic” source, contain a range of frequencies. Therefore, the total radiant energy is the sum over all photons of all frequencies:

\[ E = \sum_{f} \sum_{\text{all photons}} hf \]  

(26)

The amount of radiant energy transmitted per unit of time is called the **radiant flux** \( \Phi \):

\[ \Phi = \frac{dE}{dt} \]  

(27)

Radiant flux is also referred to as **radiant power**, and it is measured in watts (joules per second).

To obtain the lighting effects for surfaces in a scene, we calculate the radiant flux per unit area that is leaving a surface. This quantity is called the **radiosity** \( B \), or **radiant exitance**:

\[ B = \frac{d\Phi}{dA} \]  

(28)

which is measured in units of watts per meter\(^2\). The **intensity** \( I \) is often taken to be a measure of the radiant flux in a particular direction per unit solid angle per unit projected area, with units of watts/(meter\(^2\) \( \cdot \) steradians). Sometimes, however, intensity is defined simply as the radiant flux in a particular direction.

Depending on the interpretation of the term **intensity**, the **radiance** can be defined as the intensity per unit projected area. Alternatively, we can obtain radiance from the radiant flux or the radiosity per unit solid angle.
The Basic Radiosity Model

To describe diffuse reflections from a surface accurately, the radiosity model computes radiant-energy interactions between all the surfaces in a scene. Because the resulting set of equations can be extremely difficult to solve, the basic radiosity model assumes that all surfaces are small, opaque, ideal diffuse reflectors (Lambertian).

We apply the radiosity model by determining the differential amount of radiant flux $dB$ leaving each surface point in the scene, and then we sum the energy contributions over all surfaces to obtain the amount of energy transfer between the surfaces. In Figure 26, which illustrates the radiant energy transfer from a surface, $dB$ is the visible radiant flux emanating from the surface point in the direction given by angles $\theta$ and $\phi$ within a differential solid angle $d\omega$ per unit time, per unit surface area.

The intensity $I$ for the diffuse radiation in direction $(\theta, \phi)$ can be described as the radiant energy per unit time per unit projected area per unit solid angle, or

$$I = \frac{dB}{d\omega \cos \phi}$$

(29)

Assuming the surface is an ideal diffuse reflector, we can set the intensity $I$ to a constant for all viewing directions. Thus, $dB/d\omega$ is proportional to the projected surface area (Figure 27). To obtain the total rate of energy radiation from the surface point, we need to sum the radiation for all directions. That is, we want the total energy emanating from a hemisphere centered on that surface point, as in Figure 28, which is

$$B = \int_{\text{hemi}} dB$$

(30)

![Figure 26](image1)

**FIGURE 26**
Visible radiant energy emitted from a surface point in direction $(\theta, \phi)$ within solid angle $d\omega$.

![Figure 27](image2)

**FIGURE 27**
For a unit surface element, the projected area perpendicular to the direction of energy transfer is equal to $\cos \phi$.

![Figure 28](image3)

**FIGURE 28**
Total radiant energy from a surface point is the sum of the contributions in all directions over a hemisphere centered on that surface point.
For a perfect diffuse reflector, \( I \) is a constant, so we can express radiant flux \( B \) as

\[
B = I \int_{\text{hemi}} \cos \phi \, d\omega
\]

Also, the differential element of solid angle \( d\omega \) can be expressed as

\[
d\omega = \frac{dS}{r^2} = \sin \phi \, d\phi \, d\theta
\]

so that

\[
B = I \int_0^{2\pi} \int_0^{\pi/2} \cos \phi \, \sin \phi \, d\phi \, d\theta = I \pi
\]

A model for the light reflections from the various surfaces is formed by setting up an “enclosure” of surfaces (Figure 29). Each surface in the enclosure is either a reflector, an emitter (light source), or a combination reflector-emitter. We designate radiosity parameter \( B_k \) as the total rate of radiant energy leaving surface \( k \) per unit area. Incident energy parameter \( H_k \) is the sum of the radiant energy contributions from all surfaces in the enclosure arriving at surface \( k \) per unit time, per unit area. That is,

\[
H_k = \sum_j B_j F_{jk}
\]

where parameter \( F_{jk} \) is called the form factor for surfaces \( j \) and \( k \). Form factor \( F_{jk} \) is the fractional amount of radiant energy from surface \( j \) that reaches surface \( k \).

For a scene with \( n \) surfaces in the enclosure, the radiant energy from surface \( k \) is described with the following radiosity equation:

\[
B_k = E_k + \rho_k H_k = E_k + \rho_k \sum_{j=1}^n B_j F_{jk}
\]

If surface \( k \) is not a light source, then \( E_k = 0 \). Otherwise, \( E_k \) is the rate of energy emitted from surface \( k \) per unit area (watts/m\(^2\)). Parameter \( \rho_k \) is the reflectivity factor for surface \( k \) (percent of incident light that is reflected in all directions). This reflectivity factor is related to the diffuse reflection coefficient used in empirical illumination models. Plane and convex surfaces cannot “see” themselves, so no self-incidence takes place and the form factor \( F_{kk} \) for these surfaces is 0.
To obtain the illumination effects over the various surfaces in the enclosure, we need to solve the simultaneous radiosity equations for \( n \) surfaces, given the array values for \( E_k, \rho_k, \) and \( F_{jk} \). That is, we must solve

\[
(1 - \rho_k F_{kk}) B_k - \rho_k \sum_{j \neq k} B_j F_{jk} = E_k \quad k = 1, 2, 3, \ldots, n
\]  

or

\[
\begin{bmatrix}
1 - \rho_1 F_{11} & -\rho_1 F_{12} & \cdots & -\rho_1 F_{1n} \\
-\rho_2 F_{21} & 1 - \rho_2 F_{22} & \cdots & -\rho_2 F_{2n} \\
\vdots & \vdots & \ddots & \vdots \\
-\rho_n F_{n1} & -\rho_n F_{n2} & \cdots & 1 - \rho_n F_{nn}
\end{bmatrix}
\begin{bmatrix}
B_1 \\
B_2 \\
\vdots \\
B_n
\end{bmatrix}
= 
\begin{bmatrix}
E_1 \\
E_2 \\
\vdots \\
E_n
\end{bmatrix}
\]  

We then convert to intensity values \( I_k \) by dividing the radiosity values \( B_k \) by \( \pi \).

For color applications, we can calculate the individual red, green, and blue (RGB) components of the radiosity \( (B_{kr}, B_{kg}, B_{kb}) \) using the color components for \( \rho_k \) and \( E_k \).

Before we can solve Equation 35, we must determine the values for form factors \( F_{jk} \). We do this by considering the energy transfer from surface \( j \) to surface \( k \) (Figure 30). The rate of radiant energy falling on a small surface element \( dA_k \) from area element \( dA_j \) is

\[
\text{dB}_j \, dA_j = (I_j \cos \phi_j \, d\omega) \, dA_j
\]

However, solid angle \( d\omega \) can be written in terms of the projection of area element \( dA_k \) perpendicular to the direction \( dB_j \) as

\[
d\omega = \frac{dA}{r^2} = \frac{\cos \phi_k \, dA_k}{r^2}
\]

Therefore, we can express Equation 37 in the form

\[
\text{dB}_j \, dA_j = \frac{I_j \cos \phi_j \cos \phi_k \, dA_j \, dA_k}{r^2}
\]

The form factor between the two surfaces is the percent of energy emanating from area \( dA_j \) that is incident on \( dA_k \):

\[
F_{dA_j, dA_k} = \frac{\text{energy incident on } dA_k}{\text{total energy leaving } dA_j} = \frac{I_j \cos \phi_j \cos \phi_k \, dA_j \, dA_k}{r^2} \cdot \frac{1}{B_j \, dA_j}
\]
Also, $B_j = \pi I_j$, so

$$F_{dA_j, dA_k} = \frac{\cos \phi_j \cos \phi_k dA_k}{\pi r^2}$$  \hspace{1cm} (41)

and the fraction of emitted energy from area $dA_j$ incident on the entire surface $k$ is

$$F_{dA_j, A_k} = \int_{\text{surf}_j} \frac{\cos \phi_j \cos \phi_k}{\pi r^2} dA_k$$  \hspace{1cm} (42)

where $A_k$ is the area of surface $k$. We can then define the form factor between the two surfaces as the area average of the above expression, which is

$$F_{jk} = \frac{1}{A_j} \int_{\text{surf}_j} \int_{\text{surf}_k} \frac{\cos \phi_j \cos \phi_k}{\pi r^2} dA_k dA_j$$  \hspace{1cm} (43)

The two integrals in Equation 43 are evaluated using numerical integration techniques and stipulating the following conditions:

- $\sum_{k=1}^{n} F_{jk} = 1, \ for\ all\ k$ (conservation of energy)
- $A_j F_{jk} = A_k F_{kj}$ (uniform light reflection)
- $F_{jj} = 0, \ for\ all\ j$ (only plane or convex surface patches)

To apply the radiosity model, we subdivide each surface in a scene into many small polygons. The realistic appearance of the displayed scene is improved as we decrease the size of the polygon subdivisions, but more time is then needed to render the scene. We can speed up the calculation of the form factors by using a hemispheres to approximate the hemisphere. This replaces the spherical surface with a set of linear (plane) surfaces. Once the form factors are evaluated, we can solve the simultaneous linear equations 35 using a numerical technique such as Gaussian elimination or LU decomposition. Alternatively, we could start with approximate values for the $B_j$ and solve the set of linear equations equations iteratively using the Gauss-Seidel method. At each iteration, we calculate an estimate of the radiosity for surface patch $k$ using the previously obtained radiosity values in the radiosity equation

$$B_k = E_k + \rho_k \sum_{j=1}^{n} B_j F_{jk}$$

We could then display the scene at each step to observe the improvement in surface rendering. This process is repeated until there is little change in the calculated radiosity values.

**Progressive Refinement Radiosity Method**

Although the radiosity method produces highly realistic surface renderings, considerable processing time is needed to calculate the form factors and there are tremendous storage requirements. Using progressive refinement, we can restructure the iterative radiosity algorithm to speed up the calculations and reduce storage requirements at each iteration.

From the radiosity equation, the radiant energy transfer between two surface patches is calculated as

$$B_k \ \text{due to} \ B_j = \rho_k B_j F_{jk}$$  \hspace{1cm} (44)

Reciprocally,

$$B_j \ \text{due to} \ B_k = \rho_j B_k F_{kj}, \ \text{for all } j$$  \hspace{1cm} (45)
which we can rewrite as

\[ B_j \text{ due to } B_k = \rho_j B_k F_{jk} \frac{A_j}{A_k}, \text{ for all } j \]  \hspace{1cm} (46)

This relationship is the basis for the progressive refinement approach to the radiosity calculations. Using a single surface patch \( k \), we can calculate all form factors \( F_{jk} \) and consider the light transfer from that patch to all other surfaces in the environment. With this procedure, we need only compute and store parameter values for a single hemicube and the associated form factors. At the next iteration, we replace these parameter values with values for another selected patch; and we can display the progressive improvements in the surface rendering as we proceed from one selected patch to another.

Initially, we set \( B_k = E_k \) for all surface patches. We then select the patch with the highest radiosity value, which is the brightest light emitter, and calculate the next approximation to the radiosity for all other patches. This process is repeated at each step, so that light sources are chosen first in order of highest radiant energy, and then other patches are selected based on the amount of light received from the light sources. The steps in a simple progressive refinement approach are outlined in the following algorithm:

```c
for each patch k
    /* Set up hemicube and calculate form factor F[j][k]. */
    for each patch j {
    }
    dB[k] = 0;
```

At each step, the surface patch with the highest value for \( \Delta B_k A_k \) is selected, because radiosity is a measure of radiant energy per unit area. Also, we choose the initial values as \( \Delta B_k = B_k = E_k \) for all surface patches. This progressive refinement algorithm approximates the actual propagation of light through a scene as a function of time.

Displaying the rendered surfaces at each step produces a sequence of views that proceeds from a dark scene to a fully illuminated one. After the first step, the only surfaces illuminated are the light sources and those nonemitting patches that are visible to the chosen emitter. To produce more useful initial views of the scene, we could set an ambient light level so that all patches have some illumination. At each stage of the iteration, we then reduce the ambient light according to the amount of radiant energy transfer into the scene.

Once a radiosity solution (either complete, or progressive) has been calculated, it is possible to view the scene from any point without require additional computation. Thus, radiosity is inherently view-independent (as opposed to ray-tracing, which is highly view-dependent). However, a change in position of any object in the scene will result in a change in form factors, which will require re-solving the radiosity equations.

Color Plate 29 shows a scene rendered with the progressive-refinement radiosity model. Various lighting conditions in radiosity renderings are illustrated in Color Plates 30 and 31. Ray-tracing methods are often combined with the radiosity model to produce highly realistic diffuse and specular surface shadings, as in Color Plate 28.
3 Environment Mapping

An alternate procedure for modeling global reflections is to define an array of intensity values that describes the environment around a single object or a group of objects. Instead of using interobject ray tracing or the radiosity calculations to pick up the global specular and diffuse illumination effects, we simply map the environment array onto an object in relationship to the viewing direction. This procedure is called environment mapping, and it is sometimes referred to as reflection mapping (although transparency effects could also be modeled with the environment map). Another name for environment mapping is “the poor person’s ray-tracing method,” because it is a cheap and fast approximation of the more accurate global-illumination rendering techniques that we discussed in Sections 1 and 2.

The environment map is defined over the surfaces of an enclosing universe. Information in the environment map includes intensity values for light sources, the sky, and other background objects. Figure 31 shows the enclosing universe as a sphere, but a cube or a cylinder is often used to define the environment surfaces surrounding the objects in a scene.

To render the surface of an object, we project pixel areas onto the object surface and then reflect each projected pixel area onto the environment map to pick up the surface intensity values for the pixel. If the object is transparent, we can also refract the projected pixel area to the environment map. The environment-mapping process for reflection of a projected pixel area is illustrated in Figure 32. Pixel intensity is determined by averaging the intensity values within the intersected region of the environment map.

**Figure 31**
A spherical enclosing universe, with the environment map on the surface of the sphere.

**Figure 32**
Projecting a pixel area to a surface, then reflecting the area to the environment map.
4 Photon Mapping

Although the radiosity method can produce accurate displays of global illumination effects for simple scenes, the method becomes more difficult to apply as the complexity of a scene increases. Both the rendering time and the storage requirements become prohibitive for very complicated scenes, and many illumination effects are difficult to model correctly. **Photon mapping** provides a general method for modeling global illumination in complex scenes that is both efficient and accurate.

The basic concept in photon mapping is to separate the illumination information from the geometry of a scene. Ray paths are traced through the scene from all light sources, and the lighting information from the ray-object intersections is stored in a **photon map**. Distributed ray-tracing methods are then applied using incremental algorithms similar to those employed in radiosity rendering.

Light sources can be designated as points, directional spotlights, or any other configuration. The assigned intensity for a light source is divided among its rays (photons), and the ray directions are distributed randomly. A point light source is modeled by generating ray paths uniformly in all directions unless the source is directional. For other light sources, random positions on the source are selected and rays are generated in random directions. More rays are generated from brighter lights than from low-power light sources. In addition, *projection maps*, which store binary information about whether there are objects in any region of the space, can be constructed for the light sources. Bounding spheres can also be used in the algorithm to provide object information within large spatial regions. Any number of rays can be generated for a scene, and the accuracy of the illumination effects improves as more ray paths are generated.

Photon mapping is particularly adept at simulating two real-world effects that are more difficult to obtain through ray-tracing and radiosity methods: **caustics** and **diffuse inter-reflections**. Caustics are patterns that result from the reflection or refraction of light. An example of this is the rippling light effect often seen on the bottom of a swimming pool. As light passes through the water in the pool, it is focused in different ways as the water moves. Another type of caustic effect comes from the reflection of light off shiny, curved surfaces, such as the inside of a polished brass ring. In this type of caustic, the reflected light produces a cardioid pattern.

As the name suggests, diffuse inter-reflections are the result of the reflection of light between diffuse surfaces. This type of reflection is responsible for the “color bleeding” effect that is seen when light reflects from a brightly colored surface onto another surface.

Typically, photon mapping is implemented using a two-pass method. The first pass distributes photons throughout the scene. Photons are emitted from each light source in varying numbers and directions based on characteristics of the light source. The power of an individual photon is determined by the original power (wattage) of the light source and the number of photons being generated from that source. The emitted photons are traced through the scene until they strike an object, at which point they may be reflected, transmitted, or absorbed by the object, based upon a probability distribution for that object. At each intersection with an object, information about the photon’s incident direction, incoming power, and the intersection point are added to the photon map. Reflected and transmitted photons are followed as they continue through the scene.

Once the photon map has been built, the second pass renders the image from the collected photon information. This is commonly implemented using a
modified form of ray-tracing. As rays are traced into the scene, the information collected in the photon map is used to produce an estimate of the radiance exiting from the intersection point along the ray. That estimate is then used in the calculation of the pixel color.

As with a radiosity solution, a photon map is view-independent. Once the photon energy has been distributed, the scene can be viewed from any position without requiring that photon tracing be performed again.

5 Summary

Ray tracing is a method for obtaining global, specular reflection and transmission effects by tracing light paths through a scene to pixel positions. Pixel rays are traced through a scene, bouncing from object to object while accumulating intensity contributions. A ray-tracing tree is constructed for each pixel, and intensity values are combined from the terminal nodes of the tree back to the root. Object-intersection calculations in ray tracing can be reduced with space-subdivision methods that test for ray-object intersections only within subregions of the total space. Distributed ray tracing employs multiple rays per pixel, randomly assigning various ray parameters, such as direction and time. This provides an accurate method for modeling surface gloss and translucency, finite camera apertures, extended light sources, shadow effects, and motion blur.

Radiosity methods provide accurate modeling for diffuse-reflection effects by calculating radiant energy transfer between the various surface patches in a scene. Progressive refinement is used to speed up the radiosity calculations by considering energy transfer from one surface patch at a time. Highly photo-realistic scenes are generated using a combination of ray tracing and radiosity.

A fast method for approximating global illumination effects is environment mapping. An environment array is used to store information on background intensity for a scene. This array is then mapped to the objects in a scene based on the specified viewing direction.

Photon mapping provides an accurate and efficient model for global illumination in complex scenes. Random rays are generated from the light sources, and the illumination effects for each ray are stored in a photon map, which separates the lighting information from the scene geometry. The accuracy of the illumination effects improves as more rays are generated.

REFERENCES


EXERCISES
1 Write a program to implement the basic ray-tracing algorithm for a scene containing two spheres hovering over a white ground square that has evenly spaced vertical black stripes down its length. The scene is to be illuminated with a single point light source at the viewing position.
2 Write a program to implement the basic ray-tracing algorithm for a scene containing any specified arrangement of spheres and polygon surfaces illuminated by a given set of point light sources. The program should allow the user to increase or decrease the intensity of each light source via keyboard input.
3 Write a program to implement the basic ray-tracing algorithm using space-subdivision methods for any specified arrangement of spheres and polygon surfaces illuminated by a given set of point light sources. The program should allow the user to increase or decrease the intensity of each light source via keyboard input.
4 Write a program to implement the following features of distributed ray tracing: pixel sampling with 16 jittered rays per pixel, distributed reflection directions (gloss), distributed refraction directions (translucency), and extended light sources.
5 Set up an algorithm for modeling the motion blur of two spheres moving in opposite directions using distributed ray tracing.
6 Implement the basic radiosity algorithm for rendering the inside surfaces of a regular pentagon when one inside face of the pentagon is a light source.
7 Devise an algorithm for implementing the progressive-refinement radiosity method.
8 Apply the algorithm in developed in the previous exercise to the pentagon example in Exercise 6.
9 Write a routine to transform an environment map to the surface of a sphere.

IN MORE DEPTH
1 Implement the basic ray tracing algorithm to replace the existing illumination model used to render your scene. Record the average rendering time for the scene.
2 Choose one of the methods for improving the performance of the basic ray-tracing algorithm (e.g., using bounding volumes to reduce object-intersection calculations, subdivision methods). Render your scene using these improvements and record the average rendering time. How much of an improvement do you obtain by incorporating these methods?
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Global Illumination
Color Plates

**Color Plate 25**
A ray-traced scene, showing global reflection and transparency effects. (Courtesy of Evans & Sutherland.)

**Color Plate 26**
A “sphereflake” rendered with ray tracing using 7,381 spheres and 3 light sources. (Courtesy of Eric Haines. Reprinted with permission.)
**Color Plate 27**
A room scene illuminated with 5 light sources (a) was rendered using the ray-tracing light-buffer technique to process shadow rays. A closeup (b) of part of the room shown in (a) illustrates the global illumination effects. The room is modeled with 1,298 polygons, 4 spheres, 76 cylinders, and 35 quadrics. Rendering time was 246 minutes on a VAX 11/780, compared to 602 minutes without using light buffers. (Courtesy of Eric Haines. Reprinted with permission.)

**Color Plate 28**
A scene showing the focusing, antialiasing, and illumination effects possible with a combination of ray-tracing and radiosity methods. Realistic physical models of light illumination were used to generate the refraction effects, including the caustic in the shadow of the glass. (Courtesy of Peter Shirley, Computer Science Department, University of Utah.)

**Color Plate 29**
Nave of Chartres Cathedral rendered with a progressive-refinement radiosity model by John Wallace and John Lin, using the Hewlett-Packard Starbase Radiosity and Ray Tracing software. Radiosity form factors were computed with ray-tracing methods. (Courtesy of John Wallace, Autodesk, Inc. Reprinted with permission.)
Color Plate 30

Color Plate 31
In the earliest days of computer-generated imagery, virtually the entire image generation process was under the control of the programmer. Hardware vendors provided libraries of functions that facilitated low-level access to their products, but libraries of routines for drawing primitives and altering their attributes were nonexistent or very rudimentary in nature. As a result, the programmer was required to specify every aspect of the image generation process. While this provided tremendous control of the final image, it also required significant investments of time and effort. Programmers typically developed their own algorithms for implementing primitives and converting their ideas into displayable result images. Furthermore, programs developed to use one vendor’s graphics hardware were typically unusable on other hardware without significant modification.

As the field of computer graphics began to mature, standard libraries of graphics routines became common. The development of the Graphical Kernel System (GKS) in 1984 and the subsequent...
Programmable Shaders

development of the Programmer’s Hierarchical Interactive Graphics System (PHIGS) and OpenGL made it possible to create images in a more device-independent way. A program written to use the GKS application programming interface (API), for instance, could be used on any system for which a GKS library was available with relatively little modification. With the simplification of use, however, came a reduction in control over the image generation process. The graphics API provided a standardized interface to the hardware; internally, the API processed all primitive and attribute requests by the programmer in a fixed way. This internal organization is commonly known as a fixed-function rendering pipeline.

As the capabilities of widely available graphics hardware continued to increase, programmers looked for ways to access the increasingly powerful hardware beyond the constraints of the fixed-function pipeline. To facilitate this, API developers provided “hooks” into the rendering pipeline, which allow modification of the behavior of certain stages of the pipeline programmatically through the use of programmable shaders. These shaders replace the built-in fixed functionality of those stages with whatever behavior the programmer desires, making it possible to achieve effects that would be difficult or impossible with the original fixed-function pipeline.

1 A History of Shading Languages

By the mid 1980s, computer-generated imagery had become a standard tool in the movie and advertising industries. The generation of realistic-looking images was a common area of research. To a large extent, realism was limited by the sophistication of the methods used to shade the surfaces of objects. Given the algorithmic nature of shading calculations, the development of specialized languages for expressing surface shading methods was an obvious step.

Cook’s Shade Trees

One of the first shader-specific languages was described by Rob Cook in 1984. Cook discussed a system for integrating shading and texturing techniques which he called shade trees. Shaders are represented as expression tree data structures. For example, the classic Phong model describes the reflections from a surface illuminated by a single point light source as

\[ I = k_a I_a + k_d I_l (\mathbf{N} \cdot \mathbf{L}) + k_s I_l (\mathbf{N} \cdot \mathbf{V})^n \]  

This equation can be represented as a binary expression tree, shown in Figure 1.

Cook’s approach was to use a generalized tree structure to represent the way in which appearance parameters such as surface normals, material properties, texture maps, and light properties should be combined to determine the color of an object’s surface. Leaf nodes represent basic data; parent nodes represent ways in which that data can be combined. Many types of operation nodes are built into the language: arithmetic and trigonometric operations; mathematical operations such as square root and vector normalization; shading functions for calculating ambient, diffuse, and specular light values; and additional supporting operations such as proportional blending (mixing) of color information. In addition to surface shading, trees can be built to describe the characteristics of light sources and to represent atmospheric effects.
Cook developed a C-like specification language for shade trees, which could be used in conjunction with a modeling language to build shade trees and attach them to the surfaces of objects. Here is the source code for a shade tree describing metallic surfaces, taken from Cook’s 1984 paper:

```c
float a = .5, s = .5;
float roughness = .1;
float intensity;
color metal_color = (1, 1, 1);
intensity = a * ambient() +
    s * specular(normal, viewer, roughness);
final_color = intensity * metal_color;
```

The ambient and specular functions are built into the language, and they return the amount of and characteristics of each type of light striking the object surface. This code would be translated into the shade tree shown in Figure 2. The shade tree could then be applied to generate a bronze surface appearance with the following statement:

```plaintext
surface "metal",
    "metal_color", material bronze,
    "roughness", .15
```

which overrides the default values for the roughness and metal_color parameters with ones appropriate to a bronze surface.
Perlin’s Pixel Stream Editor

Another notable example of an early shading language was developed by Ken Perlin in 1985 as part of an image processing filter that he called the Pixel Stream Editor (PSE). The PSE processed an input image that contained not only pixel color information, but also data describing surfaces of objects within the image; the image was manipulated by running a program written in Perlin’s shading language on each pixel.

Like Cook’s shade tree language, Perlin’s language had built-in operators and functions for specifying the way in which color information was to be computed. Perlin’s language was a much higher-level language than Cook’s, however. It included control flow statements in the form of selection and iteration constructs as well as support for user-written functions, significantly increasing the power and flexibility of shaders.

Although the concept of procedural texturing was not new, most work in this area had been done with functions operating in a two-dimensional domain. Perlin proposed extending this to three dimensions, creating what he referred to as space functions. Conceptually, such a function can be thought of as representing a solid volume in space. If the function is evaluated at the points on the visible surface of an object, the resulting values effectively “sculpt” the object out of the material, acting as a solid texture.

Perhaps the most important innovation in Perlin’s work, though, was the concept of noise. Materials in the real world often have some sort of random or stochastic component to them. Examples of this are the grain seen in cut wood, the meandering pattern of rivers, and the branching patterns of ferns. None of these patterns are truly random, however, so traditional approaches to randomness are not appropriate for modeling them.

Perlin’s basic noise function takes a three-dimensional vector as its argument and returns a pseudo-random floating-point value, which can be used in any situation where randomness is wanted. Applying it to points on the visible surface of an object, we can introduce random variations in surface color by multiplying the base color by the noise value. It can also be used to generate pseudo-random perturbations of surface normals for bump mapping, or to actually move points in space.

What makes the noise function useful are the characteristics of the result values. Operation of the function is statistically invariant under translation and rotation, so using those transformations on the function domain does not affect the characteristics of the result distribution. The function is frequency-controlled, so it is possible to increase the detail in the result distribution by scaling the argument—for example, $\text{Noise}(2^x)$ will have twice the frequency of $\text{Noise}(x)$. Results are band-limited, so variation in results will occur within a limited range.

The concept of noise has become so valuable in the world of shaders that most shader languages today provide at least one type of noise function. In addition, some graphics hardware provide hardware acceleration for noise.

RenderMan

One of the most widely used shader languages is the RenderMan Shading Language (RSL) designed by Pixar. It dates from 1988, when Pixar first published the RenderMan Interface Specification (RISpec). Based originally on the shade tree work by Cook, RenderMan became the industry-standard shading language for batch-oriented rendering in the entertainment industry. As RenderMan is a specification rather than a particular product, a number of implementations are available, including Pixar’s own Photorealistic RenderMan (prman)
package as well as several open-source implementations such as Blender, Pixie, and Aqsis.

The RISpec actually consists of two parts. The first is the RenderMan Interface, which specifies a standard interface between modeling programs and rendering programs. It lists the features that a rendering program must support, and it defines the API to be used by programs wanting to communicate with the rendering program.

The second part of the RISpec is the RSL specification. RSL is a C-like language with data types and built-in operations and functions that are designed to facilitate the development of shaders.

The RenderMan Interface gives shader developers a tremendous amount of control over the rendering process through the different types of shaders that can be developed. Each shader type manages one aspect of the rendering process. Illumination can be managed through the use of light shaders; the geometry of objects can be modified by displacement shaders, and surface shaders can be used to compute surface color at each shading point. Volume shaders allow for simulation of atmospheric effects such as fog and dust, as well as the effect of light passing through the interior of transparent or translucent objects. Finally, imager shaders can be used to modify the pixel values produced by the rendering pipeline, allowing the implementation of post-processing effects such as simulating the brush strokes in an oil painting.

An interesting feature of the RenderMan Interface is that it can be implemented in multiple ways. The RISpec defines a C-language API, much like the OpenGL API that we have used throughout this textbook. The RenderMan Interface can be provided as a C-callable library, with a full set of functions for creating standard types of objects and manipulating the rendering state. Programmers can write programs that use these routines to perform any operation that the RenderMan Interface allows. Here is an example of a C program fragment that draws a red polygon using a plastic surface shader:

```c
#include <ri.h>

RtPoint Poly[4] = {
    { 1, 1, -1 }, { -1, 1, -1 }, { -1, -1, -1 }, { 1, -1, -1 }
};

void main()
{
    RiBegin (RI_NULL);
    RiWorldBegin ();
    RiColor (1.0, 0.0, 0.0);
    RiSurface ("plastic", RI_NULL);
    RiPolygon (4, RI_P, (RtPointer)Poly, R_NULL);
    RiWorldEnd ();
    RiEnd ();
}
```

Alternatively, the RenderMan Interface can be implemented as a standalone application, such as Pixar’s prman implementation. This type of implementation must take as its input the RenderMan Interface Bytestream (RIB), defined in the RISpec. RIB statements provide a compact, easy-to-read way to specify scene parameters; rather than calling functions to create light sources, define objects, and so on, those functions are requested through RIB statements. Because they are text files, RIB files can be created and manipulated by programs—for instance, a
Program could read in an existing RIB file that specifies one frame of an animation, modify a statement that translates the position of an object, and write the modified version to a new file that could then be used to generate the next frame of the animation. In addition, because RIB statements are often more straightforward and easy to read than C programs, they may be easier for non-programmers to use. Here is a RIB file equivalent to the C program fragment shown previously:

```
WorldBegin
  Color [ 1.0, 0.0, 0.0 ]
  Surface "plastic"
  Polygon "P" [ 1 1 -1 -1 1 -1 -1 -1 -1 1 -1 -1 ]
WorldEnd
```

### 2 The OpenGL Pipeline

OpenGL has been revised a number of times since its introduction in 1992. Each revision has typically either added functionality to OpenGL, or helped adapt it to improvements in graphics hardware. (We discuss the evolution of OpenGL in more detail in Appendix C.) One of the most significant changes was the introduction of the OpenGL Shading Language, often referred to as GLSL. To understand how GLSL fits into the internal structure of OpenGL, we begin by looking at the way in which OpenGL processes geometry and pixel information.

The Fixed-Function Pipeline

The original internal structure of OpenGL was a sequence of processing steps organized into a two-channel rendering pipeline. The pipeline stages were fixed in nature—that is, they performed specific operations in response to data sent through them—and therefore the overall structure became known as the fixed-function OpenGL pipeline. Conceptually, the processing steps can be thought of as having the structure shown in Figure 3.

Different types of information are processed by the upper and lower halves of the pipeline. Geometric primitives are processed through the upper part of the diagram (often referred to as the geometric pipeline), while pixel primitives are processed by the lower part (the pixel pipeline). Both types of information can be
saved in display lists; when a display list is executed, the information it contains is sent to the appropriate parts of the pipeline.

In the geometric pipeline, primitives are described as collections of vertices along with additional attributes such as material properties, associated texture coordinates, normal vectors, and so on. The per-vertex operations and primitive assembly stage performs a number of processing steps on this information. Vertices are transformed by the modelview matrix. If automatic texture generation is enabled, new texture coordinates are generated and replace the original texture coordinates for the vertex. Lighting calculations are performed, followed by clipping and shading operations. Finally, the primitives are rasterized, which determines which pixel positions are occupied by each primitive.

Pixel information is processed in a similar fashion. Pixel data are retrieved from main memory, pixel buffers, texture memory, or the frame buffer. Additional processing is performed on the data, which is then either written into texture memory (if texture mapping is being performed) or rasterized.

The rasterized geometry and pixel data are combined into a set of fragments. Fragments are per-pixel data structures that contain all the information needed to update the data held in the frame buffer. Once created, fragments are processed by the per-fragment operations stage, which performs the final conversion to display form. Fragments which are being texture-mapped have texels generated from information in texture memory; fog and antialiasing processing are performed if they are needed. Finally, depth-buffer testing is performed, and the resulting set of fragments are written to the frame buffer.

**Changing the Pipeline Structure**

One problem with the classic fixed-function pipeline is that it no longer matches the way that modern graphics hardware operates. It is unable to take full advantage of the power available in even low-cost graphics cards for computers, with the result that rendering performance may suffer.

Consider the modified view of the OpenGL pipeline shown in Figure 4. In this pipeline, the per-vertex and per-fragment fixed processing stages have been replaced by user-programmable processing stages. This change allows the application program to determine what processing should be done at each of these points in the rendering pipeline. It also allows the OpenGL implementation to take better advantage of improving hardware capabilities, with the result that rendering can be accelerated significantly (depending on what hardware functionality is present).
Programmable Shaders

The most recent OpenGL versions have three additional user-programmable stages: a geometry processor, and two tessellation processors.

Application programs control the operation of these programmable stages through the use of shaders. Shaders are small program fragments that are loaded into OpenGL programs and attached to the appropriate processing element in the OpenGL pipeline, replacing the fixed functionality of the pipeline.

Vertex Shaders

A vertex shader is a shading program designed to replace the fixed vertex processing stage of the pipeline. It will be executed on every vertex sent into the pipeline, and is responsible for producing all the information needed by later stages in the pipeline; minimally, it must output the vertex itself, transformed into clip space by the projection and modelview matrices. Clip space is the coordinate space used throughout the rest of the OpenGL pipeline. All vertices must be transformed into this space before they are used.

Beyond this basic functionality, a vertex shader can also associate a color with the vertex, can generate or transform texture coordinates for later use in the pipeline, and can even use lighting information and surface normals at the vertex. Vertex shaders are able to access this information through the use of special built-in global variables initialized by the OpenGL implementation and pass the modified data to the rest of the pipeline through another set of global variables.

Fragment Shaders

Like the vertex shader, a fragment shader operates on information residing in the pipeline and produces modified data for use by the rest of the pipeline’s stages. As the name implies, fragment shaders operate on the rasterized vertex and pixel information, called fragments. The fragment shader will be executed once for each fragment coming through the pipeline; depending on how the primitive has been rasterized, the fragment shader may be executed many more times than the vertex shader.

At a minimum, a fragment shader is responsible for assigning the fragment color based on the basic color of the object. However, fragment shaders are also responsible for applying textures and performing operations such as bump mapping.

Geometry Shaders

Geometry shaders perform further processing on the results from the primitive assembly stage of the pipeline. A geometry shader will be executed once for each primitive and will have access to all the vertex information associated with that primitive. Unlike the vertex and fragment shaders, however, a geometry shader is not limited to passing on a modified version of the incoming data; it may actually create new primitives, which are then processed by the rest of the pipeline.

Tessellation Shaders

The tessellation processor is controlled by a pair of shaders: a tessellation control shader and a tessellation evaluation shader. These shaders operate on a type of primitive called a patch. Patches are collections of vertices, per-vertex attributes, and additional per-patch attributes. The tessellation shaders take an incoming patch, subdivide the patch into a collection of points, lines, or triangles, and output the resulting patch information for processing by the rest of the pipeline.
3 The OpenGL Shading Language

GLSL is a C-like language designed to directly support the development of shaders. It has a wide variety of data types for representing typical shading data items such as vectors, colors, and matrices, along with a collection of built-in operators that simplify manipulation of those data items.

The designers of GLSL attempted to create a shading language that met a number of fairly ambitious goals. They wanted a high-level, easy-to-use programming language that would work well with OpenGL. It needed to be as hardware-independent as possible, to allow the same shaders to be used with graphics hardware from different manufacturers. In addition, because graphics hardware continues to evolve, the language should not be tied to a particular type or generation of hardware; rather, it should be powerful enough to take advantage of the capabilities of the underlying hardware and flexible enough in its design that it can accommodate the rapid evolution of that hardware.

Although the language looks very much like C or C++, it’s important to remember that GLSL really isn’t either one of them. There are several differences in the way that function parameters are handled, and the language is much stricter with respect to type checking issues. In addition, many familiar C and C++ data types and language constructs (such as pointer variables and widespread implicit conversion between data types) were intentionally not included in GLSL.

Like OpenGL, GLSL has evolved since it was first introduced. New features have been added, and existing features have been deprecated (that is, marked for future removal from the language) in favor of new features. Depending on which version of GLSL your implementation supports, some features may not be available. The following code can be used to determine your versions of OpenGL and GLSL:

```c
printf ("OpenGL version: %s\n",
   (char *) glGetString (GL_VERSION));

printf ("GLSL version: %s\n",
   (char *) glGetString (GL_SHADING_LANGUAGE_VERSION));
```

The first statement prints out a string containing version information for your OpenGL implementation, and the second prints out the corresponding GLSL version information.

We also note that our discussion of GLSL will, necessarily, be incomplete. The language has far too many features to allow us to explore them completely in a single chapter. Instead, we will discuss enough details of shader creation and use that you may begin to experiment with your own shaders. For more in-depth study of GLSL, any of the GLSL references described at the end of this chapter should suffice.

Shader Structure

Most GLSL programs will contain both a vertex shader and a fragment shader. Each shader contains a main routine which, in fact, is a function named `main`. Shaders may also contain supporting functions, as well as global variables to facilitate communication between the vertex and fragment shaders.

A shader’s main routine will vary depending on what functionality is required, but certain operations must be performed. As mentioned earlier, a vertex shader will be executed on every vertex that passes through the pipeline;
even if it does nothing else, it must transform the vertex into clip space. This is accomplished by multiplying the vertex by the modelview matrix, and then multiplying that result by the projection matrix. Each of these values is available to the vertex shader as a built-in global variable. These built-in global variables all have names beginning with the character sequence `gl_`; to transform the vertex into clip space, the vertex shader must use the vertex position (`gl_Vertex`) and the contents of the modelview and projection matrices (`gl_ModelViewMatrix`, `gl_ProjectionMatrix`). The transformed vertex must be placed in the global variable `gl_Position` so that the next stage in the OpenGL pipeline can use it. Here is an example of a minimal vertex shader:

```c
void main ()
{
    gl_Position = gl_ProjectionMatrix *
                 (gl_ModelViewMatrix * gl_Vertex);
}
```

There are other ways to perform this transformation. An additional global variable named `gl_ModelViewProjectionMatrix` contains the product of the projection and modelview matrices and can be used to reduce the transformation to one multiplication. Because this operation is standard, a built-in function is also available to perform it:

```c
gl_Position = ftransform ();
```

Another common operation performed in vertex shaders is the assignment of a color to the vertex. This is accomplished by assigning a color value to the global variable `gl_FrontColor`, as follows:

```c
void main ()
{
    gl_Position = gl_ProjectionMatrix *
                 (gl_ModelViewMatrix * gl_Vertex);
    gl_FrontColor = gl_Color;
}
```

The `gl_Color` variable contains whatever color the OpenGL application associated with the vertex by calling `glColor`. As you might expect, there is also a `gl_BackColor` global variable, which can be used when two-sided lighting is being used in the OpenGL program.

Fragment shaders are responsible for computing the color associated with a fragment. At minimum a fragment shader must assign that color to the global variable `gl_FragColor`. The shader can compute the color, or can retrieve whatever color was assigned by the vertex shader, as in this example:

```c
void main ()
{
    gl_FragColor = gl_Color;
}
```

It is important to note that although it appears that this fragment shader is accessing the same global variable used by the vertex shader, the contents of `gl_Color` are modified by the OpenGL pipeline between the execution of the two shaders.
The fragment shader will see either the `gl_FrontColor` or the `gl_BackColor`, depending upon which side of the primitive the fragment being processed belongs to.

**Using Shaders in OpenGL**

Unlike OpenGL programs themselves, shader programs are not precompiled. Instead, they are compiled during the execution of the OpenGL program itself. The process involves a series of steps; for example, assuming that we are using both a vertex and a fragment shader, we would do the following:

1. Create two shader objects.
2. Attach the source for each shader to its shader object.
3. Compile the shaders.
4. Create a program object.
5. Attach the shader objects to the program object.
6. Link the program.

The shader source code must be a null-terminated C-style string (a sequence of characters, followed by a trailing byte containing the value 0). Commonly, the shader source code is put into a text file, from which it is read into the OpenGL program as a single string. Here is an example function that reads the contents of a file into a dynamically allocated string buffer. The function opens the file and determines how many characters are in it. It then allocates a string buffer, reads the file into that as a single string, and returns the pointer to the string buffer.

```c
#include <stdio.h>
#include <stdlib.h>

/*
 ** Create a null-terminated string from the contents of a file
 ** whose name is supplied as a parameter. Return a pointer to
 ** the string, unless something goes wrong, in which case return
 ** a null pointer.
 */

GLchar *readTextFile( const char *name ) {
    FILE *fp;
    GLchar *content = NULL;
    int count = 0;

    /* verify that we were actually given a name */
    if (name == NULL) return NULL;

    /* attempt to open the file */
    fp = fopen( name, "rt" ); /* open the file */
    if (fp == NULL) return NULL;

    /* determine the length of the file */
    fseek (fp, 0, SEEK_END);
    count = ftell (fp);
    rewind( fp );
```

---

*Programmable Shaders*

The fragment shader will see either the `gl_FrontColor` or the `gl_BackColor`, depending upon which side of the primitive the fragment being processed belongs to.

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    FILE *fp;
    GLchar *content = NULL;
    int count = 0;

    /* verify that we were actually given a name */
    if (name == NULL) return NULL;

    /* attempt to open the file */
    fp = fopen( name, "rt" ); /* open the file */
    if (fp == NULL) return NULL;

    /* determine the length of the file */
    fseek (fp, 0, SEEK_END);
    count = ftell (fp);
    rewind( fp );
```
/* allocate a buffer and read the file into it */
if (count > 0) {
    content = (GLchar *) malloc (sizeof(char) * (count+1));
    if (content != NULL) {
        count = fread (content, sizeof(char), count, fp);
        content[count] = '\0';
    }
}
fclose(fp);
return content;
}

To create our shader program, we must first create two shader objects:

GLuint vertShader, fragShader;
vertShader = glCreateShader (GL_VERTEX_SHADER);
fragShader = glCreateShader (GL_FRAGMENT_SHADER);

Each call to glCreateShader returns a handle that is associated with a shader object. We use this handle whenever we need to refer to the shader object, such as when we want to attach source code to it.

Next, we read in the source for each shader. There is no restriction on the names of shader source files; assuming that our vertex shader is in a file named simpleShader.vert and our fragment shader is in a file named simpleShader.frag, we can read them into our program as follows:

GLchar *vertSource, *fragSource;
vertSource = readTextFile("simpleShader.vert");
if (vertSource == NULL) {
    fputs("Failed to read vertex shader\n", stderr);
    exit(EXIT_FAILURE);
}
fragSource = readTextFile("simpleShader.frag");
if (fragSource == NULL) {
    fputs("Failed to read fragment shader\n", stderr);
    exit(EXIT_FAILURE);
}

Now that we have the source strings, we must attach them to the shaders:

glShaderSource (vertShader, 1,
    (const GLchar **) &vertSource, NULL);
glShaderSource (fragShader, 1,
    (const GLchar **) &fragSource, NULL);
free (vertSource);
free (fragSource);
This function allows us to attach several shader source strings to the same shader object. The first parameter is the shader object to be used. The second parameter is the number of source strings to be attached; the third parameter is an array of pointers to the strings. Finally, the fourth parameter tells \texttt{glShaderSource} that the strings are terminated by null characters. \texttt{glShaderSource} makes a copy of the string contents, so once we have attached the source strings to the shader objects, we can deallocate the strings to reduce our memory use.

The next step is to compile the shaders:

\begin{verbatim}
glCompileShader (vertShader);
glCompileShader (fragShader);
\end{verbatim}

It is a good idea to verify that the compilation succeeded. We can retrieve the compilation status with the \texttt{glGetShaderiv} function. If the compilation succeeded, the status will be \texttt{GL_TRUE}:

\begin{verbatim}
GLint status;

glGetShaderiv (vertShader, GL_COMPILE_STATUS, &status);
if (status != GL_TRUE ) {
      fputs ("Error in vertex shader compilation\n", stderr);
      exit (EXIT_FAILURE);
}

glGetShaderiv (fragShader, GL_COMPILE_STATUS, &status);
if (status != GL_TRUE ) {
      fputs ("Error in fragment shader compilation\n", stderr);
      exit (EXIT_FAILURE);
}
\end{verbatim}

Once we have compiled them, we create our program object, attach the shaders to it, and link the program:

\begin{verbatim}
GLuint program;

program = glCreateProgram ();

glAttachShader (program, vertShader);
glAttachShader (program, fragShader);

glLinkProgram (program);
\end{verbatim}

The \texttt{glCreateProgram} function allocates a program object and returns its handle to us. Again, it is a good idea to verify that the link operation succeeded:

\begin{verbatim}
glGetProgramiv (vertShader, GL_LINK_STATUS, &status);
if (status != GL_TRUE ) {
      fputs( "Error when linking shader program\n", stderr );
      exit (EXIT_FAILURE);
}
\end{verbatim}

Our error checking here is very rudimentary—all it reveals is \texttt{whether} an error occurred, not \texttt{what} the error was. We can get more information about what
happened by retrieving the shader or program information log. To do this, we first ask for the length of the log, and then retrieve the log into a string buffer, which we can then print out. Here is an example, using dynamically allocated buffers:

```c
GLint length;
GLsizei num;
char *log;

glGetShaderiv (vertShader, GL_INFO_LOG_LENGTH, &length);
if(length > 0) {
    log = (char *) malloc (sizeof(char) * length);
    glGetShaderInfoLog (vertShader, length, &num, log);
    fprintf (stderr, "%s\n", log);
}

glGetProgramiv (program, GL_INFO_LOG_LENGTH, &length);
if(length > 0) {
    log = (char *) malloc (sizeof(char) * length);
    glGetProgramInfoLog (program, length, &num, log);
    fprintf (stderr, "%s\n", log);
}
```

The information log functions have the same parameter list. The first parameter is the object whose log we want to retrieve. The fourth parameter is the buffer into which the log will be placed, expressed as a null-terminated string; the second parameter is the size of that buffer (so that the function will not overrun the buffer). The function will place the number of bytes written into the buffer (not including the trailing null) into the third parameter.

We can have any number of shader program objects in our OpenGL program, which allows us to apply different shaders to each object in our scene. To use a shader program, we make it the active shader before drawing the object to which it will be applied:

```c
glUseProgram (program);
```

Once we activate a shader, it will be applied to every object that we draw until we activate a different shader. If we have activated a shader for one or more objects and then want to “deactivate” it, we call `glUseProgram` again but give it a value of 0 as the program object:

```c
glUseProgram (0);
```

Finally, during execution, we may want to delete shader objects or program objects when we are done with them. The functions `glDeleteShader` and `glDeleteProgram` are used to do this. Each takes an object handle for the appropriate type of object (shader or program) as its only parameter. The memory associated with the object will be deallocated, and the object handle is marked as unused. Deleting a program object detaches the shaders associated with it but does not delete them; they are still usable, and they can be attached to another program object. We can explicitly detach a shader object from a program object with the function `glDetachShader`, which takes the program object given as its first parameter and detaches the shader object given as the second parameter from it.

If we delete a shader object before the program object it is attached to is deleted, the actual deletion is deferred until the program object is deleted.

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Similarly, if we delete a program object while it is still the active shader program, the deletion will be deferred until the shader program is no longer active.

**Basic Data Types**

The set of data types provided by GLSL is significantly larger than what is found in C-family languages. At the same time, some familiar types from those languages either don’t exist in GLSL, or exist in modified form. GLSL data types can be categorized as scalar types, vectors, matrices, and samplers. Any of these can be grouped using structure and array capabilities.

In general, variable declarations have the same form as those in C and C++, and may occur anywhere wherever needed within shader source code. Variables can be initialized at declaration time; however, the syntax for initialization varies depending on the type of variable being initialized.

Scalar types are limited to integer (int), unsigned integer (uint), Boolean (bool), and floating-point (float). Boolean variables have only two possible values, true and false. Integer and floating-point variables have the usual range of values possible in most programming languages, and most C operators are available, with the exception of bitwise operators.

**Vectors**

Vectors of each of the four scalar types are available, and can have two, three, or four components. Declarations of vec2, vec3, and vec4 contain floating-point values; a one-character prefix is added for the other scalar types (for example, ivec2, uvec2, and bvec2). The vector types can be used for any kind of data—for example, a vec4 could contain a red, green, blue, and alpha (RGBA) color value, or the x, y, z, and w components of a point, and so on. Vector initialization is done using the constructor syntax of C++; for example, we could initialize the four elements of a vec4 to the values 1.0, 2.0, 3.0, and 4.0 as follows:

```glsl
vec4 a = vec4(1.0, 2.0, 3.0, 4.0);
```

GLSL provides several mechanisms for manipulating vectors. Vector variables can be subscripted like arrays, with the first element having subscript 0, the second having subscript 1, and so on. In addition, structure-like referencing of vector elements is possible. For example, the four elements of a vec4 variable named position can be accessed with the expressions position.x, position.y, position.z, and position.w, respectively, treating the variable as a point in space. However, the same four elements can also be accessed with the names r, g, b, and a, treating it as an RGBA color, or as s, t, p, and q, treating it as a texture coordinate. The only compile-time type checking done here is verifying that the vector is large enough to contain the requested element; position.y, position.b, and position.p all access the third element of the vector.

It is also possible to access collections of vector elements using a technique called swizzling. Swizzling is a generalization of the structure-access mechanism; instead of a single element name, multiple names can be used. Here are several examples:

```glsl
vec4 v;

v.xyzw // a vec4 identical to v
v.xyz // a vec3 containing the first three elements of v
v.rgb // a vec3 containing the first three elements
v.y // a float containing the second element
v.sp // a vec2 containing the first and third elements
```
Element names can also be listed in order or out of order, or they can be duplicated—the only restriction is that they must be from the same name set: (xyzw, rgba, or stpq):

\[
\text{vec4 } a = \text{vec4}(1.0, 2.0, 3.0, 4.0); \\
\text{vec3 } b = \text{v.yzx}; \quad // \quad (2.0, 3.0, 1.0) \\
\text{vec4 } c = \text{v.rrbb}; \quad // \quad (1.0, 1.0, 3.0, 3.0)
\]

Arithmetic operators are overloaded to allow multiplication between vectors and matrices.

Matrices

Matrices of floating-point values can be declared. Square matrices (that is, \( n \times n \) elements) can be declared as \text{mat2}, \text{mat3}, and \text{mat4} variables. Non-square matrices can be declared as \text{mat} \( m \times n \) variables. Elements of matrices can be accessed using array notation.

It is possible to access an entire column at once by using a single subscript, or a single element by using two subscripts. As in OpenGL, matrices are stored in \text{column-major} order, so the first subscript is the column number and the second is the row number. For example, assuming the declaration \( \text{mat4 } m \), \( m[2] \) is a \text{vec4} containing the third column, and \( m[1][3] \) is a \text{float} containing the second element of the fourth row. Initialization is done using constructor syntax, listing the elements in column-major order:

\[
\text{mat2 } m = \text{mat2}(1.0, 2.0, 3.0, 4.0);
\]

creates the matrix

\[
m = \begin{bmatrix} 1.0 & 3.0 \\ 2.0 & 4.0 \end{bmatrix}
\] (2)

Arithmetic operators are overloaded to allow matrix manipulation.

Structures and Arrays

Structures and arrays are similar to their C counterparts. Arrays can be created from any type, including vectors, matrices, structures, and scalars. Structure members can be of any type known to the shader compiler at the time of declaration, including other structures and arrays. A structure declaration is considered automatically to be a type declaration; variables of the structure type are declared simply by using the structure name tag. For example:

\[
\text{struct lightsource} \\
\quad \text{vec3 color;} \\
\quad \text{vec3 position;} \\
\];

\text{light desklamp;}
\text{light spotlights[4];}

As mentioned earlier, GLSL is a much stricter language with respect to data types than either C or C++. Because there is a Boolean type, conditional expressions must always be Boolean, unlike C and C++ (which allow the use of any
expression whose value can be implicitly converted to integer as conditional expressions). Implicit type conversions are limited to conversions from integer or unsigned integer to float, either as scalars or as vectors. All other type conversions must be explicit; rather than using C-style type casting, conversion is requested using C++ constructor syntax.

Control Structures

GLSL provides most of the usual C control structures. Looping constructs include `for`, `while`, and `do-while` loops. Variables can be declared within loops, and the `break` and `continue` statements perform the expected operations. In the original version of GLSL, selection statements were limited to `if-then` and `if-then-else` constructs. GLSL version 1.30 introduced `switch` statements, but `goto` statements and labels are not available. Unlike C and C++, variables cannot be declared inside `if` statements.

As mentioned earlier, conditional expressions must be Booleans; no implicit conversion from numeric types to Boolean types is provided. The Boolean connectives (`&&` and `||`) are short-circuited, as in C and C++, and produce Boolean results, as do the relational operators.

A special statement, `discard`, is available for use in fragment shaders. Its purpose is to prevent the fragment shader from making any change to the frame buffer. When a `discard` is executed, the fragment being processed is marked to be discarded. The shader may or may not continue to execute, but whatever operations it performs will have no effect on the frame buffer.

GLSL Functions

Function declarations and calls are much like C++ function calls, with a few differences. Every function must be declared with an explicit return type; the return type `void` is allowed, indicating that the function does not return a value. Return types can be any type, including arrays and structures. Functions cannot be recursive in any form, including indirect recursion (that is, it is illegal for a function to call itself, or for it to call another function that then calls the first function again).

Names of functions can be overloaded based on parameter type; that is, multiple declarations of a function are allowed within a shader, so long as each declaration has the same return type and the parameter lists are all clearly distinct.

Parameter type checking is always performed. All actual parameters must match exactly the type of the corresponding formal parameter. Array parameters must have explicit sizes. A function declaration with an empty parameter list is not generic, as in C, but rather indicates that the function must be called without parameters.

Function parameters in GLSL are passed using `call by value-return` (sometimes known as `call by value-result`). Parameters are qualified as `in`, `out`, or `inout` parameters; `in` parameters can be qualified further as `const`, indicating that the formal parameter cannot be modified within the function. For `in` and `inout` parameters, the actual parameter supplied in the call is copied into the formal parameter; in the case of `out` parameters, the actual parameter supplied in the call is ignored (although the formal parameter is readable in the shader, its initial contents are undefined). If no qualifier is used, `in` is assumed. For `out` and `inout` parameters, the last value assigned to the formal parameter during the execution of the function is copied back to the original actual parameter when the
function returns. (These types of actual parameters must not be literals, but must, for obvious reasons, be actual variables.)

Arrays and structures can be passed as parameters to the functions. However, arrays are not passed by reference—instead, the contents of the array is copied into the formal parameter, as with all other parameter types.

As might be expected, a large number of built-in functions are available in GLSL. These range from angle conversions (degrees and radians), trigonometric operations, exponentiation and logarithm functions, and vector and matrix geometric operations.

Communicating with OpenGL

Because the main routine of a shader takes no parameters, communication with the rest of the OpenGL program is achieved by way of global variables. As with function parameters, global variables are typically qualified based on how they are used to convey information to the shader from the OpenGL program or between the vertex and fragment shaders. In the latter case, the same global variable will be declared in both shader sources but may have different qualifiers in the two shaders. Global variable qualifiers are similar to those for function parameters, with a few differences.

The OpenGL program uses uniform global variables to communicate data into all types of shaders. Generally, they contain data that does not change frequently. Shaders can read uniform variables, but cannot write to them.

The in qualifier is used in all types of shaders to indicate data that is being given to the shader from previous stages in the pipeline. In a vertex shader, the source is typically the OpenGL program, and the type of the variable is limited to a numeric scalar or vector (Booleans are not allowed) or a matrix.

In a fragment shader, the source of data read from an in global variable can be the OpenGL program or the vertex shader; in the latter case, the variable must exactly match an out-qualified variable in the vertex shader. Commonly, this data is interpolated—for example, there may be several fragments generated by the pipeline from a set of vertices, and each fragment will be sent through the fragment shader separately, so the contents of the in variable may vary between executions of the shader.

Values being produced by any type of shader for use in later stages in the pipeline are defined with the out qualifier. Other than the built-in global variables discussed earlier, out variables are the only way that results can be sent from the vertex shader to the fragment shader. The same variable must be declared with the same size and type as an in variable in the fragment shader.

In GLSL versions prior to 1.30, the in and out qualifiers did not exist. Global variables holding per-vertex data coming from OpenGL into a vertex shader was marked with the qualifier attribute, and output global variables had the qualifier varying. In fragment shaders, global variables of any type (coming in from the vertex shader, or going out to later stages in the pipeline) were tagged as varying. While attribute and varying qualifiers are still recognized in version 4.10.6 of GLSL (the current release as this book was written), their use should be limited, as they may be removed from future versions.

Communicating information from an OpenGL program to vertex and fragment shaders through global variables is not quite as simple as we might like. Because these variables are defined in the shader source code, they aren’t known when the OpenGL program is compiled, and thus the program cannot access them directly. Instead, the OpenGL program must first request the location of the variable in the current shader program object, and only then can it write data into the global for use by the shader. We request the location of a uniform variable in
this manner:

```c
GLint location;

location = glGetUniformLocation (program, "variable");
```

where `program` is a program object handle, and `variable` is a null-terminated string containing the name of the uniform global variable that we want to access. Once we have the location, we can retrieve the contents of the variable with one of the following functions:

```c
GLint i;
GLfloat f;

glGetUniformiv (program, location, &i);
glGetUniformfv (program, location, &f);
```

If we have the location of a uniform variable in a program object, we can modify its contents. To do this, we must know not only its location, but also its type and the number of elements that it contains:

```c
GLfloat v1, v2, v3, v4;

glUniform1f (location, v1);
glUniform2f (location, v1, v2);
glUniform3f (location, v1, v2, v3);
glUniform4f (location, v1, v2, v3, v4);
```

There are also array versions of these routines:

```c
GLfloat va[4];

glUniform1fv (location, 1, va);
glUniform2fv (location, 2, va);
glUniform3fv (location, 3, va);
glUniform4fv (location, 4, va);
```

Similarly, we use `glUniform*i` and `glUniform*iv` to write to uniform integer variables. Modification of attribute variables is handled similarly, with the functions `glGetAttribLocation`, `glVertexAttrib1fv`, `glVertexAttrib1iv`, and so on.

The `glUniform` functions do not take a program object parameter. This means that they can only write to shader variables found in the active program object (that is, the one selected by the most recent call to `glUseProgram`).

## 4 Shader Effects

Now that we have some understanding of the structure and capabilities of GLSL shaders, it is time to see some examples. Again, note that these examples are relatively simple; showing the full power of shaders is beyond the scope of this text.
A Phong Shader

Recall the Phong illumination model described in Equation 1. This can be implemented quite easily in GLSL. For simplicity, we will assume that GL_LIGHT0 has been enabled as a directional light source in the scene, and that each object has been defined with appropriate material properties.

To implement Phong shading, we need to know where our light source is. Information about active OpenGL lights is available in a built-in global variable named gl_LightSource, which is a uniform array with one element per OpenGL light source. Each element of the array is a structure containing a number of fields that describe the light. For our purposes, the most important of these are ambient, diffuse, specular, and position. These are all vec4 fields; the first three contain the ambient, diffuse, and specular characteristics of the light source, and the fourth contains the light’s position. The expression

\[ \text{gl\_LightSource[0].diffuse} \]

gives us the diffuse light emitted by GL_LIGHT0.

We also need to know what the material properties are for the object being shaded. These are available through a global variable named gl_FrontMaterial. This variable is also a structure, with ambient, diffuse, and specular fields containing these characteristics for the object. To compute the interaction of the diffuse light and the surface, we multiply these two fields:

\[ \text{gl\_FrontMaterial.diffuse} \times \text{gl\_LightSource[0].diffuse} \]

There is also a gl_BackMaterial global variable which we can use for two-sided lighting.

Our Phong implementation will be a relatively simple pair of shaders. All calculations will be done in the vertex shader, and the fragment shader will only copy the computed color into the gl_FragColor variable. We start by computing the ambient light contribution as the product of the object’s ambient reflective characteristics and the ambient illumination from the light source:

\[ \text{vec4 color; color = gl\_FrontMaterial.ambient} \times \text{gl\_LightSource[0].ambient;} \]

To compute the diffuse contribution, we will need to know the surface normal, the direction to the light, and the view direction. To use the dot-product method, all three of these vectors must normalized. The surface normal is available in the global variable gl_Normal; however, like gl_Vertex, it is in object coordinates, so we must transform it before we use it. We do this by multiplying it by the global variable gl_NormalMatrix, and we normalize the result with the built-in normalize function:

\[ \text{vec3 normal; normal = normalize( gl\_NormalMatrix \times gl\_Normal );} \]

We next need to normalize the direction to the light. Our light is directional, which means that its position in OpenGL is actually the direction the light is
shining. We can take the position, convert it to a three-element vector, and normalize the result:

```cpp
vec3 lightdir;
lightdir = normalize( vec3( gl_LightSource[0].position ) );
```

If the light was positional, we could compute its direction as the difference between the vertex position and the light position.

Computing the cosine of the angle between these vectors is easy because they are normalized vectors. To ensure that we don’t get a negative cosine, we will clamp the result to 0.0:

```cpp
float NdotL;
NdotL = max( dot(normal, lightdir), 0.0 );
```

We now have enough information to compute the diffuse contribution and add it to the final color:

```cpp
color += NdotL * (glFrontMaterial.diffuse * gl_LightSource[0].diffuse);
```

If the cosine value is positive, we also want to include the specular highlight. This requires that we compute the view vector and the reflection of the light vector. In eye coordinates, the view vector can be computed by subtracting the eye position from the vertex position. However, the eye position is the origin, so we can just use the vertex position and negate it to get the vector from the vertex to the eye position. We can use the built-in `reflect` function to compute the reflection of the light vector around the surface normal; our light vector is pointing from the light to the vertex, though, so we must negate it. Once we have those, we can compute their dot-product (clamping it to 0.0), calculate the specular contribution, and add it to the computed color:

```cpp
if( NdotL > 0.0 )
{
    vec3 view, reflection;
    float RdotV;
    view = vec3( -normalize(gl_ModelViewMatrix * gl_Vertex) );
    reflection = normalize( reflect(-lightdir, normal) );
    RdotV = max( dot(reflection, view), 0.0 );
    color += gl_FrontMaterial.specular *
             gl_LightSource[0].specular *
             pow( RdotV, gl_FrontMaterial.shininess );
}
```
Finally, we must assign the computed color to the global \( gl\_FragColor \) variable and transform the vertex. Here is the complete vertex shader:

```cpp
// Phong vertex shader
void main() {
    vec3 normal, lightdir;
    vec4 color;
    float NdotL;

    color = gl_FrontMaterial.ambient * gl_LightSource[0].ambient;

    normal = normalize(gl_NormalMatrix * gl_Normal);
    lightdir = normalize( vec3(gl_LightSource[0].position) );
    NdotL = max( dot(normal, lightdir), 0.0 );

    color += NdotL *
        (gl_FrontMaterial.diffuse * gl_LightSource[0].diffuse);

    if( NdotL > 0.0 )
        {
            vec3 view, reflection;
            float RdotV;

            view = vec3( -normalize(gl_ModelViewMatrix * gl_Vertex) );
            reflection = normalize( reflect(-lightdir, normal) );
            RdotV = max( dot( reflection, view ), 0.0 );

            color += gl_FrontMaterial.specular *
                gl_LightSource[0].specular *
                pow( RdotV, gl_FrontMaterial.shininess );
        }

    gl_FrontColor = color;
    gl_Position = ftransform();
}
```

Because we performed all the color calculations in the vertex shader, the fragment shader is very simple:

```cpp
// Phong fragment shader
void main()
{
    gl_FragColor = gl_Color;
}
```

Color Plate 32 shows a scene containing three \texttt{gluSpheres}, illuminated by a single directional light, drawn using this shader pair.
All the computation in our example was done in the vertex shader, and the fragment shader simply used that result. We could have done the color calculations in the fragment shader, but we would still need to do the normal, light direction, and view vector calculations in the vertex shader because of the need to access the per-vertex variables. The resulting vectors would be communicated to the fragment shader through global variables.

Texture Mapping
Texture mapping is another operation that is relatively easy to implement using shaders. It can be implemented by directly mapping each location on the surface of an object to a point within the texture, or by modifying the Phong shader shown earlier to take color information from the texture image rather than from the object’s material properties.

We first set up the texture within our OpenGL program by creating a texture object with glGenTextures, binding it with glBindTexture, setting our desired texture parameters with calls to glTexParameter, and then defining the texture itself with glTexImage. To use the texture within a shader, however, two additional steps are required. We must tell the shader where to find the texture, and the shader itself must gain access to the texture data.

Our discussion of surface texture mapping in OpenGL assumed that we could apply only a single texture at a time to the surface of an object. This was, in fact, a simplification of OpenGL’s texture-mapping capabilities. OpenGL actually supports multitexturing—that is, the ability to apply more than one texture to the surface of an object. It does this through the use of texture units. The number of texture units is implementation-dependent; the following example code queries the OpenGL state to determine the number of texture units in this implementation:

```glsl
GLint units;

glGetIntegerv (GL_MAX_TEXTURE_UNITS, &units);
```

When we define a texture, it is defined within the active texture unit. All texture parameter settings and image data are assigned to that unit. The active unit is selected with a call to glActiveTexture, as follows:

```glsl
glActiveTexture (GL_TEXTURE0);
```

This selects texture unit 0 as the active unit. (This is actually the default texture unit, so this function call is unnecessary unless we have selected a different unit and want to switch back to unit 0.)

After we bind our texture object to the texture unit, we must tell our shader which texture unit we are using. We do this by writing into a global sampler variable in the shader. Samplers are special types of data items in GLSL that have access to all the texture information in a texture unit. The shader code uses a sampler to identify the texture unit to be accessed, but the sampler itself is opaque to the shader. It cannot be directly read or written by GLSL code—it can only be passed as a parameter to a texture access function within our shader.

Samplers come in many forms. We create samplers for one-, two-, and three-dimensional floating-point textures with the types sampler1D, sampler2D, and sampler3D. Samplers can also be created for integer or unsigned integer textures, cube-map textures, shadow map textures, and other variations. To create a
sampler for a basic two-dimensional texture, for example, we would use a declaration like this one in our fragment shader:

```glsl
uniform sampler2D textureID;
```

In the OpenGL program, we assign the texture unit sequence number to the sampler. If our active shader program is `texShader` and we want to use texture unit 0, we find the location of the sampler variable and assign the sequence number to it:

```glsl
GLint texloc;

texloc = glGetUniformLocation (texShader, "textureID");
glUniform1i (texloc, 0);
```

Note that we assign the texture unit sequence number (0), not the OpenGL symbolic constant (`GL_TEXTURE0`), to the sampler variable.

A shader program that maps a two-dimensional texture directly to the surface of an object is very straightforward. Generally, the vertex shader takes care of setting up all necessary texture coordinates, and the fragment shader accesses the texture and uses it to determine the color of the fragment. The texture coordinates for texture unit 0 that correspond to the current vertex are available to the vertex shader in a global variable named `gl_MultiTexCoord0`. These coordinates must be communicated to the rest of the pipeline for interpolation; this is achieved by assigning them to the first slot in a global array of vectors named `gl_TexCoord`.

Here is a simple vertex shader that copies the existing texture coordinates for interpolation:

```glsl
void main()
{
    gl_TexCoord[0] = gl_MultiTexCoord0;
    gl_Position = ftransform();
}
```

The fragment shader must use the interpolated coordinates to access the texture image and determine the fragment color accordingly. Because the sampler variable is opaque, the shader must use built-in functions to access the texture data. The built-in function `texture2D` takes a sampler variable and a coordinate position as its parameters, and returns the texture data as a `vec4` value. Here is a simple fragment shader that uses the texture data directly as the fragment color:

```glsl
uniform sampler2D textureID;

void main ()
{
    vec4 color = texture2D(textureID, gl_TexCoord[0].st);
    gl_FragColor = color;
}
```
The result of using this shader pair to map a texture image of the surface of the Earth to a square polygon is shown in Color Plate 33. We can use it to texture-map any object for which texture coordinates are defined. Color Plate 34 shows an application of the same texture image to a GLU quadric sphere.

This example texture-mapping shader is very simplistic because it uses the texture color information directly as the fragment color. A more realistic result could be obtained, for instance, by modifying a Phong shader to use color information from a texture image instead of the material properties of the object.

Bump Mapping

Another application of texture mapping is the simulation of surface roughness on an object. The technique known as bump mapping uses a function to perturb the normal vector at a point on the surface of an object, and then applies a standard illumination model to calculate color at that point. Bump mapping is relatively easy to implement in an interactive program using shaders, but is significantly more difficult to implement using the original fixed-function pipeline.

To bump-map an image, we must decide how far to perturb the surface normal at each point on the object. We can do this computationally as we process each fragment, or we can precompute the changes to be applied at each point and hold them in a special type of texture called a normal map. If we are also applying a texture to the surface of the object as well as bump-mapping the surface, an obvious source of bump-map information is the color variation within the image.

One complication in bump mapping is the fact that we must work with several different coordinate spaces. The incoming information that we use to compute surface colors is typically in either object coordinates or eye coordinates; however, we must do our displacement calculations in texture space. Typically, we resolve this issue by converting everything into texture space. To do this, we compute the partial derivative vector \( P_u' \). We then normalize this vector and the surface normal and take their cross-product, which produces third vector that is orthogonal to the first two. These three vectors are used to perform the transformation.

The normalized \( P_u' \) vector is called the tangent vector. The cross-product of the tangent vector and the surface normal is called the binormal vector. From the components of these vectors, we build the transformation matrix

\[
M = \begin{bmatrix} T_x & T_y & T_z \\ B_x & B_y & B_z \\ N_x & N_y & N_z \end{bmatrix}
\]

where \( (T_x, T_y, T_z) \) is the tangent vector, \( (B_x, B_y, B_z) \) is the binormal vector, and \( (N_x, N_y, N_z) \) is the normal vector. Multiplying an object-space vector by this matrix transforms it into tangent space. Tangent space is a local coordinate system around the point being shaded, and the tangent and binormal vectors may change from point to point across the surface.

Our bump-mapping shader will be a simplified form of a technique known as relief mapping. We will use the texture image to determine the surface color at the
shading point, as in our earlier texture-mapping example. However, we will also use it to calculate the displacements to be applied to our surface normals—that is, the apparent roughness of the surface will be determined by the color variations in our texture image.

To implement bump mapping as a shader pair, we must divide the work between the vertex and fragment shaders. In addition to the transformations we have seen in earlier examples, the vertex shader will compute the light and view vectors and will transform them into tangent space for use by the fragment shader. The fragment shader, in turn, will use the color information from the texture image to calculate the height variation for this fragment, and it also will perform a simple diffuse shading calculation using the texture color information.

To compute the transformation matrix in our vertex shader, we need the tangent and binormal vectors in addition to the surface normal. We are given the surface normal; we can either compute a tangent vector from it or use one supplied by the OpenGL program. We can calculate the tangent vector fairly easily by computing the cross-products between the surface normal and the $y$ and $z$ axes, and then selecting the longer of the two cross-products and normalizing it.

If we choose to have the OpenGL program supply the tangent vector, it must be written into a global attribute variable used by the vertex shader. We obtain its location using `glGetAttribLocation` and write three values into it as follows:

```c
GLfloat tangVector[3];
GLint tangentLoc;

tangentLoc = glGetAttribLocation (bumpshader, "tangent");
glVertexAttrib3fv (tangentLoc, tangent);
```

In the vertex shader, `tangent` is declared globally as an attribute variable. We must also declare the view and light vector variables as varying variables:

```c
attribute vec3 tangent;
varying vec3 light, view;
```

In the vertex shader, we transform the surface normal, and compute the binormal vector: Once we have these three vectors, we can compute the view and light vectors, and transform them into tangent space. A fast way to perform the transformation is to take advantage of the fact that we can compute the three result values from multiplying the transformation matrix by a vector using the built-in dot product function. For example, given the transformed light vector and the three tangent-space vectors, we can transform the light vector into tangent space as follows:

```c
vec3 tmp;
tmp.x = dot( light, tangent );
tmp.y = dot( light, binorm );
tmp.z = dot( light, normal );
light = tmp;
```
We complete the vertex shader by performing the usual copying of the texture location and transformation of the vertex position into clip space. Here is the completed vertex shader:

```glsl
varying vec3 light, view;
attribute vec3 tangent;

void main()
{
  vec3 normal = vec3( normalize( gl_NormalMatrix * gl_Normal ) );
  vec3 binorm = normalize( cross( normal, tangent ) );
  view = -normalize( vec3( gl_ModelViewMatrix * gl_Vertex ) );
  light = normalize( vec3( gl_LightSource[0].position ) );
  vec3 tmp;
  tmp.x = dot( light, tangent );
  tmp.y = dot( light, binorm );
  tmp.z = dot( light, normal );
  light = tmp;
  tmp.x = dot( view, tangent );
  tmp.y = dot( view, binorm );
  tmp.z = dot( view, normal );
  view = tmp;
  gl_TexCoord[0] = gl_MultiTexCoord0;
  gl_Position = ftransform();
}
```

The bump-mapping fragment shader is more complicated than our previous fragment shaders. Globally, we need to define the light and view vector variables, along with our texture sampler variable.

To compute the height offset from a color value, we will compute the average of the red and green components and then smooth out variations from point to point by taking 1.5 percent of the color average and adding that to 98.5 percent of a 50 percent gray value. We compute the blended value using the built-in `mix` function:

```glsl
float height( vec3 color )
{
  float avg = (color.r + color.g) / 2.0;
  return mix( avg, 0.5, 0.985 );
}
```

Creating the perturbed surface normal at a point on the texture is achieved by first creating a small triangle around the point on the surface. We calculate the positions of the triangle’s vertices by adding three different offsets to the texture coordinate to locate three points at 0°, 120°, and 240° around an imaginary circle centered on the coordinate point. For each of these vertices, we create a vector consisting of the s and t offsets and a height offset calculated from the color at the vertex. We then create the normal vector from these three vectors by computing the cross-product of vectors formed by taking the difference between pairs of the vectors.

The rest of our fragment shader is straightforward. We compute the modified normal vector for the texture coordinate, and then compute ambient and diffuse

---

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color contributions based on the texture color and the modified normal. Here is
the completed fragment shader:

```glsl
varying vec3 light, view;
uniform sampler2D textureID;

// Calculate height offset
float height( vec3 color ) {
    float avg = (color.r + color.g)/2.0;
    return mix( avg, .5, .985 );
}

// Create modified surface normal
vec3 modNormal( vec2 point ) {

    // Create the small triangle - first, the s and t
    // distances from the center point
    vec2 d0 = vec2( 0, 0.001 );
    vec2 d1 = vec2( -0.000866, -0.0005 );
    vec2 d2 = vec2( 0.000866, -0.0005 );

    // Calculate the triangle vertex positions
    vec2 p0 = point + d0;
    vec2 p1 = point + d1;
    vec2 p2 = point + d2;

    // Compute the height offset for each vertex
    float h0 = height( vec3( texture2D( textureID, p0 ) ) );
    float h1 = height( vec3( texture2D( textureID, p1 ) ) );
    float h2 = height( vec3( texture2D( textureID, p2 ) ) );

    // Create the three vectors
    vec3 v0 = vec3( d0, h0 );
    vec3 v1 = vec3( d1, h0 );
    vec3 v2 = vec3( d2, h0 );

    // Compute the modified normal vector
    return normalize( vec3( cross( v1-v0, v2-v0 ) ) );
}

void main() {
    vec4 base = texture2D( textureID, gl_TexCoord[0].st );
    vec3 bump = modNormal( gl_TexCoord[0].st );
    vec4 color = gl_LightSource[0].ambient * base;

    float NdotL = max( dot(bump, light), 0.0 );
    color += NdotL * ( gl_LightSource[0].diffuse * base );

    gl_FragColor = color;
}
```

Color Plate 35 shows the results of using this shader pair to apply the Earth
surface texture image used in previous examples to a square polygon. Compare
5 Summary

Computer graphics libraries have evolved over time to match the capabilities of graphics hardware. In the beginning, graphics programmers were required to work directly with the hardware available to them. Libraries of commonly used graphics routines were developed in an attempt to standardize the development of graphics programs, culminating in APIs such as the OpenGL library and its original fixed-function internal pipeline.

As graphics hardware continued to evolve, the fixed-function pipeline became more limiting because it could not take advantage of improved hardware capabilities. To solve this problem, a programmable pipeline model was developed, which allowed graphics programmers more control over the functionality of different stages in the pipeline through the use of programmable shaders. Shading languages were created to simplify the task of performing common shading operations.

The OpenGL Shading Language (GLSL) was developed as a way of integrating programmable shading operations into the OpenGL pipeline. GLSL provides programmable "hooks" into the pipeline at critical stages, allowing the manipulation of vertices, object geometries, surface tessellation, and fragment manipulation through the use of shader programs. Given the flexibility of GLSL, it is possible to perform easily shading tasks that would be difficult or impossible to accomplish using a fixed-function graphics pipeline. Table 1 lists the OpenGL functions used to create and communicate with GLSL shader programs.

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>glCreateShader</td>
<td>Creates a shader object.</td>
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<td>glShaderSource</td>
<td>Attaches shader source code to a shader object.</td>
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<td>glCompileShader</td>
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<td>glGetShaderiv</td>
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<td>glCreateProgram</td>
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<td>glAttachShader</td>
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<td>glGetProgramInfoLog</td>
<td>Retrieves shader program messages.</td>
</tr>
<tr>
<td>glUseProgram</td>
<td>Activates a shader program.</td>
</tr>
<tr>
<td>glGetUniformLocation</td>
<td>Obtains the location of a global shader uniform variable.</td>
</tr>
<tr>
<td>glGetUniform*</td>
<td>Reads the contents of a global shader uniform variable.</td>
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<td>glUniform*</td>
<td>Writes the contents of a global shader uniform variable.</td>
</tr>
<tr>
<td>glGetAttribLocation</td>
<td>Obtains the location of a vertex shader attribute variable.</td>
</tr>
<tr>
<td>glGetAttrib*</td>
<td>Reads the contents of a vertex shader attribute variable.</td>
</tr>
<tr>
<td>glVertexAttrib*</td>
<td>Writes the contents of a vertex shader attribute variable.</td>
</tr>
</tbody>
</table>
REFERENCEs
Shade trees are discussed in Cook (1984). Ken Perlin’s PSE is described in Perlin (1985), and his original noise implementation can be found on his web-site, at http://cs.nyu.edu/~perlin/. RenderMan is presented in Upstill (1989) and Apodaca and Gritz (2000). The official RISpec can be found online at https://renderman.pixar.com/products/rispec/index.htm, and a number of RenderMan implementations (both commercial and open-source) can be found on the Internet. Relief texture mapping is discussed in Oliveira, Bishop and McAllister (2000) and in Policarpo, Oliveira, and Comba (2005).
Official specifications for GLSL can be found at http://www.opengl.org/, along with sample programs and tutorial guides. GLSL is also discussed in Shreiner (2010). Finally, more complete treatments of GLSL can be found in Rost and Licea-Kane (2010) and in Bailey and Cunningham (2009).

EXERCISEs
1. Determine whether or not your OpenGL installation supports GLSL. If it does, determine the version of GLSL it supports.
2. Write a function which takes two null-terminated strings as its parameters and returns the GLuint identifier for a shader program object. The parameters contain the names of vertex and fragment shader source files.
3. Write a program using the functions in the previous exercise to draw a square in the center of the display window and use the vertex shader program to color the square red.
4. The example Phong shader performs its color calculations at each vertex. Convert it into a shader that does the color calculations in the fragment shader. Remember that some critical values must be computed in the vertex shader.
5. Write a program to display an origin-centered tetrahedron on a black background using the shader you wrote in the previous exercise to shade the object. Add the ability to rotate the object around the y-axis using keyboard input.
6. Modify the example Phong shader to use a light source specified through a global shader variable rather than just using light source 0.
7. Modify the example Phong shader to work with multiple light sources. Use a global shader variable to tell the shader how many light sources are active.
8. Modify the program in Exercise 5 to add two more light sources to the scene. Use the shader you developed in the previous exercise to shade the object in the scene. The positions and orientations of the light sources should be taken as input parameters to the program.
9. Modify the simple texture mapping shader so that it performs Phong calculations using color information from the texture image instead of the material properties of the object being shaded.
10. Write program to display an origin-centered cube on a black background using the shader you wrote in the previous exercise to shade the object with a textured image on each of the faces of the cube. Provide the ability to rotate the cube about the y and z axes using keyboard input.
11. Modify the program and shader used in the previous exercise to add two more light sources to the scene and have the shader texture the object using lighting information from all three lights. The positions and orientations of the light sources should be taken as input parameters to the program.

IN MORE DEPTH
1. By modifying the examples in the chapter, write a shader program to apply texture patterns to the objects in your scene. If you wrote a bump map for any of those objects, modify the example in the chapter to produce a shader program for that as well. Replace the existing texture and bump maps with the shader programs and note the visual differences between the two approaches to texture mapping, if any.
2. Consider the pattern created by four ceramic tiles arranged in a 2 × 2 pattern on a floor or wall. Each tile has a width and height, and the grout line between adjacent tiles also has a width. The total width of this pattern is 2 × tileWidth + 2 × groutWidth, and similarly the total height is 2 × tileHeight + 2 × groutWidth. We can apply this pattern to a surface as a procedural texture map without using an actual texture image fairly easily. The s texture coordinate specifies a position between the left edge (s = 0) and the right edge (s = 1); similarly, the t coordinate specifies a position between the bottom edge (t = 0) and the top edge (t = 1). Because we know the width and height of each tile and the width of the grout line, we can use the s and t coordinates to determine whether this point on the “texture” is covered by tile or grout. Create a shader that applies this type of procedural texture to an object. Communicate the width, height, and color of a tile and the width and color of the grout line to your shader through global uniform variables.
Color Plate 32
An example of Phong shading implemented as a GLSL shader. Each sphere has its own material properties; the scene is illuminated by a directional white light source.

Color Plate 33
An image of the surface of the Earth texture-mapped onto a square. (Earth image courtesy of James Hastings-Trew.)
**Color Plate 34**
The image of the Earth mapped onto a sphere. (Earth image courtesy of James Hastings-Trew.)

**Color Plate 35**
A bump-mapped version of the Earth image, using color information from the image to control the displacement of the surface normals. (Earth image courtesy of James Hastings-Trew.)
Solid object modeling is a broad topic. In earlier chapters, we discussed the use of boundary representations and physically based methods to represent solid objects. These techniques are fine for modeling objects that have regular shapes and smooth sides. However, many real-world objects have rough surfaces or irregular shapes, which are difficult to model using the techniques we have already studied. Algorithmic modeling techniques provide a conceptually straightforward way to represent these types of objects.
1 Fractal-Geometry Methods

Virtually all the object representations that we have considered in previous chapters used Euclidean-geometry methods; that is, object shapes were described with equations. These methods are adequate for describing manufactured objects: those that have smooth surfaces and regular shapes. However, natural objects, such as mountains and clouds, have irregular or fragmented features, and Euclidean methods do not provide realistic representations for such objects. Natural objects can be realistically described with fractal-geometry methods, where procedures rather than equations are used to model objects. As we might expect, procedurally defined objects have characteristics quite different from objects described with equations. Fractal-geometry representations for objects are commonly applied in many fields to describe and explain the features of natural phenomena. In computer graphics, we use fractal methods to generate displays of natural objects and visualizations of various mathematical and physical systems.

A fractal object has two basic characteristics: infinite detail at every point, and a certain self-similarity between the object parts and the overall features of the object. The self-similarity properties of an object can take different forms, depending on the representation we choose for the fractal. We describe a fractal object with a procedure that specifies a repeated operation for producing the detail in the object subparts. Natural objects are represented with procedures that theoretically repeat an infinite number of times. Graphics displays of natural objects are, of course, generated with a finite number of steps.

If we zoom in on a continuous Euclidean shape, no matter how complicated, we can eventually get the zoomed-in view to smooth out. However, if we zoom in on a fractal object, we continue to see more and more details in the magnifications without an eventual smoothing of the object appearance. A mountain outlined against the sky continues to have the same jagged shape as we view it from a closer and closer position (Figure 1). As we near the mountain, the smaller detail in the individual ledges and boulders becomes apparent. Moving even closer, we see the outlines of rocks, then stones, and then grains of sand. At each step, the outline reveals more twists and turns. If we took the grains of sand and put them under a microscope, we would again see the same detail repeated down through the molecular level. Similar shapes describe coastlines and the edges of plants and clouds.

To obtain a magnified view of a displayed fractal, we can select a section of the fractal for display within a viewing area of the same size. We then carry out the fractal-construction operations for that part of the object and display the increased detail for that level of magnification. As we repeat this process, we continue to display more and more of the object’s detail. Because of the infinite detail inherent in the construction procedures, a fractal object has no definite size. When we
include more of the detail in an object description, the dimensions increase without limit, but the coordinate extents for the object remain bound within a finite region of space.

We can characterize the amount of variation in object detail with a number called the fractal dimension. Unlike Euclidean dimension, this number is not necessarily an integer. The fractal dimension for an object is sometimes referred to as the fractional dimension, which is the basis for the name “fractal.”

Fractal methods have proven useful for modeling a very wide variety of natural phenomena. In graphics applications, fractal representations are used to model terrain, clouds, water, trees and other plant life, feathers, fur, and various surface textures, and sometimes just to make pretty patterns. In other disciplines, fractal patterns have been found in the distribution of stars, river islands, and moon craters; in rain-field configurations; in stock market variations; in music; in traffic flow; in urban property utilization; and in the boundaries of convergence regions for numerical-analysis techniques.

Fractal Generation Procedures
A fractal object is generated by repeatedly applying a specified transformation function to points within a region of space. If \( P_0 = (x_0, y_0, z_0) \) is a selected initial position, each iteration of a transformation function \( F \) generates successive levels of detail with the calculations

\[
P_1 = F(P_0), \quad P_2 = F(P_1), \quad P_3 = F(P_2), \quad \cdots
\]

In general, the transformation function can be applied to a specified point set or to an initial set of primitives, such as straight lines, curves, color areas, or surfaces. Also, we can use either deterministic or random procedures. The transformation function could be defined in terms of geometric transformations (scaling, translation, rotation), or it could involve nonlinear coordinate transformations and statistical decision parameters.

Although fractal objects, by definition, contain infinite detail, we apply the transformation function a finite number of times, and, of course, the objects that we display have finite dimensions. A procedural representation approaches a "true" fractal as the number of transformations is increased to produce more and more detail. The amount of detail included in the final graphical display of an object depends on the number of iterations performed and the resolution of the display system. We cannot display detail variations that are smaller than the size of a pixel. However, we can repeatedly zoom in on selected portions of an object to view more of its detail.

Classification of Fractals

**Self-similar** fractals have parts that are scaled-down versions of the entire object. Starting with an initial shape, we construct the object subparts by apply a scaling parameter \( s \) to the overall shape. We can use the same scaling factor \( s \) for all subparts, or we can use different scaling factors for different scaled-down parts of the object. If we also apply random variations to the scaled-down subparts, the fractal is said to be statistically self-similar. The parts then have the same statistical properties. Statistically self-similar fractals are commonly used to model trees, shrubs, and other vegetation.

**Self-affine** fractals have parts that are formed with different scaling parameters, \( s_x, s_y, \) and \( s_z \), in different coordinate directions. We can also include random variations to obtain statistically self-affine fractals. Terrain, water, and clouds are typically modeled with statistically self-affine fractal construction methods.
**Invariant fractal sets** are formed with nonlinear transformations. This class of fractals includes self-squaring fractals, such as the Mandelbrot set (formed with squaring functions in complex space), and self-inverse fractals, constructed with inversion procedures.

**Fractal Dimension**

The amount of variation in the structure of a fractal object can be described with a number \( D \), called the fractal dimension, which is a measure of the roughness, or fragmentation, of the object. More jagged-looking objects have larger fractal dimensions. One method for generating a fractal object is to set up an iterative procedure that uses a selected value for \( D \). Another approach is to determine the fractal dimension from the desired properties of an object, although, in general, the fractal dimension can be difficult to calculate. Methods for calculating \( D \) are based on dimension concepts developed in branches of mathematics, particularly topology.

An expression for the fractal dimension of a self-similar fractal constructed with a single scalar factor \( s \) is obtained by analogy with the subdivision of a Euclidean object. Figure 2 shows the relationships between the scaling factor \( s \) and the number of subparts \( n \) for subdivision of a unit straight-line segment, a unit square, and a unit cube. With \( s = \frac{1}{2} \), the unit line segment [Figure 2(a)] is divided into two equal-length subparts. For the same scaling factor, the square in Figure 2(b) is divided into four equal-area subparts, and the cube [Figure 2(c)] is divided into eight equal-volume subparts. For each of these

\[
\begin{align*}
L & \quad \Rightarrow \quad L' = \frac{L}{n} \\
D_E & = 1, \quad s = \frac{1}{n}, \quad n = 2 \\
& \quad n s^1 = 1 \\
& \quad \text{(a)}
\end{align*}
\]

\[
\begin{align*}
A & \quad \Rightarrow \quad A' = \frac{A}{n} \\
D_E & = 2, \quad s = \frac{1}{n^{1/2}}, \quad n = 4 \\
& \quad n s^2 = 1 \\
& \quad \text{(b)}
\end{align*}
\]

\[
\begin{align*}
V & \quad \Rightarrow \quad V' = \frac{V}{n} \\
D_E & = 3, \quad s = \frac{1}{n^{1/3}}, \quad n = 8 \\
& \quad n s^3 = 1 \\
& \quad \text{(c)}
\end{align*}
\]

**Figure 2**

Subdividing a unit line (a), a unit square (b), and a unit cube (c). The Euclidean dimension is represented as \( D_E \), and the scaling factor for each object is \( s = \frac{1}{2} \).
objects, the relationship between the number of subparts and the scaling factor is \( n \cdot s^{DE} = 1 \). In analogy with Euclidean objects, the fractal dimension \( D \) for self-similar objects can be obtained from

\[
ns^D = 1 \tag{2}
\]

Solving this expression for \( D \), the **fractal similarity dimension**, we have

\[
D = \frac{\ln n}{\ln(1/s)} \tag{3}
\]

For a self-similar fractal constructed with different scaling factors for the different subparts of the object, the fractal similarity dimension is obtained from the implicit relationship

\[
\sum_{k=1}^{n} s_k^D = 1 \tag{4}
\]

where \( s_k \) is the scaling factor for subpart \( k \).

In Figure 2, we considered the subdivision of simple shapes (straight line, rectangle, and cube). If we have more complicated shapes, including curved lines and objects with nonplanar surfaces, determining the structure and properties of the subparts is more difficult. For general object shapes, we can use **topological covering methods** that approximate object subparts with simple shapes. A subdivided curve, for example, could be approximated with straight-line sections, and a subdivided spline surface could be approximated with small squares or rectangles. Other covering shapes, such as circles, spheres, and cylinders, can also be used to approximate the features of an object divided into a number of smaller parts. Covering methods are commonly used in mathematics to determine geometric properties, such as length, area, or volume, of a complex object by summing the properties of a set of smaller covering objects. We can also use covering methods to determine the fractal dimension \( D \) of some objects.

Topological covering concepts were originally used to extend the meaning of geometric properties to nonstandard shapes. An extension of covering methods using circles or spheres led to the notion of a **Hausdorff-Besicovitch dimension**, or fractional dimension. The Hausdorff-Besicovitch dimension can be used as the fractal dimension of some objects, but in general, it is difficult to evaluate. More commonly, an object’s fractal dimension is estimated with **box-covering methods** using rectangles or parallelepipeds. Figure 3 illustrates the notion of a box covering. Here, the area inside the large irregular boundary can be approximated by the sum of the areas of the small covering rectangles.

Box-covering methods are applied by first determining the coordinate extents of an object, then subdividing the object into a number of small boxes using the given scaling factors. The number of boxes \( n \) that it takes to cover an object is called the **box dimension**, and \( n \) is related to the fractal dimension \( D \). For statistically self-similar objects with a single scaling factor \( s \), we can cover the object with squares or cubes. We then count the number \( n \) of covering boxes and use Equation 3 to estimate the fractal dimension. For self-affine objects, we cover the object with rectangular boxes because different directions are scaled differently. In this case, we estimate the fractal dimension using both the number of boxes \( n \) and the affine transformation parameters.

The fractal dimension of an object is always greater than the corresponding Euclidean dimension (or topological dimension), which is simply the least number of parameters needed to specify the object. A Euclidean curve is
one-dimensional because we can determine coordinate positions with one parameter, \( u \). A Euclidean surface is two-dimensional, with surface parameters \( u \) and \( v \). Finally, a Euclidean solid, which requires three parameters for each coordinate specification, is three-dimensional.

For a fractal curve that lies completely within a two-dimensional plane, the fractal dimension \( D \) is greater than 1 (the Euclidean dimension of a curve). The closer \( D \) is to 1, the smoother the fractal curve. If \( D = 2 \), we have a Peano curve; that is, the “curve” completely fills a finite region of two-dimensional space. For \( 2 < D < 3 \), the curve self-intersects and the area could be covered an infinite number of times. Fractal curves can be used to model natural object boundaries, such as shorelines.

Spatial fractal curves (those that do not lie completely within a single plane) also have fractal dimension \( D \) greater than 1, but \( D \) can be greater than 2 without self-intersecting. A curve that fills a volume of space has dimension \( D = 3 \), and a self-intersecting space curve has fractal dimension \( D > 3 \).

Fractal surfaces typically have a dimension within the range \( 2 < D \leq 3 \). If \( D = 3 \), the “surface” fills a volume of space. If \( D > 3 \), there is an overlapping coverage of the volume. Terrain, clouds, and water are typically modeled with fractal surfaces.

The dimension of a fractal solid is usually in the range \( 3 < D \leq 4 \). Again, if \( D > 4 \), we have a self-overlapping object. Fractal solids can be used, for example, to model cloud properties such as water vapor density or temperature within a region of space.

**Geometric Construction of Deterministic Self-Similar Fractals**

To construct a deterministic (nonrandom) self-similar fractal geometrically, we start with a given geometric shape, called the initiator. Subparts of the initiator are then replaced with a pattern, called the generator.

For example, if we use the initiator and generator shown in Figure 4, we can construct the snowflake pattern, or Koch curve, shown in Figure 5. Each straight-line segment in the initiator is replaced with the generator pattern, consisting of four equal-length line segments. Then the generator is scaled and applied to the line segments of the modified initiator, and this process is repeated for some number of steps. The scaling factor at each step is \( \frac{1}{3} \), so the fractal dimension is \( D = \frac{\ln 4}{\ln 3} \approx 1.2619 \). Also, the length of each line segment in the initiator increases by a factor of \( \frac{1}{3} \) at each step, so that the length of the fractal curve tends to infinity as more detail is added to the curve (Figure 6). Figure 7 illustrates additional generator patterns that could be used for self-similar fractal curve constructions. The generators in Figure 7(b) and (c) contain more detail than the Koch curve generator, and they have higher fractal dimensions.
**FIGURE 5**
First three iterations in the generation of the Koch curve.

**FIGURE 6**
The length of each side of the Koch curve increases by a factor of \( \frac{4}{3} \) at each step, while the line segment lengths are reduced by a factor of \( \frac{1}{3} \).

**FIGURE 7**
Generators for self-similar fractal curve constructions and their associated fractal dimensions.
We can also use generators with multiple disjoint components. Some examples of compound generators are shown in Figure 8. We could combine these patterns with random variations to model various natural objects that have multiple unconnected parts, such as island distributions along a coastline.

The generator of Figure 9 contains line segments with varying lengths, and multiple scaling factors are used in the construction of the fractal curve. Thus, the fractal dimension of the generated curve is determined from Equation 4.

Displays of trees and other plants can be constructed with self-similar geometric-construction methods. Each branch of the fern outline shown in Figure 10(a) is a scaled version of the overall fern shape. In (b) of this figure, the fern is fully rendered with a twist applied to each branch.

As an example of a self-similar fractal construction for the surfaces of a three-dimensional object, we scale the regular tetrahedron shown in Figure 11 by a factor of $\frac{1}{2}$, then place the scaled object on each of the original four surfaces of the tetrahedron. Each face of the original tetrahedron is converted into six smaller...
faces and the original face area is increased by a factor of \( \frac{3}{2} \). The fractal dimension of this surface is

\[ D = \frac{\ln 6}{\ln 2} \approx 2.58496 \]

which indicates a fairly fragmented surface.

Another way to create self-similar fractal objects is to punch holes in a given initiator, instead of adding more surface area. Figure 12 shows some examples of fractal objects created in this way.

Geometric Construction of Statistically Self-Similar Fractals

To introduce variability into the geometric construction of a self-similar fractal, we could randomly select a generator at each step from a menu of patterns. Or we could construct a self-similar fractal by computing coordinate displacements with small random variations. For example, in Figure 13, we use a probability distribution function to compute variable midpoint displacements at each step in the creation of a random snowflake pattern.

Another example of this method is shown in Figure 14. Random scaling parameters and branching directions are used in this display to model the vein patterns in a leaf.

Once a fractal object has been created, we can model a scene using several transformed instances of the object. Color Plate 36 illustrates instancing with random rotations of a fractal tree. In Color Plate 37, a fractal forest is displayed using various random transformations.

To model the gnarled and contorted shapes of some trees, we can apply twisting functions as well as scaling to create the random, self-similar branches.
Affine Fractal-Construction Methods

We can obtain highly realistic representations for terrain and other natural objects using affine fractal methods that model object features as fractional Brownian motion. This is an extension of standard Brownian motion, a form of “random walk,” which describes the erratic, zigzag movement of particles in a gas or fluid. Figure 15 illustrates a random-walk path in the $xy$ plane. Starting from a given position, we generate a straight-line segment in a random direction and with a random length. Another random line is then constructed from the endpoint of this first line, and the process is repeated for a designated number of line segments. Fractional Brownian motion is obtained by adding an additional parameter to the statistical distribution describing Brownian motion. This additional parameter sets the fractal dimension for the “motion” path.

A single fractional Brownian path can be used to model a fractal curve. With a two-dimensional array of random fractional Brownian elevations over a ground-plane grid, we can model the surface of a mountain by connecting the elevations to form a set of polygon patches. If random elevations are generated on the surface of a sphere, we can model the mountains, valleys, and oceans of a planet. In Figure 16, Brownian motion was used to create the elevation variations on the planet surface. The elevations were then color-coded from lowest (the oceans) to highest (snow on the mountains). Fractional Brownian motion was used to create the terrain features in the foreground. Craters were created with random diameters and random positions, using affine fractal procedures that closely describe the distribution of observed craters, river islands, rain patterns, and other similar systems of objects.

By adjusting the fractal dimension in the fractional Brownian-motion calculations, we can vary the ruggedness of terrain features. Values for the fractal dimension in the neighborhood of $D \approx 2.15$ produce realistic mountain features, while higher values close to 3.0 can be used to create unusual-looking extraterrestrial landscapes. We can also scale the calculated elevations to deepen the valleys and increase the height of mountain peaks. Some examples of terrain features that can be modeled with fractal procedures are given in Color Plate 38.

Random Midpoint-Displacement Methods

Fractional Brownian-motion calculations are time-consuming because the elevation coordinates of the terrain above a ground plane are calculated with Fourier series, which are sums of sine and cosine terms. Fast Fourier transform (FFT)
methods are typically used, but it is still a slow process to generate fractal-mountain scenes. Therefore, faster **random midpoint-displacement methods**, similar to the random displacement methods used in geometric constructions, have been developed to approximate fractional Brownian-motion representations for terrain and other natural phenomena. These methods were originally used to generate animation frames for science-fiction films involving unusual terrain and planet features. Midpoint-displacement methods are now commonly used in many other computer-graphics applications, including animations for television advertising.

Although random midpoint-displacement methods are faster than fractional Brownian-motion calculations, they produce less realistic-looking terrain features. Figure 17 illustrates the midpoint-displacement method for generating a random-walk path in the $xy$ plane. Starting with a straight-line segment, we calculate a displaced $y$ value for the midposition of the line as the average of the endpoint $y$ values plus a random offset:

$$y_{\text{mid}} = \frac{1}{2} [y(a) + y(b)] + r \quad (5)$$

To approximate fractional Brownian motion, we choose a value for $r$ from a Gaussian distribution with a mean of 0 and a variance proportional to $|(b - a)|^{2H}$, where $H = 2 - D$ and $D > 1$ is the fractal dimension. Another way to obtain a random offset is to take $r = sr_g|b - a|$, with parameter $s$ as a selected surface “roughness” factor and $r_g$ as a Gaussian random value with mean 0 and variance 1. Table lookups can be used to obtain the Gaussian values. The process is then repeated by calculating a displaced $y$ value for the midposition of each half of the subdivided line. We continue the subdivision to obtain a certain number of segments or until the lengths of the subdivided line sections are less than some selected length. At each step, the value of the random variable $r$ decreases because it is proportional to the width $|b - a|$ of the line section to be subdivided. Figure 18 shows a fractal curve obtained with this method.

Terrain features are generated by applying the random midpoint-displacement procedures to a rectangular ground plane (Figure 19). We begin by assigning an elevation $z$ value to each of the four corners ($a$, $b$, $c$, $d$ in Figure 19) of the ground plane. Then we divide the ground plane at the midpoint of each edge to obtain the five new grid positions: $e$, $f$, $g$, $h$, and $m$. 

**FIGURE 17**
Random midpoint displacement of a straight-line segment.

**FIGURE 18**
A random-walk path generated from a straight-line segment with four iterations of the random midpoint-displacement procedure.

**FIGURE 19**
A rectangular ground plane (a) is subdivided into four equal grid sections (b) for the first step in a random midpoint-displacement procedure to calculate terrain elevations.
Elevations at the midpositions e, f, g, and h of the ground-plane edges can be calculated as the average elevation of the nearest two vertices plus a random offset. For example, elevation $z_e$ at midposition e is calculated using vertices a and b, while elevation at midposition f is calculated using vertices b and c:

$$z_e = (z_a + z_b)/2 + r_e, \quad z_f = (z_b + z_c)/2 + r_f$$

Random values $r_e$ and $r_f$ can be obtained from a Gaussian distribution with mean 0 and variance proportional to the grid separation raised to the $2H$ power, with $H = 3 - D$ and $D > 2$. Higher values for $D$, the surface fractal dimension, produce more jagged terrain, while lower values generate smoother terrain features. We could also calculate random offsets as the product of a surface roughness factor times the grid separation times a table lookup value for a Gaussian value with mean 0 and variance 1. The elevation $z_m$ of the ground plane midposition m can be calculated using positions e and g or positions f and h. Alternatively, we could calculate $z_m$ using the assigned elevations of the four ground plane corners and a random offset as

$$z_m = (z_a + z_b + z_c + z_d)/4 + r_m$$

This process is repeated for each of the four new grid sections at each step until the grid separation becomes smaller than a selected value.

Triangular surface patches for the terrain surface can be formed as the elevations are generated. Figure 20 shows eight surface patches that could be constructed at the first subdivision step. At each level of recursion, the triangles are successively subdivided into smaller planar patches. When the subdivision process is completed, the patches are rendered using the positions selected for the light sources, the values of other illumination parameters, and the chosen colors and surface textures for the terrain.

The random midpoint-displacement method can be applied to generate other components of a scene besides the terrain. For instance, we could use the same methods to obtain surface features for water waves or cloud patterns above a ground plane.

**Controlling Terrain Topography**

One way to control the placement of peaks and valleys in a fractal-terrain scene that is modeled with a midpoint-displacement method is to constrain the calculated elevations to certain intervals over the various sections of the ground plane. We can accomplish this by designating a set of control surfaces over the ground plane, as illustrated in Figure 21. Then we calculate a random elevation at each midpoint grid position on the ground plane that depends on the difference between the control elevation and the average elevation calculated for that position. This procedure constrains elevations to be within a preset interval about the control-surface elevations.

Control surfaces can be used to model existing terrain features in the Rocky Mountains, or some other region, by constructing the plane facets using the...
elevations in a contour plot for a particular region. Alternatively, we could set the elevations for the vertices of the control polygons to design our own terrain features. Also, control surfaces can have any shape. Planes are easiest to deal with, but we could use spherical surfaces or other curve shapes.

We use the random midpoint-displacement method to calculate grid elevations, but now we select random values from a Gaussian distribution where the mean $\mu$ and standard deviation $\sigma$ are functions of the control elevations. A method for obtaining the values for $\mu$ and $\sigma$ is to make them both proportional to the difference between the calculated average elevation and the predefined control elevation at each grid position. For example, for grid position $e$ in Figure 19, we set the mean and standard deviation as

$$\mu_e = zc_e - (z_a + z_b)/2, \quad \sigma_e = s|\mu_e|$$

where $zc_e$ is the control elevation for ground-plane position $e$, and $0 < s < 1$ is the scaling factor. Small values for $s$, such as $s < 0.1$, produce tighter conformity to the terrain envelope, and larger values for $s$ allow greater fluctuations in terrain height.

To determine the values for the control elevations over a control-surface plane, we first determine the values for the plane parameters $A$, $B$, $C$, and $D$. For any ground-plane position $(x, y)$, the elevation in the plane containing that control polygon is then calculated as

$$zc = (-Ax - By - D)/C$$

Incremental methods can then be used to calculate control elevations over positions in the ground-plane grid. To carry out these calculations efficiently, we subdivide the ground plane into a smaller grid of $xy$ positions and project each polygon control surface onto the ground plane, as shown in Figure 22. From this projection, we determine which grid positions are below each control polygon. This can be accomplished using procedures similar to those in scan-line area filling. That is, for each $y$ “scan line” in the ground plane mesh that crosses the polygon edges, we calculate scan-line intersections and determine which grid positions are within the control-polygon projection. Calculations for the control elevations at these grid positions are performed incrementally as

$$zc_{i+1,j} = zc_{i,j} - \Delta x(A/C), \quad zc_{i,j+1} = zc_{i,j} - \Delta y(B/C)$$

(6)

where $\Delta x$ and $\Delta y$ are the grid separations in the $x$ and $y$ directions. This procedure is particularly fast when parallel vector methods are applied to process the control-plane grid positions.

Figure 23 shows a scene constructed using control planes to structure the surfaces for the terrain, water, and clouds above a ground plane.
Surface-rendering algorithms were then applied to smooth out the polygon edges and to provide the appropriate surface colors.

**Self-Squaring Fractals**

Another method for generating fractal objects is to apply a transformation function repeatedly to points in complex space. In two dimensions, a complex number can be represented as $z = x + iy$, where $x$ and $y$ are real numbers and $i^2 = -1$. In three-dimensional and four-dimensional space, points are represented with quaternions. A complex squaring function $f(z)$ is one that involves the calculation of $z^2$, and we can use some self-squaring functions to generate fractal shapes.

Depending on the initial position selected for the iteration, repeated application of a self-squaring function will produce one of three possible outcomes (Figure 24):

- The transformed position can diverge to infinity.
- The transformed position can converge to a finite limit point, called an attractor.
- The transformed position remains on the boundary of a region.

For example, the nonfractal squaring operation $f(z) = z^2$ in the complex plane transforms positions according to their relation to the unit circle (Figure 25). Any point $z$ whose magnitude $|z|$ is greater than 1 is transformed through a sequence of positions that tend to infinity. A point with $|z| < 1$ is transformed toward the coordinate origin. Points that are originally on the circle, $|z| = 1$, remain on the circle. Although the $z^2$ transformation does not produce a fractal, some complex squaring operations generate a fractal curve as the boundary between those positions that move toward infinity and those that tend toward a finite limit. A closed fractal boundary generated with a squaring operation is called a **Julia set**.

In general, we can locate the fractal boundary for a squaring function by testing the behavior of selected positions. If a position is transformed so that it either diverges to infinity or converges to an attractor point, we can try another...
nearby position. We repeat this process until we eventually locate a position on the fractal boundary. Then, iteration of the squaring transformation generates the fractal shape. For simple transformations in the complex plane, a quicker method for locating positions on the fractal curve is to use the inverse of the transformation function. An initial point chosen on the inside or outside of the curve will then converge to a position on the fractal curve (Figure 26).

A function that is rich in fractals is the squaring transformation

$$z' = f(z) = \lambda z(1 - z)$$

with \(\lambda\) as a complex constant. For this function, we can use the inverse method to locate the fractal curve. We first rearrange terms to obtain the quadratic equation

$$z^2 - z + \frac{z'}{\lambda} = 0$$

The inverse transformation is then the quadratic formula

$$z = f^{-1}(z') = \frac{1}{2} \left(1 \pm \sqrt{1 - \frac{4z'}{\lambda}}\right)$$

Using complex arithmetic operations, we solve this equation for the real and imaginary parts of \(z\) as

$$x = \text{Re}(z) = \frac{1}{2} \left(1 \pm \sqrt{|\text{discr}| + \text{Re}(\text{discr})}\right)$$

$$y = \text{Im}(z) = \pm \frac{1}{2} \sqrt{|\text{discr}| - \text{Re}(\text{discr})}$$

where the discriminant of the quadratic formula is \(\text{discr} = 1 - \frac{4z'}{\lambda}\). A few initial values for \(x\) and \(y\) (say, 10) can be calculated and ignored before we begin to plot the fractal curve. Also, because this function yields two possible transformed \((x, y)\) positions, we can randomly choose either the plus or the minus sign at each step of the iteration so long as \(\text{Im}(\text{discr}) \geq 0\). Whenever \(\text{Im}(\text{discr}) < 0\), the two possible positions are in the second and fourth quadrants. In this case, \(x\) and \(y\) must have opposite signs. The following program gives an implementation for this self-squaring function, and two example curves are plotted in Figure 27.

\[\text{FIGURE 26}\]
Locating the fractal boundary curve using the inverse, self-squaring function \(z = f^{-1}(z')\).

\[\text{FIGURE 27}\]
Two fractal curves generated with the inverse of the function \(f(z) = \lambda z(1 - z)\) by procedure \texttt{selfSqTransf}, using (a) \(\lambda = 3\) and (b) \(\lambda = 2 + i\). Each curve is plotted with 10,000 points.
#include <GL/glut.h>
#include <stdlib.h>
#include <math.h>

/* Set initial size of display window. */
GLsizei winWidth = 600, winHeight = 600;

/* Set coordinate limits in complex plane. */
GLfloat xComplexMin = -0.25, xComplexMax = 1.25;
GLfloat yComplexMin = -0.75, yComplexMax = 0.75;

struct complexNum
{
    GLfloat x, y;
};

void init (void)
{
    /* Set color of display window to white. */
    glClearColor (1.0, 1.0, 1.0, 0.0);
}

void plotPoint (complexNum z)
{
    glBegin (GL_POINTS);
    glVertex2f (z.x, z.y);
    glEnd ();
}

void solveQuadraticEq (complexNum lambda, complexNum * z)
{
    GLfloat lambdaMagSq, discrMag;
    complexNum discr;
    static complexNum fourOverLambda = { 0.0, 0.0 };
    static GLboolean firstPoint = true;
    if (firstPoint) {
        /* Compute the complex number: 4.0 divided by lambda. */
        lambdaMagSq = lambda.x * lambda.x + lambda.y * lambda.y;
        fourOverLambda.x = 4.0 * lambda.x / lambdaMagSq;
        fourOverLambda.y = -4.0 * lambda.y / lambdaMagSq;
        firstPoint = false;
    }
    discr.x = 1.0 - (z->x * fourOverLambda.x - z->y * fourOverLambda.y);
    discr.y = z->x * fourOverLambda.y + z->y * fourOverLambda.x;
    discrMag = sqrt (discr.x * discr.x + discr.y * discr.y);
    /* Update z, checking to avoid the square root of a negative number. */
    if (discrMag + discr.x < 0)
    {
        z->x = 0;
    }
    else
    {
        z->x = sqrt ((discrMag + discr.x) / 2.0);
    }
}
if (discrMag - discr.x < 0)
    z->y = 0;
else
    z->y = 0.5 * sqrt ((discrMag - discr.x) / 2.0);
/* For half the points, use negative root.
   * placing point in quadrant 3.
   */
if (rand ( ) < RAND_MAX / 2) {
    z->x = -z->x;
    z->y = -z->y;
}
/* When imaginary part of discriminant is negative, point
 * should lie in quadrant 2 or 4, so reverse sign of x.
 */
if (discr.y < 0)
    z->x = -z->x;
/* Complete the calculation for the real part of z. */
z->x = 0.5 * (1 - z->x);
}

void selfSqTransf (complexNum lambda, complexNum z, GLint numPoints)
{
    GLint k;
    /* Skip the first few points. */
    for (k = 0; k < 10; k++)
        solveQuadraticEq (lambda, &z);
    /* Plot the specified number of transformation points. */
    for (k = 0; k < numPoints; k++) {
        solveQuadraticEq (lambda, &z);
        plotPoint (z);
    }
}

void displayFcn (void)
{
    GLint numPoints = 10000; // Set number of points to be plotted.
    complexNum lambda = { 3.0, 0.0 }; // Set complex value for lambda.
    complexNum z0 = { 1.5, 0.4 }; // Set initial point in complex plane.
    glClear (GL_COLOR_BUFFER_BIT); // Clear display window.
    glColor3f (0.0, 0.0, 1.0); // Set point color to blue.
    selfSqTransf (lambda, z0, numPoints);
    glFlush ( );
}
void winReshapeFcn (GLint newWidth, GLint newHeight)
{
    /* Maintain an aspect ratio of 1.0, assuming that
    * width of complex window = height of complex window.
    */
    glViewport (0, 0, newHeight, newHeight);
    glMatrixMode (GL_PROJECTION);
    glLoadIdentity ();
    gluOrtho2D (xComplexMin, xComplexMax, yComplexMin, yComplexMax);
    glClear (GL_COLOR_BUFFER_BIT);
}

void main (int argc, char** argv)
{
    glutInit (&argc, argv);
    glutInitDisplayMode (GLUT_SINGLE | GLUT_RGB);
    glutInitWindowPosition (50, 50);
    glutInitWindowSize (winWidth, winHeight);
    glutCreateWindow ("Self-Squaring Fractal");
    init ( );
    glutDisplayFunc (displayFcn);
    glutReshapeFunc (winReshapeFcn);
    glutMainLoop ( );
}

A three-dimensional plot in variables $x$, $y$, and $\lambda$ of the self-squaring function $f(z) = \lambda z (1 - z)$, with $|\lambda| = 1$, is given in Figure 28. Each cross-sectional slice of this plot is a fractal curve in the complex plane.

Another squaring operation that produces a variety of fractal shapes is a slightly modified $z^2$ transformation. In this case, the fractal is the boundary region around the set of complex values $z$ that do not diverge under the squaring transformation

$$
    z_0 = z \\
    z_k = z_{k-1}^2 + z_0 \quad k = 1, 2, 3, \ldots
$$

(11)

Thus, we first select a point $z$ in the complex plane, then we compute the transformed position $z^2 + z$. In the next step, we square this transformed position and add the original $z$ value. We repeat this procedure until we can determine whether or not the transformation is diverging.

Mathematicians had been aware of the unusual features of such squaring functions for some time, but these functions were difficult to analyze without computing aids. After the development of the digital computer, the convergence boundary for Equation 11 was plotted on a line printer. As the capabilities of digital computing increased, further graphical investigation into the properties of this function were possible. Subsequently, using more sophisticated computer-graphics techniques, Benoit Mandelbrot extensively studied this function, and the set of points that do not diverge under Equation 11 has become known as the

Mandelbrot set.
To implement Equation 11, we first choose a rectangular area in the complex plane. Positions within this area are then mapped to color-coded pixel positions within a display window on a video monitor (Figure 29). The pixel colors are chosen according to the rate of divergence of the corresponding point in the complex plane under Equation 11. If the magnitude of a complex number is greater than 2, then it will diverge quickly as it is squared repeatedly. Therefore, we can set up a loop to repeat the squaring operations until either the magnitude of the complex number exceeds 2 or we have reached a preset number of iterations. The maximum number of iterations depends on the amount of detail that we want to display and the number of points to be plotted. This value is often set to some value between 100 and 1,000, although lower values can be used to speed up the calculations. With lower settings for the iteration limit, however, we do tend to lose some detail along the boundary (Julia set) of the convergence region. At the end of the loop, we select a color value according to the number of iterations executed by the loop. For example, we can color the pixel black if the iteration count is at the maximum value (a nondiverging point), and we can color the pixel red if the iteration count is near 0. Other color values can then be chosen according to the value of the iteration count within the interval from 0 to the maximum value. By choosing different color mappings and different sections of the complex plane, we can generate a variety of dramatic displays for positions in the vicinity of the fractal boundary that encloses the nondivergent points. One choice for color coding the pixel positions in the region around the Mandelbrot set is shown in Color Plate 39.

An implementation of Equation 11 for displaying the set of convergence points and its boundaries is given in the following program. The major part of the
convergence set is contained within the following region of the complex plane:

\[-2.00 \leq \text{Re}(z) \leq 0.50\]
\[-1.20 \leq \text{Im}(z) \leq 1.20\]

We can explore the details along the boundary of the Mandelbrot set by choosing successively smaller rectangular regions in the complex plane so that we can zoom in on selected areas of the display. Color Plate 39 shows a color-coded display of the region around the convergence set and a series of zooms that illustrate some of the remarkable features of this squaring transformation.

```c
#include <GL/glut.h>

/* Set initial size of the display window. */
GLsizei winWidth = 500, winHeight = 500;

/* Set limits for the rectangular area in complex plane. */
GLfloat xComplexMin = -2.00, xComplexMax = 0.50;
GLfloat yComplexMin = -1.25, yComplexMax = 1.25;

GLfloat complexWidth = xComplexMax - xComplexMin;
GLfloat complexHeight = yComplexMax - yComplexMin;

class complexNum {
    public:
        GLfloat x, y;
    };

struct color { GLfloat r, g, b; };

void init (void)
{
    /* Set display-window color to white. */
    glClearColor (1.0, 1.0, 1.0, 0.0);
}

void plotPoint (complexNum z)
{
    glBegin (GL_POINTS);
    glVertex2f (z.x, z.y);
    glEnd ( );
}

/* Calculate the square of a complex number. */
complexNum complexSquare (complexNum z)
{
    complexNum zSquare;

    zSquare.x = z.x * z.x - z.y * z.y;
    zSquare.y = 2 * z.x * z.y;
    return zSquare;
}
```
GLint mandelSqTransf (complexNum z0, GLint maxIter)
{
    complexNum z = z0;
    GLint count = 0;
    /* Quit when z * z > 4 */
    while ((z.x * z.x + z.y * z.y <= 4.0) && (count < maxIter)) {
        z = complexSquare (z);
        z.x += z0.x;
        z.y += z0.y;
        count++;
    }
    return count;
}

void mandelbrot (GLint nx, GLint ny, GLint maxIter)
{
    complexNum z, zIncr;
    color ptColor;
    GLint iterCount;
    zIncr.x = complexWidth / GLfloat (nx);
    zIncr.y = complexHeight / GLfloat (ny);
    for (z.x = xComplexMin; z.x < xComplexMax; z.x += zIncr.x)
        for (z.y = yComplexMin; z.y < yComplexMax; z.y += zIncr.y) {
            iterCount = mandelSqTransf (z, maxIter);
            if (iterCount >= maxIter)
                /* Set point color to black. */
                ptColor.r = ptColor.g = ptColor.b = 0.0;
            else if (iterCount > (maxIter / 8))
                /* Set point color to orange. */
                ptColor.r = 1.0;
                ptColor.g = 0.5;
                ptColor.b = 0.0;
            else if (iterCount > (maxIter / 10))
                /* Set point color to red. */
                ptColor.r = 1.0;
                ptColor.g = ptColor.b = 0.0;
            else if (iterCount > (maxIter /20))
                /* Set point color to dark blue. */
                ptColor.b = 0.5;
                ptColor.r = ptColor.g = 0.0;
            else if (iterCount > (maxIter /40))
                /* Set point color to yellow. */
                ptColor.r = ptColor.g = 1.0;
                ptColor.b = 0.0;
            else if (iterCount > (maxIter /100))
                /* Set point color to dark green. */
                ptColor.r = ptColor.b = 0.0;
                ptColor.g = 0.3;
        }
}
Complex-function transformations, such as Equation 7, can be extended to produce fractal surfaces and fractal solids. Methods for generating these objects use quaternion representations for transforming points in
three-dimensional and four-dimensional space. A quaternion has four components, with one real-number term and three imaginary-number terms. We can represent a quaternion in the following form, as an extension of the concept of a number in the complex plane:

\[ q = s + ia + jb + kc \]  

(12)

where \( i^2 = j^2 = k^2 = -1 \). The real-number term \( s \) is also referred to as the scalar part of the quaternion, and the imaginary terms are called the quaternion vector part \( \mathbf{v} = (a, b, c) \).

Using the rules for quaternion multiplication and addition, we can apply self-squaring functions and other iteration methods to generate surfaces of fractal objects. A basic procedure is to test points in complex space until we can identify the boundary between the diverging and nondiverging positions. For example, if we first locate a nondiverging (interior) position, then we test neighboring points from that position until a diverging (exterior) point is identified. The preceding interior point is then retained as a boundary-surface position. Neighbors of this surface point are then tested to determine whether they are inside (converging) or outside (diverging). Any inside point that connects to an outside point is a surface point. In this way, the procedure threads its way along the fractal boundary without straying far from the surface. When four-dimensional fractals are generated, three-dimensional slices are projected onto the two-dimensional surface of the video monitor.

Procedures for generating self-squaring fractals in four-dimensional space require considerable computation time for evaluating the iteration function and for testing positions for convergence or divergence. Each point on a surface can be represented as a small cube, giving the inner and outer limits of the surface. Output from such programs for the three-dimensional projections of the fractal typically contain more than a million vertices for the surface cubes. We display the fractal object by applying illumination models to determine the color for each surface cube. Visible-surface detection methods are also applied so that only visible surfaces of the object are displayed.

Self-Inverse Fractals

Various geometric inversion transformations can be used to create fractal shapes. Again, we start with an initial set of points, and we repeatedly apply nonlinear inversion operations to transform the initial points into a fractal.

For example, we consider a two-dimensional inversion transformation with respect to a circle of radius \( r \) and center position \( P_c = (x_c, y_c) \). A point \( P \) outside the circle is inverted to a position \( P' \) inside the circle (Figure 30) with the transformation

\[ (P, \mathbf{P})(P', \mathbf{P'}) = r^2 \]  

(13)

![Figure 30](image-url)  
**FIGURE 30** Inverting point \( P \) to a position \( P' \) inside a circle with radius \( r \).
where both \( P \) and \( P' \) lie on a straight line passing through the circle center \( P_c \). We can also use Equation 13 to transform positions that are inside the circle. Some inside positions transform to outside positions, while other inside positions transform to inside positions.

If the coordinates of the two points are represented as \( P = (x, y) \) and \( P' = (x', y') \), we can write Equation 13 as

\[
[(x - x_c)^2 + (y - y_c)^2]^{1/2}[(x' - x_c)^2 + (y' - y_c)^2]^{1/2} = r^2
\]

Also, because the two points are along a line passing through the circle center, we have \((y - y_c)/(x - x_c) = (y' - y_c)/(x' - x_c)\). Therefore, the transformed coordinate values for position \( P' \) are

\[
x' = x_c + \frac{r^2(x - x_c)}{(x - x_c)^2 + (y - y_c)^2}, \quad y' = y_c + \frac{r^2(y - y_c)}{(x - x_c)^2 + (y - y_c)^2}
\]  \quad (14)

Thus, points outside the circle are mapped to positions within the circle circumference, with distant points \( (\pm \infty) \) transformed to the circle center. Conversely, points near the circle center are mapped to distant points outside the circle. As we move out from the circle center, points are mapped to outside positions closer to the circle circumference. Also, inside points near the circumference are transformed to inside positions closer to the circle center. For example, outside \( x \) values in the range from \( r \) to \( +\infty \) map to \( x' \) values in the range from \( \frac{r}{2} \) to \( 0 \), for a circle centered at the origin, \((x_c, y_c) = (0, 0)\). Inside values for \( x \) in the range from \( \frac{r}{2} \) to \( 0 \) map to the \( x' \) values from \( r \) to \( +\infty \), for a circle centered at the origin, and inside \( x \) values from \( \frac{r}{2} \) to \( r \) are transformed to values in the range from \( r \) to \( \frac{r}{2} \). Similar results are obtained for negative values of \( x \).

We can apply this transformation to various objects, such as straight lines, circles, or ellipses. A straight line that passes through the circle center is invariant under this inversion transformation; it maps into itself. However, a straight line that does not pass through the circle center inverts into a circle whose circumference contains the center point \( P_c \); and any circle that passes through the center of the reference circle is inverted into a straight line that does not pass through the circle center. If the circle does not intersect the center of the reference circle, it inverts into another circle, as in Figure 31. Another invariant inversion is the transformation of a circle that is orthogonal to the reference circle. That is, the tangents of the two circles are perpendicular at the intersection points.

We can create various fractal shapes with this inversion transformation by starting with a set of circles and repeatedly applying the transformation using different reference circles. Similarly, we can apply circle inversion with respect to a set of straight lines. Comparable inversion methods can be developed for other two-dimensional shapes. In addition, we can generalize the procedure to spheres, planes, or other three-dimensional objects.

\[\text{FIGURE 31}\]

Inversion of a circle that does not pass through the origin of the reference circle.
2 Particle Systems

For some applications, it is often useful to describe one or more objects using a collection of disjoint pieces, called particle systems. This approach can be applied to describe objects with fluid-like properties that change over time by flowing, billowing, spattering, expanding, imploding, or exploding. Objects with these characteristics include clouds, smoke, fire, fireworks, waterfalls, and water spray. Particle systems have been employed, for instance, to model the planet explosion and expanding wall of fire from the “genesis bomb” in the motion picture Star Trek II: The Wrath of Khan. In addition, particle-system methods have been used to model other kinds of objects, including clumps of grass.

In a typical application, a system of particles is defined within some spatial region and then random processes are applied to vary system parameters over time. Each particle has attributes associated with it that determine its behavior, such as its appearance, size, initial color, motion path and velocity, and lifetime. Values for these attributes are determined when the particle is generated, and are used by processes within the system during the particle’s lifetime.

Particle systems are extremely flexible because particle attribute values are determined algorithmically. This means that virtually any type of behavior can be built into the system. Particle shapes could be described with small spheres, ellipsoids, or boxes, which can be fixed or can vary randomly over time. Particle transparency, color, and movement can all be randomly chosen, can be static, or can evolve. Motion paths for the particles could be described kinematically or defined with forces such as a gravity field. Lifetimes can be identical for all particles or can vary from one particle to another. The number and type of particles can be fixed, or new particles can be generated whenever existing particles reach the ends of their lives.

As each particle moves, its path is plotted and displayed in a particular color. For example, a fireworks pattern can be displayed by randomly generating particles within a spherical region of space and allowing them to move radially outward, as in Figure 32. The particle paths could be color-coded from red to yellow, for instance, to simulate the temperature of the exploding particles.

Alternatively, rather than simply displaying the current position of each particle, “trajectory” particles can be used to model realistic displays of grass clumps (Figure 33). Particles begin their lives at the ground, are shot up into the air, and fall back to earth under gravity, with each point along their trajectory being
shown. The particle paths can originate within a tapered cylinder, and particles may be color-coded from green to yellow.

Color Plate 40 illustrates a particle-system simulation of a waterfall. The water particles fall from a fixed elevation, are deflected by an obstacle, then splash up from the ground. Different colors are used to distinguish the particle paths at each stage.

A composite scene formed with a variety of representations is given in Color Plate 41. The scene is modeled using particle-system grass and fractal mountains, in addition to texture mapping and other surface-rendering procedures.

3 Grammar-Based Modeling Methods

A number of other procedural methods can be used to design object shapes or levels of surface detail. Shape grammars are sets of production rules that can be applied to an initial object to add layers of detail that are harmonious with the original shape. Transformations can be applied to alter the geometry (shape) of the object, or the transformation rules can be applied to add details for surface color or texture.

Given a set of production rules, a shape designer can experiment by applying different rules at each step of the transformation from a given initial object to the final structure. Figure 34 shows four geometric substitution rules for altering triangle shapes. The geometric transformations for these rules can be expressed algorithmically by the system based on an input picture drawn with a production-rule editor. That is, each rule can be described graphically by showing the initial and final shapes. Implementations can then be set up in Mathematica or some programming language with graphics capability.

An application of the geometric substitutions in Figure 34 is given in Figure 35, where Figure 35(d) is obtained by applying the four rules in succession, starting with the initial triangle in Figure 35(a).

Three-dimensional shape and surface features are transformed with similar operations. Figure 36 shows the results of geometric substitutions applied to polyhedra. The initial shape for the objects shown in Figure 37 is an icosahedron (a polyhedron with 20 faces). Geometric substitutions were applied to the plane faces of the icosahedron, and the resulting polygon vertices were projected to the surface of an enclosing sphere.

Another set of production rules for describing the shape of objects is called L-grammars, or graftals. These rules are typically used to generate plant displays. For instance, the topology of a tree can be described as a trunk, with some attached branches and leaves. A tree can then be modeled with rules to provide a particular connection of the branches and the leaves on the individual branches. The geometrical description is then given by placing the object structures at particular coordinate positions.

Color Plate 42 shows a scene containing various plants and trees, which was constructed with a commercial plant-generator package. Procedures in the software apply botanical laws to generate the shapes for the plants and trees.

![Figure 34](image) Four geometric substitution rules for subdividing and altering the shape of an equilateral triangle.
FIGURE 35
An equilateral triangle (a) is converted to shape (b) using substitution rules 1 and 2 in Figure 34. Rule 3 is then used to convert (b) into shape (c), which in turn is transformed to (d) using rule 4. [Courtesy of Andrew Glassner, Xerox PARC (Palo Alto Research Center). © 1992.]

FIGURE 36
A design created with geometric substitution rules for altering prism shapes. The initial shape for this design is a representation of Rubik’s Snake. [Courtesy of Andrew Glassner, Xerox PARC (Palo Alto Research Center). © 1992.]

FIGURE 37
Designs created on the surface of a sphere using triangle substitution rules applied to the plane faces of an icosahedron, followed by projections to the sphere surface. [Courtesy of Andrew Glassner, Xerox PARC (Palo Alto Research Center). © 1992.]
4 Summary

Fractal-geometry representations provide highly effective methods for describing natural phenomena. We can use these methods to model terrain, trees, bushes, water, and clouds, and to generate unusual graphics patterns. A fractal object can be described with a construction procedure and a fractal dimension. Fractal construction procedures include geometric constructions, midpoint-displacement methods, self-squaring operations in complex space, and inversion transformations. Other procedural methods for constructing object representations using transformation rules are shape grammars and graftals.

Objects that exhibit fluidity, such as clouds, smoke, fire, water, and things that explode or implode, can be modeled with particle systems. Using this representation scheme, we describe an object with a set of particles and the rules that govern the particle movements.

REFERENCES


EXERCISES

1 Using the random midpoint-displacement method, write a routine to create a mountain outline, starting with a horizontal line in the xy plane.

2 Write a routine to calculate elevations above a ground plane using the random midpoint-displacement method, given a set of corner elevations for the ground plane.

3 Write a program to display the terrain generated by the routine in the previous exercise. Set up a point light source and choose appropriate rendering parameters to display the set of triangle patches that result from the routine.

4 Write a program to display the fractal snowflake (Koch curve) for a given number of iterations.

5 Modify the program in the previous exercise to allow the user to set the number of iterations with which to draw the Koch curve using a slider in the display window. The Koch curve should be redrawn each time the value of the slider is changed.

6 Write a program to generate a fractal curve for a specified number of iterations using one of the generators in Figure 7 or Figure 8. What is the fractal dimension of the curve?

7 Modify the program in the previous exercise to allow the user to set the number of iterations with which to draw the chosen fractal using a slider in the display window. The chosen fractal should be redrawn each time the value of the slider is changed.

8 Write a program to generate fractal curves using the self-squaring function \( f(z) = z^2 + \lambda \), where the complex constant \( \lambda \) is specified as input.

9 Write a program to generate a fractal curve using the self-squaring function \( f(z) = iz(z^2 + 1) \), where \( i = \sqrt{-1} \).

10 Modify the programming example in Section 1 to use additional color levels in displaying the boundary regions around the Mandelbrot set.

11 Modify the program in the previous exercise to allow the colors and number of color levels to be given as input values.

12 Modify the program in the previous exercise to select and display any rectangular boundary region (the zoom area) around the Mandelbrot set.

13 Write a routine to implement point inversion, Equation 14, for a specified circle and a set of point positions.

14 Devise a set of geometric-substitution rules for altering the shape of an equilateral triangle.

15 Write a program for the previous exercise that displays the stages in the conversion of the triangle. Allow the user to move forward and backward through the stages using keyboard input.

16 Write a program to model and display an exploding sphere in the xy plane, using a particle system. Specify a beginning and end configuration.
for the explosion and divide the interval between
the two into several frames, each of which the user
can view by moving forward and backward using
keyboard input.

17 Modify the program of the previous exercise to
explode a firecracker (cylinder) in a similar way,
allowing the user to step back and forth through
each frame of the explosion.

IN MORE DEPTH
1. Use the concepts presented in this chapter to
incorporate a fractal design into your application
in an appropriate way. For example, if your appli-
cation contains natural terrain, you might use a
midpoint-displacement method to model the ter-
rain in a more realistic manner. Alternatively, you
might develop two-dimensional fractal patterns,
which can then be used as texture patterns to be
mapped onto the surface of some of the objects in
your application to produce more sophisticated
object textures.

2. Implement one of the two-dimensional fractals
described in this chapter as a programmable shader that you then use to shade one or more
of the objects in your scene. Modify the exam-
ple shader programs in the previous chapter to
produce a shader that exhibits self-similarity as
discussed in this chapter.
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Algorithmic Modeling
Color Plates

Color Plate 36
Modeling a scene using multiple object instancing. Fractal leaves are attached to a tree in randomly transformed positions, and several rotated and scaled instances of the tree are used to form a grove. The grass is modeled with multiple instances of green cones. (Courtesy of John C. Hart. Reprinted with permission.)
Color Plate 37
A fractal forest created with multiple instances of leaves, pine needles, grass, and tree bark. (Courtesy of John C. Hart. Reprinted with permission.)

Color Plate 38
Variations in terrain features modeled with fractional Brownian motion. (Courtesy of (a) R. V. Voss and B. B. Mandelbrot, adapted from The Fractal Geometry of Nature by Benoit B. Mandelbrot (W. H. Freeman and Co., New York, 1983); and (b) and (c) Ken Musgrave and Benoit B. Mandelbrot, Mathematics and Computer Science, Yale University.)
Color Plate 39
Zooming in on the fractal boundaries for transformation 23-n. Starting with a display of the Mandelbrot set, the black region in (a), and its surrounding areas, we zoom in on selected border regions (b) through (f). The white box outline shows the rectangular area selected for each successive zoom. Different color combinations are chosen at each step to enhance the displayed zoom patterns. (Copyright © Brian Evans. Reprinted with permission.)

Color Plate 40
Simulation of the behavior of a waterfall hitting a stone (circle). The water particles are deflected by the stone and then splash up from the ground. (Courtesy of Toby Howard. Reprinted with permission.)
Color Plate 41
A scene, entitled Road to Point Reyes, showing particle-system grass, fractal mountains, and texture-mapped surfaces. (Courtesy of Pixar © 1983 Pixar.)

Color Plate 42
Realistic scenery generated with the TDI-AMAP software package, which can generate over 100 varieties of plants and trees using procedures based on botanical laws. (Courtesy of Thomson Digital Image.)
The use of computer-graphics methods as an aid in scientific and engineering analysis is commonly referred to as **scientific visualization**. This involves the visualization of data sets and processes that may be difficult or impossible to analyze without graphical methods. For example, visualization techniques are needed to deal with the output of high-volume data sources such as computer monitors, satellite and spacecraft scanners, radio-astronomy telescopes, and medical scanners. Millions of data points are often generated from numerical solutions of computer simulations and from observational equipment, and it is difficult to determine trends and relationships by simply scanning the raw data. Similarly, visualization techniques are useful for analyzing processes that occur over a long time period or that cannot be observed directly, such as quantum-mechanical phenomena and special-relativity effects produced by objects traveling near the speed of light. Scientific visualization uses methods from computer graphics,
Visualization of Data Sets

image processing, computer vision, and other areas to visually display, enhance, and manipulate information to allow better understanding of the data. Similar methods employed by commerce, industry, and other nonscientific areas are sometimes referred to as **business visualization**.

Data sets are classified according to their spatial distribution and data type. Two-dimensional data sets have values distributed over a surface, and three-dimensional data sets have values distributed over the interior of a cube, a sphere, or some other region of space. Data types include scalars, vectors, tensors, and multivariate data.

1 Visual Representations for Scalar Fields

A **scalar quantity** is one that has a single value. Scalar data sets contain values that may be distributed in time, as well as over spatial positions, and the data values may also be functions of other scalar parameters. Some examples of physical scalar quantities are energy, density, mass, temperature, pressure, electric charge, electrical resistance, reflectivity, frequency, and water content.

A common method for visualizing a scalar data set is to use graphs or charts that show the distribution of data values as a function of other parameters, such as position and time. If the data are distributed over a surface, we could plot the data values as vertical bars rising from the surface, or we can interpolate the data values in some other way at selected surface positions. **Pseudo-color methods** are also used to distinguish different values in a scalar data set, and color-coding techniques can be combined with graph and chart methods. To color-code a scalar data set, we choose a range of colors and map the range of data values to the color range. For example, blue could be assigned to the lowest scalar value, and red could be assigned to the highest value. Color-coding a data set sometimes requires careful consideration because certain color combinations can lead to misinterpretations of the data.

**Contour plots** are used to display **isolines** (lines of constant value) for a scalar data set distributed over a surface. The isolines are spaced at a convenient interval to show the range and variation of the data values over the region of space. A typical application is a contour plot of elevations over a ground plane. Usually, contouring methods are applied to a set of data values that are distributed over a regular grid, as in Figure 1. Regular grids have equally spaced grid lines, and data values are known at the grid intersections. Numerical solutions of computer simulations are usually set up to produce data distributions on a regular grid, while observational data sets are often irregularly spaced. Contouring methods have been devised for various kinds of nonregular grids, but nonregular data distributions are often converted to regular grids. A two-dimensional contouring algorithm traces the isolines from cell to cell within the grid by checking the four corners of grid cells to determine which cell edges are crossed by a particular isoline. The isolines are usually plotted as straight line sections across each cell, as illustrated in Figure 2. Sometimes isolines are plotted with spline curves, but spline-fitting can lead to inconsistencies and misinterpretation of a data set. For example, two spline isolines could cross, or curved isoline paths might not be a true indicator of the data trends because data values are known only at the cell corners. Contouring packages can allow interactive adjustment of isolines by a researcher to correct any inconsistencies.

For three-dimensional scalar data fields, we can take cross-sectional slices and display the two-dimensional data distributions over the slices. We could either
color code the data values over a slice, or we could display isolines. Visualization packages typically provide a slicer routine that allows cross sections to be taken at any angle. Color Plate 43 shows a display generated by a commercial slicer-dicer package.

Instead of looking at two-dimensional cross-sections, we can plot one or more isosurfaces, which are simply three-dimensional contour plots. When two overlapping isosurfaces are displayed, the outer surface is made transparent so that we can view the shapes of both isosurfaces. Constructing an isosurface is similar to plotting isolines except that now we have three-dimensional grid cells and we need to check the data values at the eight corners of a cell to locate sections of an isosurface. Figure 3 shows some examples of isosurface intersections with grid cells. Isosurfaces are usually modeled with triangle meshes, and then surface-rendering algorithms are applied to display the final shape.

**Volume rendering**, which is often somewhat like an X-ray picture, is another method for visualizing a three-dimensional data set. The interior information about a data set is projected to a display screen using ray-casting methods. Along the ray path from each screen pixel (see Figure 4), interior data values are examined and encoded for display. Often, data values at the grid positions are averaged so that one value is stored for each voxel of the data space. How the data are encoded for display depends on the application. Seismic data, for example, are often examined to find the maximum and minimum values along each ray. The values can then be color-coded to give information about the width of the interval and the minimum value. In medical applications, the data values are opacity factors in the range from 0 to 1 for the tissue and bone layers. Bone layers are completely opaque, while tissue is somewhat transparent (that is, it has low opacity). Along each ray, the opacity factors are accumulated until either the total is greater than or equal to 1, or until the ray exits at the back of the three-dimensional data grid. The accumulated opacity value is then encoded and displayed as a pixel color or as a grayscale value.
2 Visual Representations for Vector Fields

A vector quantity \( \mathbf{V} \) in three-dimensional space has three scalar values \((V_x, V_y, V_z)\), one for each coordinate direction; and a two-dimensional vector has two components \((V_x, V_y)\). Another way to describe a vector quantity is by giving its magnitude \(|\mathbf{V}|\) and its direction as a unit vector \(\mathbf{u}\). As with scalars, vector quantities may be functions of position, time, and other parameters. Some examples of physical vector quantities are velocity, acceleration, force, electric current, and electric, magnetic, and gravitational fields.

One way to visualize a vector field is to plot each data point as a small arrow that shows the magnitude and direction of the vector. This method is most often used with cross-sectional slices, as in Color Plate 44, because it can be difficult to see the data trends in a three-dimensional region that is cluttered with overlapping arrows. Magnitudes for vector values can be represented as variations in the lengths of the arrows, or we could display color-coded arrows that are all the same size.

We can also represent vector values by plotting field lines, also called streamlines. Field lines are commonly used for electric, magnetic, and gravitational fields. The magnitude of the vector values is indicated by the spacing between field lines, and the direction of the field is represented by the tangents (slopes) of the field lines, as shown in Figure 5. Streamlines can be displayed as wide arrows, particularly when a whirlpool, or vortex, effect is present. For animations of fluid flow, the behavior of the vector field can be visualized by tracking particles along the flow direction.

Sometimes, only the magnitudes of the vector quantities are displayed. This is often done when multiple quantities are to be visualized at a single position, when the directions do not vary much in some region of space, or when vector directions are of less interest.

3 Visual Representations for Tensor Fields

A tensor quantity in three-dimensional space has nine components and can be represented with a \(3 \times 3\) matrix. Actually, this representation is used for a second-order tensor, and higher-order tensors do occur in some applications, particularly general-relativity studies. Some examples of physical second-order tensors are stress and strain in a material subjected to external forces, conductivity (or resistivity) of an electrical conductor, and the metric tensor, which gives the properties of a particular coordinate space. The stress tensor in Cartesian coordinates, for example, can be represented as

\[
\begin{bmatrix}
\sigma_x & \sigma_{xy} & \sigma_{xz} \\
\sigma_{yx} & \sigma_y & \sigma_{yz} \\
\sigma_{zx} & \sigma_{zy} & \sigma_z
\end{bmatrix}
\]  

(1)

Tensor quantities are encountered frequently in anisotropic materials, which have different properties in different directions. The \(x, xy,\) and \(xz\) elements of the...
conductivity tensor, for example, describe the contributions of electric-field components in the \( x \), \( y \), and \( z \) directions to the current in the \( x \) direction. Usually, physical tensor quantities are symmetric, so that the tensor has only six distinct values. For instance, the \( xy \) and \( yx \) components of the stress tensor have the same value.

Visualization schemes for representing all six components of a symmetric, second-order tensor quantity are based on devising shapes that have six parameters. One such graphical representation for a tensor is shown in Color Plate 45. The three diagonal elements of the tensor are used to construct the magnitude and direction of the arrow, and the three off-diagonal terms are used to set the shape and color of the elliptical disk.

Instead of trying to visualize all six components of a symmetric tensor quantity, we can reduce the tensor to a vector or a scalar. Using a vector representation, we can simply display the values for the diagonal elements of the tensor. In addition, by applying tensor-contraction operations, we can obtain a scalar representation. For example, stress and strain tensors can be contracted to generate a scalar strain-energy density that can be plotted at points in a material subject to external forces.

### 4 Visual Representations for Multivariate Data Fields

In some applications, we may want to represent multiple data values at each grid position over a region of space. This data often contains a mixture of scalar, vector, and tensor values. For example, fluid-flow data includes the fluid velocity, temperature, and density values at each three-dimensional position. Thus, we have five scalar values to display at each position, and the situation is similar to displaying a tensor field.

A method for displaying multivariate data fields is to construct graphical objects, sometimes referred to as *glyphs*, with multiple parts. Each part of a glyph represents a particular physical quantity. The size and color of each part can be used to display information about scalar magnitudes. To give directional information for a vector field, we can use a wedge, a cone, or some other pointing shape for the glyph part representing the vector. An example of the visualization of a multivariate data field using a glyph structure at selected grid positions is shown in Color Plate 46.

### 5 Summary

Visualization techniques use computer-graphics methods to analyze data sets, which can include scalar, vector, and tensor values in various combinations. Data representations can be accomplished with color-coding or with the display of different object shapes.

**REFERENCES**

EXERCISES

1. Write a routine to visualize a two-dimensional scalar data set using a pseudo-color representation.
2. Write a routine to visualize a two-dimensional scalar data set using contour lines.
3. Write a routine to visualize a two-dimensional vector data set using an arrow representation for the vector values. Use a fixed-size arrow with different color codings.
4. Write a routine to visualize a three-dimensional scalar data set using isosurfaces.
5. Write a routine to visualize a three-dimensional scalar data set using volume rendering. Provide the ability to choose between displaying the maximum, minimum, and average value of the voxels intersected along a given ray.
6. Write a routine to visualize a three-dimensional vector data set using an arrow representation that allows the view of a two-dimensional cross-section of data specified by the user. Use color coded arrows to indicate vector magnitude.
7. Find a moderately sized data set of interest to you on the internet that contains two-dimensional scalar data. Using the routines developed in the previous exercises, visualize the data using a pseudo-color representation and again using a contour line representation. Compare the two visualizations and list the advantages and disadvantages of displaying the data in each format.
8. Find a moderately sized data set of interest to you on the internet that contains three-dimensional scalar data. Using the routines developed in the previous exercises, visualize the data using isosurfaces and again using volume rendering. Compare the two visualizations and list the advantages and disadvantages of displaying the data in each format. Also explore the different possible schemes for dealing with multiple voxel intersections in volume rendering (e.g., value-averaging, min-max values).
9. Find a moderately sized data set of interest to you on the Internet that contains two-dimensional vector data. Using the routine developed in a previous exercise, visualize the data using an arrow representation for the vector values. Explore the difference between using color-coded arrows to indicate vector magnitude and using arrow size.

IN MORE DEPTH

1. Find a large data set of interest to you containing either two- or three-dimensional scalar data. Experiment with at least two different ways of presenting the data and write a program that allows switching between each of these display methods. You may include input routines to allow the user to interact with the data to make it more understandable.
2. Carry out the same procedure as the previous exercise using a large data set containing either two- or three-dimensional vector data.
Visualization of Data Sets

Color Plates

**Color Plate 43**
Cross-sectional slices of a three-dimensional data set. (Courtesy of Spyglass, Inc.)

**Color Plate 44**
Arrow representation for a vector field over cross-sectional slices. (Courtesy of the National Center for Supercomputing Applications (NCSA) and the Board of Trustees of the University of Illinois.)

**Color Plate 45**
Representing stress and strain tensors with an elliptical disk and an arrow over the surface of a stressed material. (Courtesy of the National Center for Supercomputing Applications (NCSA) and the Board of Trustees of the University of Illinois.)

**Color Plate 46**
One frame from an animated visualization of a time-varying multivariate data field using glyphs. The wedge-shaped part of the glyph indicates the direction of a vector quantity at each point. (Courtesy of the National Center for Supercomputing Applications (NCSA) and the Board of Trustees of the University of Illinois.)
A variety of mathematical concepts and techniques are employed in computer-graphics algorithms. Here, we provide a brief reference for the methods from analytic geometry, linear algebra, vector analysis, tensor analysis, complex numbers, quaternions, calculus, numerical analysis, and other areas that are referred to in the discussions throughout this book.

1 Coordinate Reference Frames

Both Cartesian and non-Cartesian reference frames are often useful in computer-graphics applications. We typically specify coordinates in a graphics program using a Cartesian reference system, but the initial specification of a scene could be given in a non-Cartesian frame of reference. Spherical, cylindrical, or other symmetries often can be exploited to simplify expressions involving object descriptions or manipulations.

Two-Dimensional Cartesian Screen Coordinates

For the device-independent commands within a graphics package, screen-coordinate positions are referenced within the first quadrant of a two-dimensional Cartesian frame in standard position, as shown in Figure 1(a). The coordinate origin for this reference frame is at the lower-left screen corner. Scan lines, however, are numbered from 0 at the top of the screen, so that screen positions are represented internally with respect to the upper-left corner of the screen. Therefore, device-dependent commands, such as those for interactive input and display-window manipulations, often reference screen coordinates using the inverted Cartesian frame shown in Figure 1(b). Horizontal coordinate values in the two systems are the same, and an inverted $y$ value is converted to a $y$ value measured from the bottom of the screen with the calculation

$$y = y_{\text{max}} - y_{\text{invert}}$$  

In some application packages, the screen-coordinate origin can be placed at an arbitrary position, such as the center of the screen.

FIGURE 2
A polar-coordinate reference frame, formed with concentric circles and radial lines.

FIGURE 3
Relationship between polar and Cartesian coordinates.

Standard Two-Dimensional Cartesian Reference Frames

We use Cartesian systems in standard position for world-coordinate specifications, viewing coordinates, and other references within the two-dimensional viewing pipeline. Coordinates in these frames can be positive or negative, with any range of values. To display a view of a two-dimensional picture, we designate a clipping window and a viewport to map a section of the picture to screen coordinates.

Polar Coordinates in the $xy$ Plane

A frequently used two-dimensional non-Cartesian system is a polar-coordinate reference frame (Figure 2), where a coordinate position is specified with a radial distance $r$ from the coordinate origin and an angular displacement $\theta$ from the horizontal. Positive angular displacements are counterclockwise, and negative angular displacements are clockwise. The relation between Cartesian and polar coordinates is shown in Figure 3. Considering the right triangle in Figure 4, and using the definition of the trigonometric functions, we transform from polar coordinates to Cartesian coordinates with the expressions

$$x = r \cos \theta, \quad y = r \sin \theta \quad (2)$$

The inverse transformation from Cartesian to polar coordinates is

$$r = \sqrt{x^2 + y^2}, \quad \theta = \tan^{-1}\left(\frac{y}{x}\right) \quad (3)$$

Angular values can be measured either in degrees or in dimensionless units (radians). One radian is defined as a measure for an angle that is subtended by a circular arc that has a length equal to the circle radius. This definition is illustrated in Figure 5, which shows two intersecting lines in a plane and a circle centered on the intersection point $P$. For any circle centered on $P$, the value of angle $\theta$ in radians is given by the ratio

$$\theta = \frac{s}{r} \quad (\text{radians}) \quad (4)$$

where $s$ is the length of the circular arc subtending $\theta$, and $r$ is the radius of the circle. Total angular distance around point $P$ is the length of the circle perimeter ($2\pi r$) divided by $r$, or $2\pi$ radians. In terms of degrees, a circle circumference is
divided into 360 arcs of equal length, so that each arc subtends an angle of 1 degree. Therefore, $360^\circ = 2\pi$ radians.

Other conics, besides circles, can be used to specify coordinate positions. For example, using concentric ellipses instead of circles, we can give coordinate positions in elliptical coordinates. Similarly, other types of symmetries can be exploited with hyperbolic or parabolic plane coordinates.

**Standard Three-Dimensional Cartesian Reference Frames**

Figure 6(a) shows the conventional orientation for the coordinate axes in a three-dimensional Cartesian reference system. This is called a right-handed system because the right-hand thumb points in the positive $z$ direction when we imagine grasping the $z$ axis with the fingers curling from the positive $x$ axis to the positive $y$ axis (through $90^\circ$), as illustrated in Figure 6(b). In most computer-graphics programs, we specify object descriptions and other coordinate parameters in right-handed Cartesian coordinates. For discussions throughout this book, we assume that all Cartesian reference frames are right-handed unless specifically stated otherwise.

Cartesian reference frames are **orthogonal coordinate systems**, which just means that the coordinate axes are perpendicular to each other. Also, in Cartesian frames, the axes are straight lines. But coordinate systems with curved axes are useful in many applications. Most of these systems are also orthogonal in the sense that the axial directions at any point in the space are mutually perpendicular.

**Three-Dimensional Cartesian Screen Coordinates**

When a view of a three-dimensional scene is displayed on a video monitor, depth information is stored for each screen position. The three-dimensional position corresponding to each screen point is often referenced with the left-handed system shown in Figure 7. In this case, the left-hand thumb points in the positive
z direction when we imagine grasping the z axis so that the fingers of the left hand curl from the positive x axis to the positive y axis through 90°. Positive z values indicate positions behind the screen for each point in the xy plane, and larger values along the positive z axis are interpreted as being farther from the viewer.

Three-Dimensional Curvilinear-Coordinate Systems

Any non-Cartesian reference frame is referred to as a curvilinear-coordinate system. The choice of coordinate system for a particular graphics application depends on a number of factors, such as symmetry, ease of computation, and visualization advantages. Figure 8 shows a general curvilinear-coordinate reference frame formed with three coordinate surfaces, where each surface has one coordinate held constant. For instance, the $x_1x_2$ surface is defined with $x_3 = \text{const}$. Coordinate axes in any reference frame are the intersection curves of the coordinate surfaces. If the coordinate surfaces intersect everywhere at right angles, we have an orthogonal curvilinear-coordinate system. Nonorthogonal, curvilinear reference frames are useful for some applications, such as visualizations of motions governed by the laws of general relativity, but they are used less frequently in computer graphics than orthogonal systems.

A cylindrical-coordinate specification of a spatial position is shown in Figure 9 in relation to a Cartesian reference frame. The surface of constant $\rho$ is a vertical cylinder; the surface of constant $\theta$ is a vertical plane containing the z axis; and the surface of constant z is a horizontal plane parallel to the Cartesian xy plane. We transform from a cylindrical-coordinate specification to a Cartesian reference frame with the calculations

\[
    x = \rho \cos \theta, \quad y = \rho \sin \theta, \quad z = z \tag{5}
\]

Another commonly used curvilinear-coordinate specification is the spherical-coordinate system shown in Figure 10. Spherical coordinates are sometimes referred to as polar coordinates in three-dimensional space. The surface of constant $r$ is a sphere; the surface of constant $\theta$ is again a vertical plane containing the z axis; and the surface of constant $\phi$ is a cone with apex at the coordinate origin. If $\phi < 90^\circ$, the cone is above the xy plane. If $\phi > 90^\circ$, the cone is below the xy plane.
We transform from a spherical-coordinate specification to a Cartesian reference frame with the calculations
\[ x = r \cos \theta \sin \phi, \quad y = r \sin \theta \sin \phi, \quad z = r \cos \phi \] 
(6)

Solid Angle

The definition for a solid angle \( \omega \) is formulated by analogy with the definition for a two-dimensional radian-angle \( \theta \) between two intersecting lines (Eq. 4). For a three-dimensional angle, however, we consider a cone with its apex at a point \( P \) and a sphere centered at \( P \), as shown in Figure 11. The solid angle \( \omega \) within the cone-shaped region with apex at \( P \) is defined as
\[ \omega = \frac{A}{r^2} \] 
(7)
where \( A \) is the area of the spherical surface intersected by the cone and \( r \) is the radius of the sphere.

Also, in analogy with two-dimensional surface coordinates, the dimensionless unit for solid angles is called the steradian. The total solid angle about point \( P \) is the total area of the spherical surface \((4\pi r^2)\) divided by \( r^2 \), or \( 4\pi \) steradians.

2 Points and Vectors

There is a fundamental difference between the concept of a geometric point and that of a vector. A point is a position specified with coordinate values in some reference frame, where the coordinates and other properties for the point depend on our choice for the frame of reference. A vector, on the other hand, has properties that are independent of any particular coordinate system.

Point Properties

Figure 12 illustrates the coordinate specification for a two-dimensional point position \( P \) in two reference frames. In frame A, the point has coordinates that are given by the ordered pair \((x, y)\), and its distance from the origin is \( \sqrt{x^2 + y^2} \). In frame B, the same point has coordinates \((0, 0)\), and the distance to the coordinate origin of frame B is 0.
Vector Properties

In a chosen coordinate system, we can define a vector as the difference between two point positions. Thus, for the two-dimensional points \( P_1 \) and \( P_2 \) in Figure 13, we can specify a vector as

\[
V = P_2 - P_1 = (x_2 - x_1, y_2 - y_1) \tag{8}
\]

where the Cartesian components (or Cartesian elements) \( V_x \) and \( V_y \) are the projections of \( V \) onto the \( x \) and \( y \) axes. We could also obtain these same vector components using two other point positions in this coordinate reference frame. In fact, there are an infinite number of point pairs that will produce the same vector components, and a vector is often defined with a single point position relative to the current frame of reference. Therefore, a vector has no fixed position within a coordinate system. Also, if we transform the representation for \( V \) to another reference frame, the coordinates for the positions \( P_1 \) and \( P_2 \) change, but the basic properties of the vector remain unchanged.

We can describe a vector as a directed line segment that has two fundamental properties: magnitude and direction. For the two-dimensional vector in Figure 13, we calculate the vector magnitude using the Pythagorean theorem, which gives us the distance along the vector direction between its two endpoint positions.

\[
|V| = \sqrt{V_x^2 + V_y^2} \tag{9}
\]

We can specify the vector direction in various ways. For example, we can give the direction in terms of the angular displacement from the horizontal as

\[
\alpha = \tan^{-1} \left( \frac{V_y}{V_x} \right) \tag{10}
\]

A vector has the same magnitude and direction no matter where we position the vector within a single coordinate system. And the vector magnitude is independent of the coordinate representation. However, if we transform the vector to another reference frame, the values for its components and direction within that reference frame may change. For example, we could transform the vector to a rotated Cartesian frame so that the vector direction is along the new \( y \) direction.
For a three-dimensional Cartesian vector representation \( \mathbf{V} = (V_x, V_y, V_z) \), the vector magnitude is
\[
|\mathbf{V}| = \sqrt{V_x^2 + V_y^2 + V_z^2}
\]  
(11)
And we can give the vector direction in terms of the direction angles, \( \alpha \), \( \beta \), and \( \gamma \), that the vector makes with each of the coordinate axes (Figure 14). Direction angles are the positive angles that the vector makes with each of the positive coordinate axes. We calculate these angles as
\[
\cos \alpha = \frac{V_x}{|\mathbf{V}|}, \quad \cos \beta = \frac{V_y}{|\mathbf{V}|}, \quad \cos \gamma = \frac{V_z}{|\mathbf{V}|}
\]  
(12)
The values \( \cos \alpha \), \( \cos \beta \), and \( \cos \gamma \) are called the direction cosines of the vector. Actually, we need specify only two of the direction cosines to give the direction of \( \mathbf{V} \), because
\[
\cos^2 \alpha + \cos^2 \beta + \cos^2 \gamma = 1
\]  
(13)
Vectors are used to represent any quantities that have the properties of magnitude and direction. Two common examples are force and velocity (Figure 15). A force can be thought of as an amount of push or pull along a particular direction. A velocity vector specifies how fast (speed) an object is moving in a certain direction.

Vector Addition and Scalar Multiplication
By definition, the sum of two vectors is obtained by adding corresponding components:
\[
\mathbf{V}_1 + \mathbf{V}_2 = (V_{1x} + V_{2x}, V_{1y} + V_{2y}, V_{1z} + V_{2z})
\]  
(14)
Two-dimensional vector addition is illustrated geometrically in Figure 16. We obtain the vector sum by placing the start position of one vector at the tip of the other vector and drawing the representation for the vector sum from the start of the first vector to the tip of the second. Addition of a vector with a scalar is undefined, because a scalar has only one numerical value while a vector has \( n \) numerical components in an \( n \)-dimensional space.

Multiplication of a vector by a scalar value \( s \) is defined as
\[
s \mathbf{V} = (sV_x, sV_y, sV_z)
\]  
(15)
For example, if the scalar parameter \( s \) has the value 2, each component of \( \mathbf{V} \) is doubled and its magnitude is doubled.

We can also combine vectors using multiplicative processes in various ways. One highly useful method is to multiply the magnitudes of two vectors so that this product is used to form either another vector or a scalar quantity.
Scalar Product of Two Vectors

We obtain a scalar value from two vectors with the calculation

$$V_1 \cdot V_2 = |V_1||V_2| \cos \theta, \quad 0 \leq \theta \leq \pi$$ (16)

where $\theta$ is the smaller of the two angles between the vector directions (Figure 17). This multiplication scheme is called the scalar product, or dot product, of two vectors. It is also referred to as the inner product, particularly in discussing scalar products in tensor analysis. Equation 16 is valid in any coordinate representation and can be interpreted as the product of the parallel components of the two vectors, where $|V_2| \cos \theta$ is the projection of vector $V_2$ in the direction of $V_1$.

In addition to the coordinate-independent form of the scalar product, we can express this calculation in specific coordinate representations. For a Cartesian reference frame, the scalar product is calculated as

$$V_1 \cdot V_2 = V_{1x}V_{2x} + V_{1y}V_{2y} + V_{1z}V_{2z}$$ (17)

The scalar product is a generalization of the Pythagorean theorem, and the scalar product of a vector with itself produces the square of the vector magnitude. Also, the scalar product of two vectors is zero if and only if the two vectors are perpendicular (orthogonal).

Dot products are commutative:

$$V_1 \cdot V_2 = V_2 \cdot V_1$$ (18)

because this operation produces a scalar. And dot products are distributive with respect to vector addition:

$$V_1 \cdot (V_2 + V_3) = V_1 \cdot V_2 + V_1 \cdot V_3$$ (19)

Vector Product of Two Vectors

We use the following calculation to combine two vectors to produce another vector:

$$V_1 \times V_2 = u|V_1||V_2| \sin \theta, \quad 0 \leq \theta \leq \pi$$ (20)

Parameter $u$ in this expression is a unit vector (magnitude 1) that is perpendicular to both $V_1$ and $V_2$ (Figure 18). The direction for $u$ is determined by the right-hand rule: We grasp an axis that is perpendicular to the plane containing $V_1$ and $V_2$ so that the fingers of the right hand curl from $V_1$ to $V_2$. Vector $u$ is then in the direction of the right thumb. This calculation is called the vector product, or cross product, of two vectors, and Equation 20 is valid in any coordinate representation. The cross product of two vectors is a vector that is perpendicular to the plane of the two vectors, and the magnitude of the cross-product vector is equal to the area of the parallelogram formed by the two vectors.

We can also express the cross product in terms of vector components in a specific reference frame. In a Cartesian-coordinate system, we calculate the components of the cross product as

$$V_1 \times V_2 = (V_{1y}V_{2z} - V_{1z}V_{2y}, V_{1z}V_{2x} - V_{1x}V_{2z}, V_{1x}V_{2y} - V_{1y}V_{2x})$$ (21)

If we denote the unit vectors (magnitude 1) along the $x$, $y$, and $z$ axes as $u_x, u_y,$ and $u_z$, we can write the cross product in terms of Cartesian components using a determinant notation (Section 5):

$$V_1 \times V_2 = \begin{vmatrix} u_x & u_y & u_z \\ V_{1x} & V_{1y} & V_{1z} \\ V_{2x} & V_{2y} & V_{2z} \end{vmatrix}$$ (22)
The cross product of any two parallel vectors is zero. Therefore, the cross product of a vector with itself is zero. Also, the cross product is not commutative; it is anticommutative:

$$V_1 \times V_2 = -(V_2 \times V_1) \tag{23}$$

And the cross product is not associative; that is,

$$V_1 \times (V_2 \times V_3) \neq (V_1 \times V_2) \times V_3 \tag{24}$$

But the cross product is distributive with respect to vector addition or subtraction:

$$V_1 \times (V_2 + V_3) = (V_1 \times V_2) + (V_1 \times V_3) \tag{25}$$

### 3 Tensors

A generalization of the concept of a vector is the class of objects called tensors. Formally, a **tensor** is defined as a quantity with a specified **rank** and with certain transformation properties when the tensor is converted from one coordinate representation to another. For orthogonal-coordinate systems, the transformation properties are straightforward and the same as those for vectors. Various physical properties of objects, such as stress, strain, and conductivity, are tensors.

The rank of a tensor, along with the dimension of the space in which the tensor is defined, determines the number of components (also called **elements** or **coefficients**) in that tensor. Scalar quantities and vectors are special cases of the more general class of tensors. A scalar is a tensor of rank zero, and a vector is a tensor of rank one. Basically, the rank of a tensor specifies the number of subscripts used to designate the tensor elements, and the spatial dimension determines the number of values that can be assigned to each subscript. Thus, a scalar quantity (tensor of rank zero) has zero subscripts, and a vector (tensor of rank one) has one subscript. Sometimes any parameter with one subscript is mislabeled as “one-dimensional,” and any parameter with two subscripts is mislabeled as “two-dimensional.” However, the **dimension** of a quantity depends on the spatial representation, not on the number of subscripts. In two-dimensional space, the single vector subscript can be assigned two values and the two-dimensional vector has two components. In three-dimensional space, the single vector subscript can be assigned three values and the three-dimensional vector has three components. Similarly, a tensor of rank two has two subscripts, and in three-dimensional space, this tensor has nine components (three values for each subscript).

### 4 Basis Vectors and the Metric Tensor

We can specify the coordinate directions for an \( n \)-dimensional reference frame using a set of axis vectors, labeled \( \mathbf{u}_k \), where \( k = 1, 2, \ldots, n \), as in Figure 19, which illustrates the axis vectors at the origin of a three-dimensional curvilinear space. Each coordinate-axis vector gives the direction for one of the spatial axes at any point along that axis. These axis tangent vectors form a linearly independent vector set. That is, the axis vectors cannot be written as linear combinations of each other. Also, any other vector in that space can be written as a linear combination of the axis vectors, and the set of axis vectors is called a **basis**, or a set of **base vectors**, for the space. In general, the space is referred to as a vector space and the basis contains the minimum number of vectors needed to represent any other vector in the space as a linear combination of the base vectors.

![Figure 19](Image)
Determining Basis Vectors for a Coordinate Space

The basis vectors in any space are determined from the position vector \( \vec{r} \), which is the vector representation for any spatial position. For example, in three-dimensional Cartesian space, the position vector for any point \((x, y, z)\) is

\[ \vec{r} = x \mathbf{u}_x + y \mathbf{u}_y + z \mathbf{u}_z \]  

(26)

where \( \mathbf{u}_x \), \( \mathbf{u}_y \), and \( \mathbf{u}_z \) are the unit base vectors for the \( x \), \( y \), and \( z \) axes. Unlike other coordinate representations, Cartesian basis vectors are constants that are independent of the spatial coordinates, so that we have

\[ \mathbf{u}_x = \frac{\partial \vec{r}}{\partial x}, \quad \mathbf{u}_y = \frac{\partial \vec{r}}{\partial y}, \quad \mathbf{u}_z = \frac{\partial \vec{r}}{\partial z} \]  

(27)

Similarly, for any other three-dimensional space, we formulate the expression for the position vector \( \vec{r}(x_1, x_2, x_3) \) in terms of the coordinates for that space, and then we determine the basis vectors as

\[ \vec{u}_k = \frac{\partial \vec{r}}{\partial x_k}, \quad k = 1, 2, 3 \]  

(28)

In general, the base vectors \( \vec{u}_k \) are neither constants nor unit vectors. They are functions of the spatial coordinates.

As an example, the position vector in two-dimensional polar-coordinate space is

\[ \vec{r} = r \cos \theta \mathbf{u}_x + r \sin \theta \mathbf{u}_y \]  

(29)

and the polar-coordinate basis vectors are

\[ \vec{u}_r = \frac{\partial \vec{r}}{\partial r} = \cos \theta \mathbf{u}_x + \sin \theta \mathbf{u}_y \]

\[ \vec{u}_\theta = \frac{\partial \vec{r}}{\partial \theta} = -r \sin \theta \mathbf{u}_x + r \cos \theta \mathbf{u}_y \]  

(30)

In this space, \( \vec{u}_r \), which is a function of \( \theta \), is a unit vector. But \( \vec{u}_\theta \), which is a function of both \( r \) and \( \theta \), is not a unit vector.

Orthonormal Basis

Often, vectors in a basis are normalized so that each vector has a magnitude of 1. We obtain unit basis vectors in any three-dimensional space with the calculations

\[ \mathbf{u}_k = \frac{\vec{u}_k}{|\vec{u}_k|}, \quad k = 1, 2, 3 \]  

(31)

and this set of unit vectors is called a normal basis. Also, for Cartesian, cylindrical, spherical, and other commonly used reference frames, including polar coordinates, the coordinate axes are mutually perpendicular at each point in space, and the set of base vectors is then referred to as an orthogonal basis. A set of unit, orthogonal base vectors is called an orthonormal basis, and these base vectors satisfy the following conditions:

\[ \mathbf{u}_k \cdot \mathbf{u}_k = 1, \quad \text{for all } k \]

\[ \mathbf{u}_j \cdot \mathbf{u}_k = 0, \quad \text{for all } j \neq k \]  

(32)

Although we primarily deal with orthogonal systems, nonorthogonal coordinate reference frames are useful in some applications, including relativity theory and visualization schemes for certain data sets.
A two-dimensional Cartesian system has the orthonormal basis
\[ u_x = (1, 0), \quad u_y = (0, 1) \] (33)
And the orthonormal basis for a three-dimensional Cartesian reference frame is
\[ u_x = (1, 0, 0), \quad u_y = (0, 1, 0), \quad u_z = (0, 0, 1) \] (34)

**Metric Tensor**

For “ordinary” coordinate spaces (that is, those in which we can define distances, which are formally referred to as Riemannian spaces), the scalar products of the basis vectors form the elements of the **metric tensor** for that space:
\[ g_{jk} = \vec{u}_j \cdot \vec{u}_k \] (35)
Thus, the metric tensor is of rank two and it is symmetric: \( g_{jk} = g_{kj} \). Metric tensors have several useful properties. The elements of a metric tensor can be used to determine (1) the distance between two points in that space, (2) the transformation equations for conversion to another space, and (3) the components of various differential vector operators (such as gradient, divergence, and curl) within that space.

In an orthogonal space,
\[ g_{jk} = 0, \quad \text{for} \ j \neq k \] (36)
For example, in a Cartesian coordinate system, where the basis vectors are constant unit vectors, the metric tensor has the components
\[ g_{jk} = \begin{cases} 1, & \text{if} \ j = k \\ 0, & \text{otherwise} \end{cases} \] (37)
And for the polar-coordinate basis vectors (Eqs. 30), we can write the metric tensor in the matrix form
\[ g = \begin{bmatrix} 1 & 0 & 0 \\ 0 & r^2 \sin^2 \phi & 0 \\ 0 & 0 & 1 \end{bmatrix} \] (polar coordinates) (38)

For a cylindrical-coordinate reference frame, the base vectors are
\[ \vec{u}_\rho = \cos \theta \, u_x + \sin \theta \, u_y, \quad \vec{u}_\phi = -\rho \sin \theta \, u_x + \rho \cos \theta \, u_y, \quad \vec{u}_z = u_z \] (39)
And the matrix representation for the metric tensor in cylindrical coordinates is
\[ g = \begin{bmatrix} 1 & 0 & 0 \\ 0 & \rho & 0 \\ 0 & 0 & 1 \end{bmatrix} \] (cylindrical coordinates) (40)

In spherical coordinates, the basis vectors are
\[ \vec{u}_r = \cos \theta \sin \phi \, u_x + \sin \theta \sin \phi \, u_y + \cos \phi \, u_z \]
\[ \vec{u}_\theta = -r \sin \theta \sin \phi \, u_x + r \cos \theta \sin \phi \, u_y \]
\[ \vec{u}_\phi = r \cos \theta \cos \phi \, u_x + r \sin \theta \cos \phi \, u_y - r \sin \phi \, u_z \] (41)
Using these base vectors in Equation 35, we obtain the following matrix representation for the metric tensor:
\[ g = \begin{bmatrix} 1 & 0 & 0 \\ 0 & r^2 \sin^2 \phi & 0 \\ 0 & 0 & r^2 \end{bmatrix} \] (spherical coordinates) (42)
5 Matrices

A matrix is a rectangular array of quantities (numerical values, expressions, or functions), which are called the elements of the matrix. Some examples of matrices are

\[
\begin{bmatrix}
3.60 & -0.01 & 2.00 \\
-5.46 & 0.00 & 1.63 \\
\end{bmatrix},
\begin{bmatrix}
e^x & x \\
e^{2x} & x^2 \\
\end{bmatrix},
\begin{bmatrix}
a_1 & a_2 & a_3 \\
x & y & z \\
\end{bmatrix}
\]

(43)

We identify matrices according to the number of rows and number of columns. For the preceding examples, the matrices in left-to-right order are \(2 \times 3\), \(2 \times 2\), \(1 \times 3\), and \(3 \times 1\). When the number of rows is the same as the number of columns, as in the second example, the matrix is called a square matrix.

In general, we can write an \(r \times c\) matrix as

\[
M = \begin{bmatrix}
m_{11} & m_{12} & \cdots & m_{1c} \\
m_{21} & m_{22} & \cdots & m_{2c} \\
\vdots & \vdots & \ddots & \vdots \\
m_{r1} & m_{r2} & \cdots & m_{rc} \\
\end{bmatrix}
\]

(44)

where \(m_{jk}\) represent the elements of matrix \(M\). The first subscript of any element gives the row number, and the second subscript gives the column number.

A matrix with a single row or a single column represents a vector. Thus, the last two matrix examples in 43 are, respectively, a row vector and a column vector. In general, a matrix can be viewed as a collection of row vectors or as a collection of column vectors.

When various operations are expressed in matrix form, the standard mathematical convention is to represent a vector with a column matrix. Following this convention, we write the matrix representation for a three-dimensional vector in Cartesian coordinates as

\[
V = \begin{bmatrix}
v_x \\
v_y \\
v_z \\
\end{bmatrix}
\]

(45)

Although we use this standard matrix representation for both points and vectors, there is an important distinction between the two. The vector representation for a point always assumes that the vector is from the origin to that point. And the distance of the point from the origin is not invariant when we switch from one coordinate system to another. Also, we cannot “add” points, and we cannot apply vector operations, such as the dot product and cross product, to points.

Scalar Multiplication and Matrix Addition

To multiply a matrix \(M\) by a scalar value \(s\), we multiply each element \(m_{jk}\) by the scalar. As an example, if

\[
M = \begin{bmatrix}
1 & 2 & 3 \\
4 & 5 & 6 \\
\end{bmatrix}
\]

then

\[
3M = \begin{bmatrix}
3 & 6 & 9 \\
12 & 15 & 18 \\
\end{bmatrix}
\]
Matrix addition is defined only for matrices that have the same number of rows $r$ and the same number of columns $c$. For any two $r \times c$ matrices, the sum is obtained by adding corresponding elements. For example,

\[
\begin{bmatrix}
1 & 2 & 3 \\
4 & 5 & 6
\end{bmatrix}
\begin{bmatrix}
0.0 & 1.5 & 0.2 \\
-6.0 & 1.1 & -10.0
\end{bmatrix}
= 
\begin{bmatrix}
1.0 & 3.5 & 3.2 \\
-2.0 & 6.1 & -4.0
\end{bmatrix}
\]

**Matrix Multiplication**

The product of two matrices is defined as a generalization of the vector dot product. We can multiply an $m \times n$ matrix $A$ by a $p \times q$ matrix $B$ to form the matrix product $AB$, providing that the number of columns in $A$ is equal to the number of rows in $B$. In other words, we must have $n = p$. We then obtain the product matrix by forming sums of the products of the elements in the row vectors of $A$ with the corresponding elements in the column vectors of $B$. Thus, for the following product:

\[
C = AB
\]

we obtain an $m \times q$ matrix $C$, whose elements are calculated as

\[
c_{ij} = \sum_{k=1}^{n} a_{ik}b_{kj}
\]

In the following example, a $3 \times 2$ matrix is postmultiplied by a $2 \times 2$ matrix to produce a $3 \times 2$ product matrix:

\[
\begin{bmatrix}
0 & -1 \\
5 & 7 \\
-2 & 8
\end{bmatrix}
\begin{bmatrix}
1 & 2 \\
3 & 4
\end{bmatrix}
= 
\begin{bmatrix}
0 \cdot 1 + (-1) \cdot 3 & 0 \cdot 2 + (-1) \cdot 4 \\
5 \cdot 1 + 7 \cdot 3 & 5 \cdot 2 + 7 \cdot 4 \\
-2 \cdot 1 + 8 \cdot 3 & -2 \cdot 2 + 8 \cdot 4
\end{bmatrix}
= 
\begin{bmatrix}
-3 & -4 \\
26 & 38 \\
22 & 28
\end{bmatrix}
\]

Vector multiplication in matrix notation produces the same result as the dot product, provided that the first vector is expressed as a row vector and the second vector is expressed as a column vector. For example,

\[
\begin{bmatrix}
1 & 2 & 3
\end{bmatrix}
\begin{bmatrix}
4 \\
5 \\
6
\end{bmatrix}
= [32]
\]

This vector product results in a matrix with a single element (a $1 \times 1$ matrix). However, if we multiply the vectors in reverse order, we obtain the following $3 \times 3$ matrix:

\[
\begin{bmatrix}
4 \\
5 \\
6
\end{bmatrix}
\begin{bmatrix}
1 & 2 & 3
\end{bmatrix}
= 
\begin{bmatrix}
4 & 8 & 12 \\
5 & 10 & 15 \\
6 & 12 & 18
\end{bmatrix}
\]

As the previous two vector products illustrate, matrix multiplication, in general, is not commutative. That is,

\[
AB \neq BA
\]

But matrix multiplication is distributive with respect to matrix addition:

\[
A(B + C) = AB + AC
\]
Matrix Transpose

The transpose $M^T$ of a matrix is obtained by interchanging rows and columns. For example,

$$
\begin{bmatrix}
1 & 2 & 3 \\
4 & 5 & 6
\end{bmatrix}^T = 
\begin{bmatrix}
1 & 4 \\
2 & 5 \\
3 & 6
\end{bmatrix},
\begin{bmatrix}
a & b & c
\end{bmatrix}^T = 
\begin{bmatrix}
a \\
b \\
c
\end{bmatrix}
$$

(50)

For a matrix product, the transpose is

$$(M_1M_2)^T = M_2^T M_1^T
$$

(51)

Determinant of a Matrix

If we have a square matrix, we can combine the matrix elements to produce a single number called the determinant of the matrix. Determinant evaluations are useful in analyzing and solving a wide range of problems. For a $2 \times 2$ matrix $A$, the second-order determinant is defined to be

$$\det A = \begin{vmatrix} a_{11} & a_{12} \\ a_{21} & a_{22} \end{vmatrix} = a_{11}a_{22} - a_{12}a_{21}
$$

(52)

Higher-order determinants are obtained recursively from lower-order determinant values. To calculate a determinant of order 2 or greater, we can select any column $k$ of an $n \times n$ matrix and compute the determinant as

$$\det A = \sum_{j=1}^{n} (-1)^{i+k} a_{jk} \det A_{jk}
$$

(53)

where $\det A_{jk}$ is the $(n-1) \times (n-1)$ determinant of the submatrix obtained from $A$ by deleting the $j$th row and the $k$th column. Alternatively, we can select any row $j$ and calculate the determinant as

$$\det A = \sum_{k=1}^{n} (-1)^{j+k} a_{jk} \det A_{jk}
$$

(54)

Evaluating determinants for large matrices ($n > 4$, say) can be accomplished more efficiently using numerical methods. One way to compute a determinant is to decompose the matrix into two factors: $A = LU$, where all elements of matrix $L$ above the diagonal are zero, and all elements of matrix $U$ below the diagonal are zero. We then compute the product of the diagonals for both $L$ and $U$, and we obtain $\det A$ by multiplying the two diagonal products. This method is based on the following property of determinants:

$$\det(AB) = (\det A)(\det B)
$$

(55)

Another numerical method for calculating determinants is based on the Gaussian-elimination procedures discussed in Section 14.

Matrix Inverse

With square matrices, we can obtain an inverse matrix if and only if the determinant of the matrix is nonzero. If an inverse exists, the matrix is said to be a nonsingular matrix. Otherwise, the matrix is called a singular matrix. For most practical applications, where a matrix represents a physical operation, we can expect the inverse to exist.

The inverse of an $n \times n$ (square) matrix $M$ is denoted as $M^{-1}$ and

$$MM^{-1} = M^{-1}M = I
$$

(56)
where $I$ is the identity matrix. All diagonal elements of $I$ have the value 1, and all other (off-diagonal) elements are zero.

Elements for the inverse matrix $M^{-1}$ can be calculated from the elements of $M$ as

$$m^{-1}_{jk} = \frac{(-1)^{j+k} \det M_{kj}}{\det M} \tag{57}$$

where $m^{-1}_{jk}$ is the element in the $j$th row and $k$th column of $M^{-1}$, and $M_{kj}$ is the $(n - 1) \times (n - 1)$ submatrix obtained by deleting the $k$th row and $j$th column of matrix $M$. For large values of $n$, we can compute values for the determinants and the elements of the inverse matrix more efficiently using numerical methods.

6 Complex Numbers

By definition, a complex number $z$ is an ordered pair of real numbers, represented as

$$z = (x, y) \tag{58}$$

where $x$ is called the real part of $z$ and $y$ is called the imaginary part of $z$. Real and imaginary parts of a complex number are designated as

$$x = \text{Re}(z), \quad y = \text{Im}(z) \tag{59}$$

Geometrically, a complex number can be described as a point in the complex plane, as illustrated in Figure 20.

When $\text{Re}(z) = 0$, the complex number $z$ is said to be a pure imaginary number. Similarly, any real number can be represented as a complex number with $\text{Im}(z) = 0$. Thus, we can write any real number in the form

$$x = (x, 0)$$

Complex numbers arise from solutions of equations such as

$$x^2 + 1 = 0, \quad x^2 - 2x + 5 = 0$$

which have no real-number solutions. Thus, the concept of a complex number and the rules for complex arithmetic have been devised as extensions of real number manipulations that provide solutions to such problems.

Basic Complex Arithmetic

Addition, subtraction, and scalar multiplication of complex numbers are carried out using the same rules as for two-dimensional vectors. For example, the sum of two complex numbers is

$$z_1 + z_2 = (x_1, y_1) + (x_2, y_2) = (x_1 + x_2, y_1 + y_2)$$

![Complex Number Diagram](image-url)
and we can express any complex number as the summation:

\[ z = (x, y) = (x, 0) + (0, y) \]

The product of two complex numbers, \( z_1 \) and \( z_2 \), is defined as

\[ (x_1, y_1)(x_2, y_2) = (x_1x_2 - y_1y_2, x_1y_2 + x_2y_1) \]  \((60)\)

This definition for complex multiplication gives the same result as for real-number multiplication when the imaginary parts are zero:

\[ (x_1, 0)(x_2, 0) = (x_1x_2, 0) \]

**Imaginary Unit**

The pure imaginary number with \( y = 1 \) is called the **imaginary unit**, and it is denoted as

\[ i = (0, 1) \]  \((61)\)

(Electrical engineers often use the symbol \( j \) for the imaginary unit, because the symbol \( i \) is used to represent electrical current.)

From the rule for complex multiplication, we have

\[ i^2 = (0, 1)(0, 1) = (-1, 0) \]

Therefore, \( i^2 \) is the real number \(-1\), and

\[ i = \sqrt{-1} \]  \((62)\)

We can represent a pure imaginary number using either of the following two forms:

\[ z = iy = (0, y) \]

And a general complex number can be expressed in the form

\[ z = x + iy \]  \((63)\)

Using the definition for \( i \), we can verify that this representation satisfies the rules for complex addition, subtraction, and multiplication.

**Complex Conjugate and Modulus of a Complex Number**

Another concept associated with a complex number is the **complex conjugate**, which is defined to be

\[ \overline{z} = x - iy \]  \((64)\)

Thus, the complex conjugate \( \overline{z} \) is the reflection of \( z \) about the \( x \) (real) axis.

The **modulus**, or absolute value, of a complex number is defined as

\[ |z| = \sqrt{z\overline{z}} = \sqrt{x^2 + y^2} \]  \((65)\)

This number gives the distance in the complex plane of point \( z \) from the origin, which is sometimes referred to as the “vector length” of the complex number. Therefore, the absolute value of a complex number is simply a representation for the Pythagorean theorem in the complex plane.

**Complex Division**

To evaluate the ratio of two complex numbers, we can simplify the expression by multiplying the numerator and denominator by the complex conjugate of the denominator. Then we use the multiplication rules to determine the values for
the components of the resulting complex number. Thus, the real and imaginary parts for the ratio of two complex numbers are obtained as

\[
\begin{align*}
\frac{z_1}{z_2} &= \frac{z_1\overline{z_2}}{z_2\overline{z_2}} \\
&= \frac{(x_1, y_1)(x_2, -y_2)}{x_2^2 + y_2^2}, \\
&= \left( \frac{x_1x_2 + y_1y_2}{x_2^2 + y_2^2}, \frac{x_2y_1 - x_1y_2}{x_2^2 + y_2^2} \right)
\end{align*}
\]

Polar-Coordinate Representation for a Complex Number

Multiplication and division operations for complex numbers are greatly simplified if we express the real and imaginary parts in terms of polar coordinates (Figure 21):

\[
z = r(\cos \theta + i \sin \theta)
\]

We can also write the polar form of \( z \) as

\[
z = re^{i\theta}
\]

where \( e \) is the base of the natural logarithms \( e \approx 2.718281828 \), and

\[
e^{i\theta} = \cos \theta + i \sin \theta
\]

which is Euler’s formula.

Using the polar-coordinate form, we compute the product of two complex numbers by multiplying their absolute values and adding their polar angles. Thus,

\[
z_1z_2 = r_1r_2e^{i(\theta_1 + \theta_2)}
\]

To divide one complex number by another, we divide their absolute values and subtract the polar angles:

\[
\frac{z_1}{z_2} = \frac{r_1}{r_2}e^{i(\theta_1 - \theta_2)}
\]

We can also use the polar representation to obtain roots of complex numbers. The \( n \)th roots of a complex number are calculated as

\[
\sqrt[n]{z} = \sqrt[n]{r} \left[ \cos \left( \frac{\theta + 2k\pi}{n} \right) + i \sin \left( \frac{\theta + 2k\pi}{n} \right) \right], \quad k = 0, 1, 2, \ldots, n - 1
\]

These roots lie on a circle of radius \( \sqrt[n]{r} \) with center at the origin of the complex plane, and they form the vertices for a regular polygon with \( n \) sides.

7 Quaternions

Complex number concepts are extended to higher dimensions using quaternions, which are quantities with one real part and three imaginary parts, written as

\[
q = s + ia + jb + kc
\]
where the coefficients $a$, $b$, and $c$ in the imaginary terms are real numbers, and parameter $s$ is a real number called the \textit{scalar part}. Parameters $i$, $j$, $k$ are defined with the properties
\begin{equation}
  i^2 = j^2 = k^2 = -1, \quad ij = -ji = k
\end{equation}
From these properties, it follows that
\begin{equation}
  jk = -kj = i, \quad ki = -ik = j
\end{equation}
Scalar multiplication is defined in analogy with the corresponding operations for vectors and complex numbers. That is, each of the four components of the quaternion is multiplied by the scalar value. Similarly, quaternion addition is defined as the addition of corresponding elements:
\begin{equation}
  q_1 + q_2 = (s_1 + s_2) + i(a_1 + a_2) + j(b_1 + b_2) + k(c_1 + c_2)
\end{equation}
Multiplication of two quaternions is carried out using the operations in Equations 74 and 75.
We can also use the following ordered-pair notation for a quaternion, which is similar to the ordered-pair representation for a complex number:
\begin{equation}
  q = (s, v)
\end{equation}
Parameter $v$ in this representation is the vector $(a, b, c)$. Using the ordered-pair notation, we can express quaternion addition in the form
\begin{equation}
  q_1 + q_2 = (s_1 + s_2, v_1 + v_2)
\end{equation}
We can write the expression for quaternion multiplication relatively compactly in terms of the vector dot product and cross product operations as
\begin{equation}
  q_1 q_2 = (s_1 s_2 - v_1 \cdot v_2, s_1 v_2 + s_2 v_1 + v_1 \times v_2)
\end{equation}
The magnitude squared of a quaternion is defined by analogy with complex-number operations, using the following sum of the squares of the quaternion components.
\begin{equation}
  |q|^2 = s^2 + v \cdot v
\end{equation}
And the inverse of a quaternion is evaluated using the expression
\begin{equation}
  q^{-1} = \frac{1}{|q|^2} (s, -v)
\end{equation}
so that
\begin{equation}
  qq^{-1} = q^{-1}q = (1, 0)
\end{equation}

8 \textbf{Nonparametric Representations}
When we write object descriptions directly in terms of the coordinates for the reference frame in use, the representation is called \textit{nonparametric}. For example, we can describe a surface with either of the following Cartesian functions.
\begin{equation}
  f_1(x, y, z) = 0, \quad \text{or} \quad z = f_2(x, y)
\end{equation}
The first form in 82 is called an \textit{implicit} expression for the surface, and the second form is called an \textit{explicit} representation. In the explicit representation, $x$ and $y$ are referred to as the \textit{independent variables}, and $z$ is called the \textit{dependent variable}. 

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Similarly, we can represent a three-dimensional curved line in nonparametric form as the intersection of two surface functions, or we could represent the curve with the pair of functions

\[ y = f(x) \quad \text{and} \quad z = g(x) \] (83)

with coordinate \( x \) as the independent variable. Values for the dependent variables \( y \) and \( z \) are then determined from Equations 83 as we step through values for \( x \) for some prescribed number of intervals.

Nonparametric representations are useful in describing objects within a given reference frame, but they have some disadvantages when used in graphics algorithms. If we want a smooth plot, we must change the independent variable whenever the first derivative (slope) of either \( f(x) \) or \( g(x) \) becomes greater than 1. This requires continual checks on the derivative values to determine when we need to change the roles of the independent and dependent variables. Also, Equations 83 provide an awkward format for representing multiple-valued functions. For instance, the implicit equation for a circle centered on the origin in the \( xy \) plane is

\[ x^2 + y^2 - r^2 = 0 \]

and the explicit expression for \( y \) is the multivalued function

\[ y = \pm \sqrt{r^2 - x^2} \]

In general, a more convenient representation for object descriptions in graphics algorithms is in terms of parametric equations.

9 Parametric Representations

We can classify objects according to the number of parameters needed to describe the coordinate positions on the objects. A curve, for example, in a Cartesian reference frame is classified as a one-dimensional Euclidean object, and a surface is a two-dimensional Euclidean object. When an object description is given in terms of its dimensionality parameter, the description is called a parametric representation.

The Cartesian description for positions along the path of a curve can be given in a parametric form using the following vector point function:

\[ \mathbf{P}(u) = (x(u), y(u), z(u)) \] (84)

where each of the Cartesian coordinates is a function of parameter \( u \). In most cases, we can normalize the three coordinate functions so that parameter \( u \) varies over the range from 0 to 1.0. For example, a circle in the \( xy \) plane with radius \( r \) and center position at the coordinate origin can be defined in parametric form with the following three functions:

\[ x(u) = r \cos(2\pi u), \quad y(u) = r \sin(2\pi u), \quad z(u) = 0, \quad 0 \leq u \leq 1 \] (85)

Because this curve is defined in the \( xy \) plane, we could eliminate the \( z(u) \) function, which has the constant value 0.

In a similar way, we can represent coordinate positions on a surface using the following Cartesian vector point function:

\[ \mathbf{P}(u, v) = (x(u, v), y(u, v), z(u, v)) \] (86)
Each of the Cartesian coordinates is now a function of the two surface parameters $u$ and $v$. A spherical surface with radius $r$ and center at the coordinate origin, for example, can be described with the equations

$$
\begin{align*}
  x(u, v) &= r \cos(2\pi u) \sin(\pi v) \\
y(u, v) &= r \sin(2\pi u) \sin(\pi v) \\
z(u, v) &= r \cos(\pi v)
\end{align*}
$$

(87)

Parameter $u$ describes lines of constant longitude over the surface, while parameter $v$ describes lines of constant latitude. The parametric equations are again normalized so that $u$ and $v$ are assigned values in the range from 0 to 1.0. By keeping one of these parameters fixed while varying the other over a subrange of the unit interval, we could plot latitude and longitude lines for any spherical section (Figure 22).

### 10 Rate-of-Change Operators

For a continuous function of a single independent variable, such as $f(x)$, we determine the rate at which the function is changing at any particular $x$ value using a function called the derivative of $f(x)$ with respect to $x$. This derivative function is defined as

$$\frac{df}{dx} = \lim_{\Delta x \to 0} \frac{f(x + \Delta x) - f(x)}{\Delta x} \quad (88)$$

and this definition is the basis for obtaining numerical solutions for problems involving rate-of-change operations. The functional forms for the derivatives of commonly occurring functions, such as polynomials and trigonometric functions, are available in derivative tables. And for rate-of-change problems involving simple functions, we can typically obtain closed-form solutions. But, in many cases, we need to solve rate-of-change problems using numerical methods.

When we have a function of several variables, rate-of-change operations with respect to the individual variables are called partial derivatives. For example, with a function such as $f(x, y, z, t)$, we can determine the rate of change of the function with respect to any one of the coordinate directions, $x$, $y$, or $z$, or the time parameter $t$. A partial derivative for a particular independent variable is defined with Equation 88, where all other independent variables are held constant. Thus, for example, the partial derivative of $f$ with respect to time is defined as

$$\frac{\partial f}{\partial t} = \lim_{\Delta t \to 0} \frac{f(x, y, z, t + \Delta t) - f(x, y, z, t)}{\Delta t}$$

which is evaluated at some particular spatial position and time.

A number of partial-derivative operators occur frequently enough that they are given special names, such as gradient, Laplacian, divergence, and curl. These operators are useful in various applications, such as determining the geometry and orientation of objects, describing the behavior of objects in certain situations, calculating electromagnetic-radiation effects, and analyzing data sets in scientific visualization studies.

#### Gradient Operator

The vector operator with the following Cartesian components is called the gradient operator:

$$\nabla = \left( \frac{\partial}{\partial x}, \frac{\partial}{\partial y}, \frac{\partial}{\partial z} \right) \quad (89)$$
Symbol $\nabla$ is referred to as nabla, del, or simply the grad operator. One important use for the gradient operator is in calculating a surface normal vector. When a surface is described with the nonparametric representation $f(x, y, z) = \text{constant}$, the surface normal at any position is calculated as

$$N = \nabla f$$  \hspace{1cm} \text{(Normal Vector for a Nonparametric Surface Representation)} \hspace{1cm} (90)$$

As an example, a spherical surface with radius $r$ can be represented in local coordinates with the nonparametric Cartesian representation $f(x, y, z) = r^2$, and the gradient of $f$ produces the surface normal vector $(2x, 2y, 2z)$. But if a surface is represented with a parametric point function $P(u, v)$, then we can determine the surface normal using the vector cross-product calculation

$$N = \frac{\partial P}{\partial u} \times \frac{\partial P}{\partial v}$$  \hspace{1cm} \text{(Normal Vector for a Parametric Surface Representation)} \hspace{1cm} (91)$$

Directional Derivative

We can also use the gradient operator and the vector dot product to form a scalar product called the directional derivative of a function $f$:

$$\frac{\partial f}{\partial u} = \mathbf{u} \cdot \nabla f$$  \hspace{1cm} (92)$$

This gives us the rate-of-change of $f$ in a direction specified by the unit vector $\mathbf{u}$. To illustrate, we can determine the directional derivative for the spherical surface function $f = x^2 + y^2 + z^2$ in the $z$ direction as

$$\frac{\partial f}{\partial z} = \mathbf{u}_z \cdot \nabla f = 2z$$

where $\mathbf{u}_z$ is the unit vector along the positive $z$ direction. And for the following unit vector in the $xy$ plane

$$\mathbf{u} = \frac{1}{\sqrt{2}} \mathbf{u}_x + \frac{1}{\sqrt{2}} \mathbf{u}_y$$

the directional derivative of $f$ from Equation 92 is

$$\frac{\partial f}{\partial u} = \frac{1}{\sqrt{2}} \frac{\partial f}{\partial x} + \frac{1}{\sqrt{2}} \frac{\partial f}{\partial y} = \sqrt{2}x + \sqrt{2}y$$

General Form of the Gradient Operator

Within any three-dimensional, orthogonal-coordinate system, we obtain the components for the gradient operator using the calculations

$$\nabla = \sum_{k=1}^{3} \frac{\mathbf{u}_k}{\sqrt{g_{kk}}} \frac{\partial}{\partial x_k}$$  \hspace{1cm} (93)$$

In this expression, each $\mathbf{u}_k$ represents the unit basis vector in the $x_k$-coordinate direction, and $g_{kk}$ are the diagonal components of the metric tensor for the space.

Laplace Operator

We can use the gradient operator and the vector dot product to form a scalar differential operator called the Laplacian or the Laplace operator, which has the Cartesian-coordinate form

$$\nabla^2 = \nabla \cdot \nabla = \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} + \frac{\partial^2}{\partial z^2}$$  \hspace{1cm} (94)$$
The symbol $\nabla^2$ is often referred to as *grad squared*, *del squared*, or *nabla squared*. And in any three-dimensional, orthogonal-coordinate system, the Laplacian of a function $f(x, y, z)$ is computed as

$$\nabla^2 f = \frac{1}{\sqrt{g_{11}g_{22}g_{33}}} \left[ \frac{\partial}{\partial x_1} \left( \sqrt{g_{22}g_{33}} \frac{\partial f}{\partial x_1} \right) + \frac{\partial}{\partial x_2} \left( \sqrt{g_{33}g_{11}} \frac{\partial f}{\partial x_2} \right) + \frac{\partial}{\partial x_3} \left( \sqrt{g_{11}g_{22}} \frac{\partial f}{\partial x_3} \right) \right]$$  \hspace{1cm} (95)

Equations involving the Laplacian arise in many applications, including the description of electromagnetic-radiation effects.

**Divergence Operator**

The vector dot product can also be used to combine the gradient operator with a vector function to produce a scalar quantity called the *divergence of a vector*, which has the following Cartesian form:

$$\text{div} \mathbf{V} = \nabla \cdot \mathbf{V} = \frac{\partial V_x}{\partial x} + \frac{\partial V_y}{\partial y} + \frac{\partial V_z}{\partial z}$$  \hspace{1cm} (96)

In this expression, $V_x$, $V_y$, and $V_z$ are the Cartesian components of the vector $\mathbf{V}$. Divergence is a measure of the rate of increase or decrease of a vector function, such as an electric field, at a point in space. In any three-dimensional, orthogonal-coordinate system, the divergence of a vector $\mathbf{V}$ is calculated as

$$\text{div} \mathbf{V} = \nabla \cdot \mathbf{V} = \frac{1}{\sqrt{g_{11}g_{22}g_{33}}} \left[ \frac{\partial}{\partial x_1} (\sqrt{g_{22}g_{33}} V_1) + \frac{\partial}{\partial x_2} (\sqrt{g_{33}g_{11}} V_2) + \frac{\partial}{\partial x_3} (\sqrt{g_{11}g_{22}} V_3) \right]$$  \hspace{1cm} (97)

with parameters $V_1$, $V_2$, and $V_3$ as the components of vector $\mathbf{V}$ with respect to the coordinate directions $x_1$, $x_2$, and $x_3$, and $g_{kk}$ are the diagonal elements of the metric tensor.

**Curl Operator**

Another very useful differential operator is the *curl of a vector*, which is applied using the gradient operator and the vector cross product. The Cartesian components for the curl of a vector are

$$\text{curl} \mathbf{V} = \nabla \times \mathbf{V} = \left( \frac{\partial V_z}{\partial y} - \frac{\partial V_y}{\partial z}, \frac{\partial V_x}{\partial z} - \frac{\partial V_z}{\partial x}, \frac{\partial V_y}{\partial x} - \frac{\partial V_x}{\partial y} \right)$$  \hspace{1cm} (98)

This operation gives us a measure of rotational effects associated with a vector quantity, such as in the scattering of electromagnetic radiation. For any three-dimensional, orthogonal-coordinate system, we can express the components of the curl in terms of the metric tensor components using the following determinant representation:

$$\text{curl} \mathbf{V} = \nabla \times \mathbf{V} = \frac{1}{\sqrt{g_{11}g_{22}g_{33}}} \left| \begin{array}{ccc} \sqrt{g_{11}} \mathbf{u}_1 & \sqrt{g_{22}} \mathbf{u}_2 & \sqrt{g_{33}} \mathbf{u}_3 \\ \frac{\partial}{\partial x_1} & \frac{\partial}{\partial x_2} & \frac{\partial}{\partial x_3} \\ \sqrt{g_{11}} V_1 & \sqrt{g_{22}} V_2 & \sqrt{g_{33}} V_3 \end{array} \right|$$  \hspace{1cm} (99)

Vectors $\mathbf{u}_k$ are the unit basis vectors for the space, and variables $g_{kk}$ are the diagonal elements of the metric tensor.

### 11 Rate-of-Change Integral Transformation Theorems

In many applications, we encounter problems that involve rate-of-change operations that are to be integrated (summed) over some region of space, which can be along a line path, across a surface, or throughout a volume of space. Often the
problem can be simplified by applying a transformation theorem that converts a surface integral to a line integral, a line integral to a surface integral, a volume integral to a surface integral, or a surface integral to a volume integral. These transformation theorems are of tremendous importance in solving a wide range of practical problems.

**Stokes’s Theorem**

For a continuous vector function \( \mathbf{F}(x, y, z) \) defined over some surface region, Stokes’s theorem states that the integral of the perpendicular component of the curl of \( \mathbf{F} \) is equal to the line integral of \( \mathbf{F} \) around the perimeter curve \( C \) for the surface. That is,

\[
\int \int_{\text{surf}} (\text{curl} \ \mathbf{F}) \cdot \mathbf{n} \, dA = \oint_{C} \mathbf{F} \cdot \mathbf{r} \, ds
\]

(100)

where the boundary \( C \) must be “piecewise smooth,” which means that \( C \) must be a continuous curve or a curve composed of a finite number of continuous sections, such as circular arcs or straight-line segments. In this expression, \( \mathbf{n} \) is the unit surface normal at any point, \( dA \) is a differential element of surface area, \( \mathbf{r} \) is a unit tangent vector to the boundary curve \( C \) at any point, and \( ds \) is a differential line segment along \( C \). The integration direction around \( C \) is counterclockwise when we view the front of the surface, as shown in Figure 23.

**Green’s Theorem for a Plane Surface**

If we consider a region of the \( xy \) plane bounded by a piecewise smooth curve \( C \) (as in Stokes’s theorem), we can express Green’s plane theorem in the Cartesian form

\[
\int \int_{\text{area}} \left( \frac{\partial f_2}{\partial x} - \frac{\partial f_1}{\partial y} \right) \, dx \, dy = \oint_{C} (f_1 \, dx + f_2 \, dy)
\]

(101)

Here, \( f_1(x, y) \) and \( f_2(x, y) \) are two continuous functions defined throughout the planar area bounded by curve \( C \), and the direction of integration around \( C \) is counterclockwise. We can also apply Green’s theorem to a region with internal holes, as in Figure 24, but we then must integrate in a clockwise direction around the interior boundary curves.

**Figure 23**
Integration around the boundary curve \( C \) is counterclockwise in Stokes’s theorem when we view the surface from the “outside” region of space.

**Figure 24**
Line integrals in Green’s plane theorem are evaluated by traversing the boundary curves \( C_1 \) and \( C_2 \) so that the interior region (shaded) is always on the left.
Although developed independently, Green’s plane theorem is a special case of Stokes’s theorem. To demonstrate this, we define a vector function \( \mathbf{F} \) with Cartesian components \((f_1, f_2, 0)\). Then, Green’s theorem can be written in the vector form

\[
\iint_{\text{area}} (\nabla \times \mathbf{F}) \cdot \mathbf{u}_z \, dA = \oint_{\mathcal{C}} \mathbf{F} \cdot \mathbf{r} \, ds
\]  

where \( \mathbf{u}_z \) is the unit vector perpendicular to the \( xy \) plane (in the \( z \) direction), \( dA = dx \, dy \), and the other parameters are the same as in Equation 100.

We can use Green’s plane theorem to compute the area of a planar region by setting \( f_1 = 0 \) and \( f_2 = x \). Then, from Green’s theorem, the area \( A \) of a plane figure is

\[
A = \iint_{\text{area}} dx \, dy = \oint_{\mathcal{C}} x \, dy
\]  

Similarly, if we set \( f_1 = -y \) and \( f_2 = 0 \), we have

\[
A = \iint_{\text{area}} dx \, dy = -\oint_{\mathcal{C}} y \, dx
\]  

Adding the two previous area equations, we obtain

\[
A = \frac{1}{2} \oint_{\mathcal{C}} (x \, dy - y \, dx)
\]

We can also convert this Cartesian expression for the area into the following polar-coordinate form:

\[
A = \frac{1}{2} \oint_{\mathcal{C}} r^2 \, d\theta
\]

Green’s plane theorem can be expressed in many other useful forms. For example, if we define \( f_1 = \partial f / \partial y \) and \( f_2 = \partial f / \partial x \), for some continuous function \( f \), then we have

\[
\iint_{\text{area}} \nabla^2 f \, dx \, dy = \oint_{\mathcal{C}} \frac{\partial f}{\partial n} \, ds
\]

where \( \partial f / \partial n \) is the directional derivative of \( f \) in the direction of the outward normal to the boundary curve \( \mathcal{C} \).

**Divergence Theorem**

The previous two theorems give us methods for converting between surface integrals and line integrals. The **divergence theorem** provides an equation for converting a volume integral into a surface integral, or vice versa. This theorem is also known by various other names, including **Green’s theorem in space** and **Gauss’s theorem**. For a continuous, three-dimensional, vector function \( \mathbf{F} \), defined over a volume of space, we can express the divergence theorem in the vector form

\[
\iiint_{\text{vol}} \text{div} \, \mathbf{F} \, dV = \iint_{\text{surf}} \mathbf{F} \cdot \mathbf{n} \, dA
\]

where \( dV \) is a differential volume element, \( \mathbf{n} \) is the normal vector for the bounding surface, and \( dA \) is a differential element of surface area.

We can use the divergence theorem to obtain several other useful integral transformations. For instance, if \( \mathbf{F} = \nabla f \) for some continuous three-dimensional
function $f$, we have the volume version of Equation 107, which is

$$\iiint_{\text{vol}} \nabla^2 f \, dV = \iint_{\text{surf}} \frac{\partial f}{\partial n} \, dA \quad (109)$$

In this equation, $\frac{\partial f}{\partial n}$ is the directional derivative of $f$ in the direction of the surface normal.

From the divergence theorem, we can derive expressions for calculating the volume of a spatial region using a surface integral. Depending upon how we represent the vector function $\mathbf{F}$, we can obtain any one of the following Cartesian forms for the surface integral:

$$V = \iiint_{\text{vol}} \, dx \, dy \, dz = \iint_{\text{surf}} y \, dz \, dx = \iint_{\text{surf}} z \, dx \, dy = \frac{1}{3} \iint_{\text{surf}} (x \, dy \, dz + y \, dz \, dx + z \, dx \, dy) \quad (110)$$

Green’s Transformation Equations

A number of other integral transformations can be derived from the divergence theorem. The following two integral equations are generally referenced as Green’s transformation equations, Green’s first and second formulas, or Green’s identities:

$$\iiint_{\text{vol}} (f_1 \nabla^2 f_2 + \nabla f_1 \cdot \nabla f_2) \, dV = \iint_{\text{surf}} f_1 \frac{\partial f_2}{\partial n} \, dA \quad (111)$$

$$\iiint_{\text{vol}} (f_1 \nabla^2 f_2 - f_2 \nabla^2 f_1) \, dV = \iint_{\text{surf}} \left( f_1 \frac{\partial f_2}{\partial n} - f_2 \frac{\partial f_1}{\partial n} \right) \, dA \quad (112)$$

In these equations, $f_1$ and $f_2$ are continuous, three-dimensional scalar functions, and $\frac{\partial f_1}{\partial n}$ and $\frac{\partial f_2}{\partial n}$ are their directional derivatives in the direction of the surface normal.

12 Area and Centroid of a Polygon

We can use the integral transformations from Section 11 to calculate various properties of objects for computer-graphics applications. For polygons, we often use the area and the centroid coordinates in programs involving geometric transformations, simulations, system design, and animations.

Area of a Polygon

From Equation 103, we can compute the area of a polygon by expressing the Cartesian coordinates in parametric form and evaluating the line integral around the perimeter of the polygon. The parametric equations for the $n$ edges of a polygon with $n$ vertices in the $xy$ plane (Figure 25) can be expressed in the form

$$x = x_k + (x_{k+1} - x_k)u$$
$$y = y_k + (y_{k+1} - y_k)u$$

where $x_{n+1} = x_1$ and $y_{n+1} = y_1$. 

**Figure 25**
A polygon defined with $n$ vertices in the $xy$ plane.
Substituting the differential expression \(dy = (y_{k+1} - y_k)\ du\) and the parametric expression for \(x\) into Equation 103, we have

\[
A = \oint_C x \, dy
\]

\[
= \sum_{k=1}^{n} \int_0^1 \left[ x_k + (x_{k+1} - x_k)u \right](y_{k+1} - y_k) \, du
\]

\[
= \sum_{k=1}^{n} (y_{k+1} - y_k) \left[ x_k + (x_{k+1} - x_k)/2 \right]
\]

\[
= \frac{1}{2} \sum_{k=1}^{n} (x_k y_{k+1} - x_k y_k + x_{k+1} y_{k+1} - x_{k+1} y_k)
\]

(114)

For each line segment, the second and third terms in this sum are canceled by similar terms with opposite signs in the expressions for successive values of \(k\). Therefore, the area of the polygon is computed with:

\[
A = \frac{1}{2} \sum_{k=1}^{n} (x_k y_{k+1} - x_k y_k)
\]

(115)

**Centroid of a Polygon**

By definition, the centroid is the position of the center of mass for a constant-density object (all points in the object have the same mass). Thus, the coordinates for the centroid are simply the mean values for the coordinates over all positions within the object boundaries.

For some simple polygon shapes, we can obtain the centroid by averaging the vertex positions. But, in general, vertex averaging does not locate the centroid correctly, because it neglects the positions for all other points in the polygon. As illustrated in Figure 26, the average vertex position is near the greatest concentration of vertices, while the centroid is at the center position for the entire polygon area.

We calculate the centroid position \((\bar{x}, \bar{y})\) for a polygon in the \(xy\) plane by averaging the coordinates for all positions within the polygon boundaries:

\[
\bar{x} = \frac{1}{A} \int \int_{\text{area}} x \, dxdy = \frac{\mu_x}{A}
\]

\[
\bar{y} = \frac{1}{A} \int \int_{\text{area}} y \, dxdy = \frac{\mu_y}{A}
\]

(116)

In these expressions, \(\mu_x\) and \(\mu_y\) are called the moments of the area with respect to the \(x\) and \(y\) axes, respectively, where the area is assumed to have unit mass per unit area.

**Figure 26**

Coordinate positions for the centroid of a polygon and the average of the vertex coordinates.
We can evaluate each of the moments of the polygon using the same procedures we employed to compute the area of the polygon. From Green’s plane theorem, we obtain a line integral equivalent to the area integral, and we evaluate the line integral using parametric representations for the Cartesian coordinates along the polygon edges. Green’s theorem for a plane surface (Eq. 101) states that

$$\int \int_{\text{area}} \left( \frac{\partial f_2}{\partial x} - \frac{\partial f_1}{\partial y} \right) \, dx \, dy = \oint_C (f_1 \, dx + f_2 \, dy) \quad (117)$$

For the evaluation of $\mu_x$, we can take $f_2 = \frac{1}{2}x^2$ and $f_1 = 0$ in the preceding transformation, so that

$$\mu_x = \int \int_{\text{area}} x \, dx \, dy = \frac{1}{2} \oint_C x^2 \, dy \quad (118)$$

From the parametric representations 113 for the polygon edges, we have

$$x^2 = x_k^2 + 2x_k(x_{k+1} - x_k)u + (x_{k+1} - x_k)^2u^2$$

and

$$dy = (y_{k+1} - y_k) \, du$$

for each of the $n$ edges, labeled $k = 1, 2, \ldots, n$. Therefore,

$$\mu_x = \frac{n}{2} \int_0^1 \left[ x_k^2 + 2x_k(x_{k+1} - x_k)u + (x_{k+1} - x_k)^2u^2 \right] \, du$$

$$= \frac{1}{6} \sum_{k=1}^n (x_{k+1} + x_k)(x_k y_{k+1} - x_{k+1}y_k) \quad (119)$$

For the evaluation of $\mu_y$, we make the substitutions $f_1 = -\frac{1}{2}y^2$ and $f_2 = 0$ in Green’s theorem, and we obtain

$$\mu_y = \int \int_{\text{area}} y \, dx \, dy = \frac{1}{2} \oint_C y^2 \, dx \quad (120)$$

Using the parametric representations in 113 to evaluate the line integral, we have

$$\mu_y = \frac{1}{6} \sum_{k=1}^n (y_{k+1} + y_k)(x_k y_{k+1} - x_{k+1}y_k) \quad (121)$$

Given any set of polygon vertices, we then use the expressions for $A, \mu_x,$ and $\mu_y$ in Equations 116 to compute the polygon centroid coordinates. Because the expression $(x_k y_{k+1} - x_{k+1}y_k)$ appears in the calculations for all three quantities, $A$, $\mu_x$, and $\mu_y$, we compute this once for each line segment.

### 13 Calculating Properties of Polyhedra

Methods similar to those for polygons are used to obtain polyhedra properties. But now we compute the spatial volume, instead of an area, and the centroid is obtained by averaging the coordinate positions throughout the volume of a polyhedron.

The volume of any spatial region is defined in Cartesian coordinates as

$$V = \int \int \int_{\text{vol}} dx \, dy \, dz \quad (122)$$
This integral can be converted to a surface integral using one of the transformation equations 110. For a polyhedron, the surface integral can then be evaluated using a parametric representation for positions across each face of the solid.

We calculate the centroid positions for polyhedra using methods similar to those for polygons. By definition, the Cartesian-coordinate centroid position for a region of space (with unit mass per unit volume) is the average of all positions within the region:

\[
\begin{align*}
\mathbf{x} &= \frac{1}{V} \iiint_{\text{vol}} x \, dx \, dy \, dz = \frac{\mu_x}{V} \\
\mathbf{y} &= \frac{1}{V} \iiint_{\text{vol}} y \, dx \, dy \, dz = \frac{\mu_y}{V} \\
\mathbf{z} &= \frac{1}{V} \iiint_{\text{vol}} z \, dx \, dy \, dz = \frac{\mu_z}{V}
\end{align*}
\] (123)

Again, we can convert the volume integrals to surface integrals, substitute parametric representations for the Cartesian coordinates, and evaluate the surface integrals over the faces of a polyhedron.

14 Numerical Methods

In computer-graphics algorithms, it is often necessary to solve sets of linear equations, nonlinear equations, integral equations, and other functional forms. Also, to visualize a discrete set of data points, it may be useful to display a continuous curve or surface function that approximates the points of the data set. In this section, we briefly summarize some common algorithms for solving various numerical problems.

Solving Sets of Linear Equations

For variables \(x_k\), with \(k = 1, 2, \ldots, n\), we can write a system of \(n\) linear equations as

\[
\begin{align*}
a_{11}x_1 + a_{12}x_2 + \cdots + a_{1n}x_n &= b_1 \\
a_{21}x_1 + a_{22}x_2 + \cdots + a_{2n}x_n &= b_2 \\
& \vdots \\
a_{n1}x_1 + a_{n2}x_2 + \cdots + a_{nn}x_n &= b_n
\end{align*}
\] (124)

where the values for parameters \(a_{jk}\) and \(b_j\) are known. This set of equations can be expressed in the matrix form

\[
AX = B
\] (125)

with \(A\) as an \(n \times n\) square matrix whose elements are the coefficients \(a_{jk}\), \(X\) as the column matrix of \(x_j\) values, and \(B\) as the column matrix of \(b_j\) values. Solving this matrix equation for \(X\), we obtain

\[
X = A^{-1}B
\] (126)

This system of equations can be solved if and only if \(A\) is a nonsingular matrix; that is, its determinant is nonzero. Otherwise, the inverse of matrix \(A\) does not exist.

One method for solving the set of equations is Cramer’s rule:

\[
x_k = \frac{\det A_k}{\det A}
\] (127)

where \(A_k\) is the matrix \(A\) with the \(k\)th column replaced with the elements of \(B\). This method is adequate for problems with a few variables. For more than three or four variables, however, the method is extremely inefficient due to the large
number of multiplications needed to evaluate each determinant. Evaluation of a single $n \times n$ determinant requires more than $n!$ multiplications.

We can solve the system of equations more efficiently using variations of **Gaussian elimination**. The basic ideas in Gaussian elimination can be illustrated with the following set of two simultaneous equations:

\[
\begin{align*}
    x_1 + 2x_2 &= -4 \\
    3x_1 + 4x_2 &= 1
\end{align*}
\]  

(128)

To solve this set of equations, we can multiply the first equation by $-3$, then add the two equations to eliminate the $x_1$ term, yielding the equation

\[-2x_2 = 13\]

which has the solution $x_2 = -13/2$. This value is then substituted into either of the original equations to obtain the solution for $x_1$, which is $9$. We can use this basic approach to solve any set of linear equations, but algorithms have been devised to perform the elimination and back substitution steps more efficiently.

A modification of Gaussian elimination is the **LU decomposition** (or **LU factorization**) method for solving sets of linear equations. In this algorithm, we first factor matrix $A$ into two matrices, called a lower-diagonal matrix $L$ and an upper-diagonal matrix $U$, such that

\[
A = LU
\]  

(129)

All elements of matrix $L$ above its diagonal have the value $0$, and all diagonal elements have the value $1$. All elements of matrix $U$ below the diagonal have the value $0$. We can then write Equation 125 as

\[
LUX = B
\]  

(130)

This allows us to solve the following two very much simpler sets of equations:

\[
LY = B, \quad UX = Y
\]  

(131)

Once we have the values for the elements of matrix $Y$ in Equation 131, we use these in the second set of equations to solve for the elements of matrix $X$. As an example, the following equation demonstrates the factorization for a $2 \times 2$ coefficient matrix:

\[
A = \begin{bmatrix} 2 & 3 \\ 8 & 5 \end{bmatrix} = \begin{bmatrix} 1 & 0 \\ 4 & 1 \end{bmatrix} \begin{bmatrix} 2 & 3 \\ 0 & -7 \end{bmatrix}
\]

One method for computing the elements of the factorization matrices is given in the following set of equations, where $u_{ij}$ are the elements for the upper-triangular matrix $U$ and $l_{ij}$ are the elements for the lower-triangular matrix $L$.

\[
\begin{align*}
    u_{1j} &= a_{1j}, \quad j = 1, 2, \ldots, n \\
    l_{i1} &= \frac{a_{i1}}{u_{11}}, \quad i = 2, 3, \ldots, n \\
    u_{ij} &= a_{ij} - \sum_{k=1}^{i-1} l_{ik}u_{kj} \quad j = i, i + 1, \ldots, n; \quad i \geq 2 \\
    l_{ij} &= \frac{1}{u_{jj}} \left( a_{ij} - \sum_{k=1}^{j-1} l_{ik}u_{kj} \right) \quad i = j + 1, j + 2, \ldots, n; \quad j \geq 2
\end{align*}
\]  

(132)

Gaussian elimination is sometimes susceptible to high round-off errors, and other methods might not produce an accurate solution. In such cases, we may be able to obtain a solution using the **Gauss-Seidel method**. This method is also an efficient way to solve the set of linear equations when we know the approximate...
In the Gauss-Seidel approach, we start with an initial "guess" for the values of variables $x_k$, then repeatedly calculate successive approximations until the difference between two successive values for each $x_k$ is small. At each step, we calculate the approximate values for the variables as

$$
x_1 = \frac{b_1 - a_{12}x_2 - a_{13}x_3 - \cdots - a_{1n}x_n}{a_{11}}$
$$
x_2 = \frac{b_2 - a_{21}x_1 - a_{23}x_3 - \cdots - a_{2n}x_n}{a_{22}}$
$$
\vdots
$$

If we can rearrange matrix $A$ so that each diagonal element has a magnitude greater than the sum of the magnitudes of the other elements across that row, than the Gauss-Seidel method is guaranteed to converge to a solution.

Finding Roots of Nonlinear Equations

A root of a function $f(x)$ is a value for $x$ that satisfies the equation $f(x) = 0$. In general, the function $f(x)$ can be an algebraic expression, such as a polynomial, or it can involve transcendental functions. An algebraic expression is one that contains only the arithmetic operators, exponents, roots, and powers. Transcendental functions, such as the trigonometric functions, log functions, and exponential functions, are represented with infinite power series.

Roots of a nonlinear equation can be real numbers, complex numbers, or a combination of real and complex numbers. Sometimes we can obtain exact solutions for all roots, depending on the complexity of the equation. For example, we know how to find an exact solution for any polynomial up to degree 4, and the roots of a simple transcendental equation such as $\sin x = 0$ are known to be $x = k\pi$ for any integer value of $k$. But in most cases of practical interest, we need to apply numerical procedures to obtain the roots of a nonlinear equation.

One of the most popular methods for finding roots of nonlinear equations is the **Newton-Raphson Algorithm**. This is an iterative procedure that approximates $f(x)$ as a linear function at each step of the iteration, as shown in Figure 27. We start with an initial "guess" of $x_0$ for the value of the root, and then calculate the next approximation to the root, $x_1$, by determining where the tangent line from $x_0$ crosses the $x$ axis. At $x_0$, the slope (first derivative) of the curve is

$$
\frac{df}{dx} = \frac{f(x_0)}{x_0 - x_1}
$$

Thus, the next approximation to the root is

$$
x_1 = x_0 - \frac{f(x_0)}{f'(x_0)}
$$

where $f'(x_0)$ denotes the derivative of $f(x)$ evaluated at $x = x_0$. We repeat this procedure at each calculated approximation until the difference between successive approximations is "small enough."

**Figure 27**

Approximating a curve at an initial value $x_0$ with a straight line that is tangent to the curve at that point.
In addition to solving problems involving real variables, the Newton-Raphson algorithm can be applied to a function of a complex variable \( f(z) \), to a function of several variables, and to sets of simultaneous nonlinear functions, real or complex. And, if the Newton-Raphson algorithm converges to a root, it will converge faster than any other root-finding method. But it may not always converge. For example, the method fails if the derivative \( f'(x) \) evaluates to 0 at some point in the iteration. In addition, depending on the oscillations of the curve, successive approximations may diverge from the position of a root.

Another method, slower but guaranteed to converge, is the **bisection method**. In this algorithm, we must identify an \( x \) interval that contains a root. Then we apply a binary-search procedure within that interval to close in on the root. We first look at the midpoint of the interval to determine whether the root is in the lower or upper half of the interval. This procedure is repeated for each successive subinterval until the difference between successive midpoint positions is smaller than some preset value. A speedup can be attained by interpolating successive \( x \) positions instead of halving each subinterval (false-position method).

### Evaluating Integrals

Integration is a summation process. For a function of a single variable \( x \), the integral of \( f(x) \) is equal to the area “under” the curve, as illustrated in Figure 28. For simple integrands, we can often determine a functional form for an integral, but, in general, we evaluate integrals using numerical methods.

From the definition of an integral, we can form the following numerical approximation:

\[
\int_a^b f(x) \, dx \approx \sum_{k=1}^n f_k(x) \Delta x_k
\]  

(136)

The function \( f_k(x) \) is an approximation to \( f(x) \) over the interval \( \Delta x_k \). For example, we can approximate the curve with a constant value in each subinterval and add the areas of the resulting rectangles (Figure 29). This approximation improves, up to a point, as we decrease the size of the subdivisions across the interval from \( a \) to \( b \). If the subdivisions are too small, the values of successive rectangular areas can be lost in the round-off error.

Polynomial approximations for the function in each subinterval generally give better results than the rectangle approach. Using a linear approximation, the
resulting subareas are trapezoids, and the approximation method is then referred to as the **trapezoid rule**. If we use a quadratic polynomial (parabola) to approximate the function in each subinterval, the method is called **Simpson’s rule** and the integral approximation is

\[
\int_a^b f(x) \, dx \approx \frac{\Delta x}{3} \left[ f(a) + f(b) + 4\sum_{\text{odd}k=1}^{n-1} f(x_k) + 2\sum_{\text{even}k=2}^{n-2} f(x_k) \right] \tag{137}
\]

In this expression, the interval from \( a \) to \( b \) is divided into the \( n \) equal-width intervals

\[
\Delta x = \frac{b - a}{n} \tag{138}
\]

where \( n \) is a multiple of 2, and with

\[
x_0 = a, \quad x_k = x_{k-1} + \Delta x, \quad k = 1, 2, \ldots, n
\]

For a function with a rapidly varying amplitude, such as the example in Figure 30, it may be difficult to approximate the function accurately over the subintervals. Also, multiple integrals (involving several integration variables) are not easy to evaluate with Simpson’s rule or the other approximation methods. In these cases, we can apply **Monte Carlo** integration techniques. The term **Monte Carlo** is used to describe any method that employs random-number procedures to solve a deterministic problem.

We apply a Monte Carlo method to evaluate an integral by generating \( n \) random positions within a rectangular area that contains \( f(x) \) over the interval from \( a \) to \( b \) (Figure 31). An approximation for the integral is then calculated as

\[
\int_a^b f(x) \, dx \approx h(b - a) \frac{n_{\text{count}}}{n} \tag{139}
\]

where parameter \( h \) is the rectangle height and parameter \( n_{\text{count}} \) is the number of random points that are between \( f(x) \) and the \( x \) axis. A random position \((x, y)\) in the rectangular region is computed by first generating two random numbers, \( r_1 \) and \( r_2 \), then carrying out the calculations

\[
h = y_{\text{max}} - y_{\text{min}}, \quad x = a + r_1(b - a), \quad y = y_{\text{min}} + r_2 h \tag{140}
\]

Similar methods can be applied to multiple integrals.

In the calculations for \( x \) and \( y \) in 140, we assume that the random numbers \( r_1 \) and \( r_2 \) are uniformly distributed over the interval \((0, 1)\). We can obtain \( r_1 \) and \( r_2 \) from a random-number function in a mathematics or statistical library, or we
can use the following algorithm, called the **linear congruential generator**:

\[ i_k = a \cdot i_{k-1} + c \pmod{m}, \quad k = 1, 2, 3, \ldots \]

\[ r_k = \frac{i_k}{m} \quad (141) \]

where parameters \( a, c, m, \) and \( i_0 \) are integers, and \( i_0 \) is a starting value called the *seed*. Parameter \( m \) is chosen to be as large as possible on a particular machine, with values for \( a \) and \( c \) chosen to make the string of random numbers as long as possible before a value is repeated. For example, on a machine with 32-bit integer representations, we can set \( m = 2^{32} - 1, a = 1664525, \) and \( c = 1013904223. \)

**Solving Ordinary Differential Equations**

Any equation containing differential rate-of-change operators is referred to as a *differential equation*. Quantities can change values in some continuous way from one coordinate position to another. They can also change over time at a fixed position, and they can change with respect to many other parameters, such as temperature or rotational acceleration. An equation that involves derivatives of a function of a single variable is called an *ordinary differential equation*. We solve a differential equation either by determining a functional form that satisfies the equation or by using numerical approximation methods to determine the values for the quantity at selected intervals.

To solve a differential equation, we also need to know one or more starting values. An equation involving only the first derivative of a quantity, called a *first-order differential equation*, requires one starting value. An equation that contains both first and second derivatives, called a *second-order differential equation*, requires two starting values. And similarly for equations involving higher-order derivatives. There are two basic classifications for specifying starting values. An *initial-value problem* is one in which the known conditions are specified for a single value of the independent variable. A *boundary-value problem* is one in which the known conditions are specified at the boundaries for the dependent variable.

A simple example of an initial-value problem is the first-order differential equation

\[ \frac{dx}{dt} = f(x, t), \quad x(t_0) = x_0 \quad (142) \]

where \( x \) represents some dependent variable that varies with time \( t \) (the independent variable), \( f(x, t) \) is the known time-variation function for the first derivative of \( x \), and \( x_0 \) is the given value for \( x \) at the initial time \( t_0 \). We can also write this equation in the form

\[ dx = f(x, t) \, dt \]

From the definition of a derivative, we can use finite intervals to approximate the differentials as

\[ \Delta x_k \approx f(x_k, t_k) \Delta t_k, \quad k = 0, 1, \ldots, n \quad (143) \]

where \( \Delta x_k = x_{k+1} - x_k \) and \( \Delta t_k = t_{k+1} - t_k \), for \( n \) time steps. Typically, we take equal time intervals and use the following incremental calculations to determine the \( x \) values at each time step, given the value of \( x_0 \) at \( t_0 \):

\[ x_{k+1} = x_k + f(x_k, t_k) \Delta t \quad (144) \]

This numerical procedure is called the **Euler method**, and it approximates \( x \) with straight-line segments over each time interval \( \Delta t \).
Although the Euler method is a simple procedure to implement, it is not very accurate. Improvements to this basic numerical algorithm have been developed from the following Taylor series expansion by incorporating higher-order terms into the approximation for the differential equation:

\[ x(t + \Delta t) = x(t) + x'(t)\Delta t + \frac{1}{2} x''(t)\Delta t^2 + \cdots \]  

(145)

Because \( x'(t) = f(x, t) \), we have \( x''(t) = f'(x, t) \), and so forth, for the higher derivatives.

A more accurate, and more widely used, method for evaluating a first-order differential equation is the \textbf{Runge-Kutta algorithm}, also called the \textbf{fourth-order Runge-Kutta algorithm}. This procedure is based on a Taylor series expansion up to fourth order. The algorithm for the Runge-Kutta method is

\[
\begin{align*}
    a &= f(x_k, t_k)\Delta t \\
    b &= f(x_k + a/2, t_k + \Delta t/2)\Delta t \\
    c &= f(x_k + b/2, t_k + \Delta t/2)\Delta t \\
    d &= f(x_k + c, t_k + \Delta t)\Delta t \\
    t_{k+1} &= t_k + \Delta t \\
    x_{k+1} &= x_k + (a + 2b + 2c + d)/6, \quad k = 0, 1, \ldots, n - 1
\end{align*}
\]

(146)

We can apply similar methods to obtain solutions for ordinary differential equations of higher order. The general approach is to use the Taylor series expansion for \( x \) to include terms in \( x'' \), \( x''' \), and so forth, depending on the order of the differential equation. For example, from the Taylor series, we can obtain the following approximation for the second derivative:

\[ x''(t_k) \approx \frac{x_{k+1} - 2x_k + x_{k-1}}{\Delta t^2} \]  

(147)

\textbf{Solving Partial Differential Equations}

As we might expect, partial differential equations are generally more difficult to solve than ordinary differential equations. But we can apply similar methods and replace the partial derivatives with finite differences.

We first consider a function \( f(x, t) \) that depends only on the \( x \) coordinate and time \( t \). We can reduce a partial differential equation involving \( \partial f / \partial x \) and \( \partial f / \partial t \) to an ordinary differential equation by replacing the spatial derivative with finite differences. This allows us to replace the function of two variables with a one-subscript function of a single variable, \( t \):

\[ f(x, t) \rightarrow f(x_k, t) \rightarrow f_k(t) \]  

(148)

The partial derivatives are then replaced with the following expressions:

\[
\begin{align*}
    \frac{\partial f}{\partial x} &= \frac{f_{k+1} - f_k}{\Delta x} \\
    \frac{\partial f}{\partial t} &= \frac{df_k}{dt}
\end{align*}
\]

(149)

Then we solve the equations at a finite number of \( x \) positions using the given initial or boundary conditions.

For higher-order derivatives, we can use Taylor series expansions to obtain the finite-difference approximations. As an example, we can use the following
approximation for the second partial derivative of $f$ with respect to $x$:

$$\frac{\partial^2 f(x, t)}{\partial x^2} \rightarrow \frac{f_{k+1}(t) - 2f_k(t) + f_{k-1}(t)}{\Delta x^2} \quad (150)$$

When we have functions defined over surfaces or volumes of space, we can divide the space into a regular grid and use finite differences for each spatial coordinate.

Another approach that is applied to partial differential equations is the finite-element method. A grid of coordinate positions is set up over the domain of interest, which could be a surface or a volume of space, and then the coupled equations are solved at the node positions using variational techniques. In this method, an approximating functional solution is used instead of finite-difference equations. Depending on the problem, an integral is set up for some quantity such as potential energy or residual error. Then some procedure, such as a least-square analysis, is applied to minimize the potential energy or residual error. This minimization yields values for the unknown parameters in the approximating function for the solution.

**Least-Squares Curve-Fitting Methods for Data Sets**

When a computer simulation or scientific-visualization study produces a set of data values, we almost always want to determine a functional form that will describe the data set. The standard method for producing a function that fits the given data is the **least-squares algorithm**. To apply this method, we first choose a general type for the function, such as a linear function, a polynomial function, or an exponential function. We then must determine the values for the parameters in the functional form that we have chosen. A two-dimensional straight-line function, for example, can be described with two parameters: the slope and the $y$ intercept. We obtain the function parameters by minimizing the sum of the squares of the differences between the theoretical function values and the actual data values.

To illustrate this method, we first consider a two-dimensional set of $n$ data points, labeled $(x_k, y_k)$ with $k = 1, 2, \ldots, n$. After we have selected the functional form $f(x)$ that we want to use to describe the data distribution, we set up an expression for an error function $E$, which is the sum of the squares of the differences between $f(x_k)$ and the data values $y_k$:

$$E = \sum_{k=1}^{n} [y_k - f(x_k)]^2 \quad (151)$$

Parameters in the function $f(x)$ are then determined by minimizing the error expression $E$.

As an example, if the data set is to be described with the linear function

$$f(x) = a_0 + a_1 x$$

then

$$E = \sum_{k=1}^{n} [y_k^2 - 2y_k(a_0 + a_1 x_k) + a_0^2 + 2a_0a_1 x_k + a_1^2 x_k^2] \quad (152)$$

Because the error $E$ is a function of two variables ($a_0$ and $a_1$), we minimize $E$ with the following two coupled equations:

$$\frac{\partial E}{\partial a_0} = \sum_{k=1}^{n} [-2y_k + 2a_0 + 2a_1 x_k] = 0$$

$$\frac{\partial E}{\partial a_1} = \sum_{k=1}^{n} [-2y_k x_k + 2a_0 x_k + 2a_1 x_k^2] = 0 \quad (153)$$
We can then solve this set of two linear equations using Cramer’s rule, which yields

\[
\begin{align*}
    a_0 &= \frac{(\sum_k x_k^2)(\sum_k y_k) - (\sum_k x_k)(\sum_k x_k y_k)}{D} \\
    a_1 &= \frac{n \sum_k x_k y_k - (\sum_k x_k)(\sum_k y_k)}{D}
\end{align*}
\]  

(154)

where the denominator in these two expressions is

\[
D = \left| \begin{array}{cc}
    n & \sum_k x_k \\
    \sum_k x_k & \sum_k x_k^2
\end{array} \right| = \sum_{k=1}^{n} x_k^2 - \left( \sum_{k=1}^{n} x_k \right)^2
\]

(155)

Similar calculations are carried out for other functions. For the polynomial

\[f(x) = a_0 + a_1 x + a_2 x^2 + \cdots + a_n x^n\]

for instance, we need to solve a set of \( n \) linear equations to determine values for parameters \( a_k \). And we can also apply least-squares curve-fitting to functions of several variables \( f(x_1, x_2, \ldots, x_m) \) which could be linear or nonlinear in each of the variables.
Any stored pictorial representation is called a graphics file or an image file. For raster-graphics systems, a color screen display is represented in the frame buffer as a set of red, green, and blue (RGB) pixel values. The contents of the frame buffer, or any rectangular section of it, is called a pixmap. Although monochromatic images can be stored in a bitmap form (using a single bit for each pixel), most raster pictures are now stored as pixmaps. In general, any raster representation for a picture is referred to as a raster file. Many formats have been developed for organizing the information in an image file in various ways, and full-color raster files can be quite large, so most file formats apply some type of compression to reduce the file size, both for archiving and for transmission. In addition, the number of color values in a full-color image file must be reduced when the picture is to be displayed on a system with limited color capabilities, or when the file is to be stored in a format that does not support 24 bits per pixel. Here, we provide a brief introduction to graphics file formats and the commonly used methods for reducing the size of both an image file and the number of colors to be used in the display of an image.

1 Image-File Configurations

Pixel color values in a raster image file are typically stored as nonnegative integers, and the range of color values depends on the number of available bits per pixel position. For a full-color (24 bits per pixel) RGB image, the value for each color component is stored in 1 byte, with $R$, $G$, and $B$ values ranging from 0 to 255. An uncompressed raster-graphics file composed of RGB color values is sometimes referred to as raw data or a raw raster file. Other color models, including HSV, HSB, and $YCbCr$, are used in compressed file formats. And the number of available bits per pixel depends on the format.

File formats typically include a header that provides information about the structure of the file. For compressed files, the header may also contain tables and other details needed to decode and display the compressed image. The header can include a variety of information, such as the file size (number of scan lines and number of pixels per scan line), the number of bits or bytes allocated per pixel, the compression method used to reduce the size of the file, the color range for the pixel values, and the image background color.

Another characteristic of raster image files is the ordering for the bytes within the file. Most computer processors store multibyte integers with the most significant byte first, but some processors store multibyte integers with the least significant byte first. The term big endian is used to refer to the ordering with the most significant byte first, and the term little endian refers to the ordering with the least significant byte first.

Some file formats store a picture in a geometric representation, which is a list of the coordinate positions and other information for straight-line segments, fill areas, circular arcs, spline curves, and other primitives. Geometric representations
can also contain attribute information and viewing parameters. This type of image representation is commonly referred to as a vector format, even though not all geometric structures are defined with straight-line segments. Originally, the term “vector” file was used to describe a list of line segments for display on a vector (random-scan) system. Although vector systems have been replaced by raster systems, and nonlinear object descriptions have been added to the “vector” files, the name continues to be applied to any file using a geometric representation for a picture. File formats that support both geometric and raster image representations are referred to as hybrid formats or metafiles.

Scientific-visualization applications often use an image file that is a set of data values generated from measuring instruments or from numerical computer simulations. Various programs are then used to provide particular data visualizations, such as pseudo-color displays, isosurface representations, or volume renderings.

2 Color-Reduction Methods

Several methods have been devised for reducing the number of colors used in the display of an image. The most popular methods are those that attempt to generate a color sampling that closely approximates the original set of colors.

Sometimes color-reduction methods are referred to as “quantization,” which is a term used in areas of physics and mathematics (such as quantum mechanics and sampling theory) for a process that produces a discrete set of values from a continuous distribution. However, a raster image file is not a continuous distribution; it contains a finite, discrete set of color values. Therefore, any color-reduction method simply replaces one discrete set of colors with a smaller discrete set of colors. Furthermore, the color-reduction processes in common use do not generate a set of colors such that each color in the set is a multiple of some selected value. In other words, color reduction does not produce a set of quantized colors.

Uniform Color Reduction

A simple method for reducing colors in a raster file is to divide each of the R, G, and B color levels by an integer and truncate the result. For example, if we divide by 2, we reduce each of the R, G, and B components in a full-color representation to 128 levels. Thus, uniform color reduction replaces groups of contiguous color levels with a reduced color level, as illustrated in Figure 1.

Another approach is to replace a group of pixel values with the value of the middle pixel in the group. Alternatively, we could replace the group of pixels with the average color for the group.

In general, we can expect that not all 256 values will be present in the image file for each of the RGB components. Therefore, we can apply a uniform color-reduction method to the color levels between the minimum and maximum levels that actually occur in the image file.

We can also apply different reduction criteria to the different RGB components. For instance, we could reduce a full-color image so that the red and green color components are represented with 3 bits each (8 levels) and the blue component is represented with 2 bits (4 levels).

Popularity Color Reduction

Another approach to color reduction is to retain only the color values that occur most frequently in an image representation. We can first process the input image file to reduce the bit representation for the individual RGB components. Then we scan this modified set of colors to produce a count, or histogram, of the frequency
of occurrence for each RGB color component. To produce a reduced color file with $k$ colors, we select the $k$ most frequently occurring colors in the image file.

**Median-Cut Color Reduction**

In this algorithm, we subdivide the color space for the image file into $k$ subregions and calculate the average color for each of the subregions. To form the subregions, we first determine the minimum and maximum values for each of the RGB components: $R_{\min}$, $R_{\max}$, $G_{\min}$, $G_{\max}$, $B_{\min}$, and $B_{\max}$. These values give us the bounds on the block of colors within the RGB color cube that are present in the image. For the largest of these three intervals, we determine the median value and use this value to form two smaller blocks of colors. As an example, if the red component has the largest range, we compute the value $R_{\text{median}}$ such that half of the pixel colors are above this value and half are below. We then slice the image color block into two subblocks at the $R_{\text{median}}$ position, as shown in Figure 2. Each of the two color subblocks is then processed using the same subdivision procedure. This process continues until we have subdivided the original image color block into $k$ subblocks. At each step, we can apply the subdivision procedure to the largest subblock. An average color at the desired precision is calculated for each subblock, and all image colors within a subblock are replaced with the average subblock color.

**3 File-Compression Techniques**

Several compression techniques are available for reducing the number of bytes in an image file, but the effectiveness of a particular compression method depends on the type of image. Simple methods that look for patterns in the image file are most effective with geometric designs that contain large single-color areas, while the more complex compression schemes produce better results with photo-realistic computer-graphics images and digitized photographs. The general technique employed to reduce the size of a graphics file is to replace the color values with an encoding that occupies fewer bytes than the original file. In addition, codes are incorporated into compressed files to indicate such things as the end of a scan line and the end of the image file.

Some compression algorithms involve floating-point operations, which can introduce round-off errors. In addition, some methods use approximations that also modify the image colors. As a result, a file that has been decoded from a compressed file often contains color values that are not exactly the same as in the original image. For instance, an integer RGB color that is specified as (247, 108, 175) in an input image file could become the color (242, 111, 177) after decoding.
the compressed file. But such color changes are often tolerable because our eyes are not sensitive to small color differences.

File-reduction methods that do not change the values in an image file are described as **lossless compression** techniques, and those that create color changes are referred to as **lossy compression** techniques. In most cases, lossy compression methods produce a much greater compression ratio for a file, where the compression ratio is the number of bytes in the original file divided by the number of bytes in the compressed file.

**Run-Length Encoding**

This compression scheme simply searches the image file for contiguous, repeated values. A reduced file is formed by storing each sequence of repeated values as a repetition count (the *run length*) followed by a single copy of the repeated value. For example, if the value 125 occurs eight times in succession along a scan line, we store the two values, 8 and 125, in the compressed file. This reduces the original eight bytes of storage to two bytes. For images with large single-color areas, this encoding scheme works well. But images such as digitized photographs have frequent color changes and few consecutive repeating values, so that many color values would be stored with a repetition factor of 1.

Variations have been developed to improve the efficiency of the basic run-length encoding algorithm. For instance, we could use a negative repetition factor to indicate a sequence of nonrepeating values rather than just storing a repetition factor of 1 with each of the values in the nonrepeating sequence. As an example of this, the following list of values

\{20, 20, 20, 20, 99, 68, 31, 40, 40, 40, 40, 40, 40, 40, 40, ...

could be encoded as

\{4, 20, -3, 99, 68, 31, 8, 40, ...

which indicates that the value 20 occurs four times, followed by the three non-repeating values 99, 68, and 31, which in turn are followed by eight occurrences of the value 40. In this encoding example, the first 15 bytes of the input file are compressed into 8 bytes.

**LZW Encoding**

Developed by Lempel, Ziv, and Welch, the LZW method is a modification of the earlier LZ, LZ77, and LZ78 pattern-recognition algorithms. In the LZW scheme, repeated patterns in an image file are replaced with a code. For instance, the following list of 12 values contains two occurrences for each of the patterns \{128, 96\} and \{200, 30, 10\}:

\{128, 96, 200, 30, 10, 128, 96, 50, 240, 200, 30, 10, ...

We can replace these two patterns with the codes c1 and c2, and the remaining pattern \{50, 240\} can be assign the code c3. This reduces the first 12 values in the input list to the following 5 bytes:

\{c1, c2, c1, c3, c2, ...

Alternatively, any nonrepeating sequence of values, such as \{50, 240\} could be stored in the compressed file without assigning a code to the sequence.
Basically, the LZW algorithm searches for repeated sequences and constructs a table of such sequences along with their assigned codes. Thus, this encoding scheme is called a substitutional algorithm or a dictionary-based algorithm. The compressed file is then decoded from the code table.

Other Pattern-Recognition Compression Methods

We can use pattern-recognition schemes to locate repetitions for particular black-and-white or RGB color combinations throughout an image file. Duplicated scan lines and other patterns can be detected and encoded to reduce further the size of image files. In addition, fractal methods have been applied to obtain small encoded self-similar sets of color values.

Huffman Encoding

File compression is accomplished with the Huffman approach by using a variable-length code for the values in an image file. The Huffman-encoding method assigns the shortest code to the most frequently occurring value in the file, and the longest code is assigned to the least frequently occurring value.

The basic idea in the Huffman algorithm is the same as in the Morse code, which assigns variable-length character codes to letters of the alphabet. High-frequency letters in the Morse scheme are assigned one-character codes, and the lowest-frequency letters are assigned four-character codes. For example, the letter E is coded as a “dot” (·), the letter T is coded as a “dash” (–), and the letter Q is coded as a four-character sequence with one dot and three dashes (– – · –). Instead of using character codes, however, the Huffman code assigns variable-length bit codes to the values in an image file, which provides greater compression ratios.

The first step in the Huffman algorithm is to count the number of occurrences of each value in the input image file. Then, bit codes are assigned to the values according to the frequency count. One method for assigning the variable-length bit codes is to construct a binary tree with the high-frequency values near the top of the tree and the lowest-frequency values as the leaf nodes. Starting with the low-frequency values, we create the subtrees from the bottom up. Each root node of a subtree is assigned a numerical label that is the sum of the frequency counts or node labels of its two children. When the tree is complete, all left subtrees are labeled with the binary value 0, and all right trees are labeled with the binary value 1. The bit code for each file value is then formed by concatenating the branch bit labels from the top of the tree down to the node position of that file value in the tree.

To illustrate the general tree-construction steps, we use the set of six values in Table 1. This set represents a short example image file containing 21 items, with the value 96 occurring eight times, the value 177 occurring four times, and so forth for the other four values in the file.

The values 210 and 43 in this table have the lowest frequency count, so we use these two values to form the first subtree (Figure 3). The root of this subtree is assigned a node label that is equal to the sum of the number of occurrences of its two offspring: \(3 = 2 + 1\). We delete these two file values (210 and 43) from the active list so that the next lowest frequency count is 3. But we just created a subtree that also has the node label 3. Therefore, we can form the next subtree using any two of the three items that have the label 3. We choose the two file values to form the subtree shown in Figure 4, and we delete the values 141 and 85 from the active list. The next subtree is constructed with the file value 177, which has a count of 4, and the subtree whose root has the label 3 (Figure 5). We delete the file value 177 and the tree node with the label 3 from the active list, and now the

<table>
<thead>
<tr>
<th>File Value</th>
<th>Frequency Count</th>
</tr>
</thead>
<tbody>
<tr>
<td>96</td>
<td>8</td>
</tr>
<tr>
<td>177</td>
<td>4</td>
</tr>
<tr>
<td>141</td>
<td>3</td>
</tr>
<tr>
<td>85</td>
<td>3</td>
</tr>
<tr>
<td>210</td>
<td>2</td>
</tr>
<tr>
<td>43</td>
<td>1</td>
</tr>
</tbody>
</table>

Total Values in File: 21
two lowest “counts” in the list represent subtrees. These two subtrees are then merged to produce the new subtree shown in Figure 6. Finally, we complete the construction of the binary tree (Figure 7) by joining the file value 96 to the last subtree we created. The value assigned to the root of the tree is the total count (21) for all values in the image file.

Now that we have all file values in a binary tree, we can label left branches in the tree with the binary value 0 and right branches with the binary value 1, as in Figure 8. Starting at the root of the tree, we concatenate the branch labels down to each of the leaf nodes. This forms the set of variable-length binary codes for each of the file values, and we then set up Table 2, which will be stored with the compressed file. For this example, there is one file value with a one-digit binary code, three file values with a three-digit binary code, and two file values with a four-digit binary code. The low-frequency values have longer codes, and the higher-frequency values have shorter codes.
An important characteristic of the Huffman codes is that no bit code is a prefix for any other bit code. This allows us to decode a list of encoded file values by providing Table 3 along with Table 2. To demonstrate the decoding algorithm, we suppose that the compressed file contains the bit stream \{100100100 \ldots \}. The first bit value in this file is 1, so it must represent the file value 96 because there is a bit code of 1, and this cannot be a prefix for any other code. Next, we have a bit value of 0. There is no one-bit code other than 1 and there are no two-bit codes, so the next code must be either 001 or 0010. Checking the indexed code table, we find a file value 210 with the code 0010, which means that there cannot be a file value with the code 001. At this point, we have decoded the first two file values, 96 and 210. The next code in the bit stream must be either 010 or 0100. There is a file value with the code 010, so there cannot be a four-bit code with that prefix. Thus the third decoded file value is 141. We continue analyzing the bit stream in this manner until the compressed file has been fully decoded.

We can also use other schemes for generating and assigning Huffman bit codes. Once we have the frequency count, we could assign a code length to each file value. Using the code length and the frequency count, we can then use a list-merging algorithm to devise the specific bit codes. A predefined code set can also be used to assign codes to the file values, which eliminates the need to store the codes with the compressed file.

**Arithmetic Encoding**

In this compression scheme, the frequency count in a file is used to obtain numerical codes for sequences of the file values. The arithmetic-encoding algorithm first computes the fraction of the file that is occupied by each value. This creates a set of subintervals within the unit interval from 0.0 to 1.0. Then each file fraction is repeatedly mapped onto these subintervals to establish numerical
TABLE 4

Frequency Count and Fraction of Occurrences for Values in a Small Example File

<table>
<thead>
<tr>
<th>File Value</th>
<th>Frequency Count</th>
<th>File Fraction</th>
<th>Unit-Interval Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>V₁</td>
<td>16</td>
<td>0.20</td>
<td>0.00–0.20</td>
</tr>
<tr>
<td>V₂</td>
<td>24</td>
<td>0.30</td>
<td>0.20–0.50</td>
</tr>
<tr>
<td>V₃</td>
<td>40</td>
<td>0.50</td>
<td>0.50–1.00</td>
</tr>
</tbody>
</table>

Total: 80 1.00

intervals for various combinations of the file values. The numerical bounds for the subintervals are used to encode these combinations.

To illustrate the method, we consider a file with 80 entries and just three distinct values. The frequency count and corresponding file fractions for the three values are listed in Table 4. Thus, value $V₁$ is associated with the subinterval from 0.00 to 0.20 within the unit interval, value $V₂$ is associated with the subinterval from 0.20 to 0.50, and value $V₃$ is associated with the subinterval from 0.50 to 1.00.

In other words, 20 percent of the unit interval is associated with $V₁$, 30 percent with $V₂$, and 50 percent with $V₃$. If we now map $V₁$ onto the $V₃$ subinterval, it will occupy 20 percent of the top half of the unit interval. This new subinterval, with a range from 0.50 to 0.60, represents the sequence $V₃ V₁$. Similar results are obtained for the mappings of $V₂$ and $V₃$ onto the $V₃$ subinterval. Table 5 lists ranges for these three two-value sequences. Continuing in this manner, we can map the intervals for the two-value sequences onto other subintervals to obtain the sequences for longer combinations of the file values. The boundary values for the subintervals are then used to encode and decode the sequences within the file.

Various algorithms can be used to terminate the unit-interval subdivisions and to assign numerical codes to the file-value combinations. And the arithmetic-encoding algorithm is typically implemented using binary numbers instead of the floating-point values within the unit interval. The compressed file is then a sequence of binary values.

**Discrete Cosine Transform**

A number of numerical transform methods, including the Fourier and Hadamard transforms, have been applied to file compression, but the discrete cosine transform is the most commonly used method. Efficient implementation algorithms for the discrete cosine transform provide faster execution and better color fidelity in a reconstructed picture at higher compression ratios.

For a list of $n$ numerical values $Vₖ$, with $k = 0, 1, \ldots, n - 1$, the discrete cosine method generates the following set of transformed values:

$$V'_j = c_j \sum_{k=0}^{n-1} Vₖ \cos \left( \frac{(2k + 1)j\pi}{2n} \right), \quad \text{for } j = 0, 1, \ldots, n - 1$$

(1)

where

$$c_j = \begin{cases} 
\frac{1}{\sqrt{n}}, & \text{for } j = 0 \\
\frac{\sqrt{2}}{n}, & \text{for } j \neq 0
\end{cases}$$
Thus, this transform method computes a discrete sum of cosine terms with increasing frequency and with amplitudes that are proportional to the input values. Except for possible round-off errors, the original values are recovered with the inverse transformation

$$V_k = \sum_{j=0}^{n-1} c_j V'_j \cos \left[ \frac{(2k + 1)j\pi}{2n} \right], \quad \text{for } k = 0, 1, \ldots, n - 1 \quad (2)$$

Very often, the transform values $V'_j$ are alluded to as the “coefficients” of the cosine functions in the inverse transform equation. But this is incorrect terminology because the coefficients of the cosine terms in the summation are the products $c_j V'_j$.

To illustrate this transform method, we consider the following list of 8 input values:

$$\{215, 209, 211, 207, 192, 148, 88, 63\}$$

The transformed values, computed to two decimal places, for this input are

$$\{471.29, 143.81, -67.76, 16.33, 7.42, -4.73, 5.49, 0.05\}$$

In this example, we note that the amplitudes of the transformed values markedly decrease, so the higher frequency cosine terms contribute less to the recovery of the input values. This is a basic characteristic of the discrete cosine transform, which allows us to approximate closely the original values using only the first several transformed values. Therefore, to obtain a compressed image file, we could calculate and store just the first half or so of the transformation values. Table 6 shows the results from Equation 2 when we use four, five, or all eight of the transformed values to regain the input values. All calculated values in the table are rounded to two decimal places.

We can improve the efficiency of this compression technique by transforming rectangular blocks of input values, rather than transforming linear sets of values across a single scan line. For a square block of $n \times n$ input values, the transformed values are calculated as

$$V'_{lm} = c_{lm} \sum_{j=0}^{n-1} \sum_{k=0}^{n-1} V_{jk} \cos \left[ \frac{(2k + 1)l\pi}{2n} \right] \cos \left[ \frac{(2j + 1)m\pi}{2n} \right]$$

(3)

with

$$l, m = 0, 1, \ldots, n - 1$$

### Table 6

**Inverse Discrete Cosine Transformation Calculations**

<table>
<thead>
<tr>
<th>Terms in Sum</th>
<th>Inverse Discrete Cosine Transform Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>212.63, 211.85, 211.53, 207.42, 188.43, 147.65, 95.47, 58.02</td>
</tr>
<tr>
<td>5</td>
<td>215.26, 209.23, 208.91, 210.04, 191.06, 145.02, 92.84, 60.64</td>
</tr>
<tr>
<td>8</td>
<td>215.00, 209.00, 211.00, 207.00, 192.00, 148.00, 88.00, 63.00</td>
</tr>
</tbody>
</table>
and

\[
c_{lm} = \begin{cases} 
\frac{1}{n}, & \text{if } l = 0 \text{ or } m = 0 \\
\frac{2}{n}, & \text{if } l \neq 0 \text{ and } m \neq 0 
\end{cases}
\]

Also, the \(n \times n\) set of input values are recovered using the inverse transformation:

\[
V_{jk} = \sum_{l=0}^{n-1} \sum_{m=0}^{n-1} c_{jm} V'_{lj} \cos \left( \frac{(2j+1)l\pi}{2n} \right) \cos \left( \frac{(2k+1)m\pi}{2n} \right) \tag{4}
\]

where

\[
j, k = 0, 1, \ldots, n - 1
\]

This transform and its inverse are typically implemented using \(8 \times 8\) groups of input values, so that groups of color values along eight scan lines are processed simultaneously.

4 Composition of the Major File Formats

Hundreds of file formats have been developed for representing graphical data within different contexts for different systems. Operating systems, for example, typically use a number of specially designed formats within the various system-processing routines. And individual formats exist for specific applications, such as three-dimensional modeling, animations, graphical user interfaces, ray-tracing software, video recording, scientific-visualization software, paint programs, word-processing systems, spreadsheet packages, Internet communications, television broadcasting, and fax transmissions. In addition, the International Standards Organization (ISO) and American National Standards Institute (ANSI) standards committees have proposed several formats and file-compression systems for general use.

Most raster file formats are designed to accommodate color images, but some apply only to bitmaps. However, the format name is often misleading, because the term bitmap is frequently used to reference color images (pixmaps). This situation is simply a result of the continued use of the older label, “bitmap,” for a raster file. Before the development of color displays, all raster images were stored as bitmaps (1 bit per pixel), representing the black-and-white pixel patterns in a picture. As color techniques developed, pixmap files (multiple bits per pixel) replaced the bitmaps. But very often these files were still referred to as bitmaps. As a result, many color-encoding schemes in use today for image files are labeled as “bitmap formats,” even though they are actually pixmap formats (multiple bits per pixel). However, the documentation for such formats can be consulted to determine the number of bits actually allotted to each pixel position in the file.

For the most part, the file formats described in this section are not static. They undergo constant revisions and updates, and many variants often exist for a particular format.

JPEG: Joint Photographic Experts Group

In its basic form, the JPEG standard, a widely-used and complex system developed by the JPEG committee of the ISO, consists of a large collection of file-compression options. More than two dozen variations are given in the JPEG definition, so that it can be implemented in a number of different ways, from simple lossless algorithms to very high-compression lossy methods. But the basic JPEG definition does not completely specify how the compressed image file should be structured...
so that it can be used on different computer systems or by different applications. For instance, there is no specified organization for the header information and there is no specification for the color model that should be used in the compressed file.

The JPEG standard defines four general file-compression modes, which are called the lossless, sequential, progressive, and hierarchical modes. In the JPEG \textbf{lossless mode}, a pattern-recognition scheme is combined with either Huffman encoding or arithmetic encoding. However, the original JPEG lossless mode is not as efficient as other available lossless formats, so it is rarely implemented. The \textbf{JPEG baseline sequential mode} is the most commonly used version of JPEG. Numerical values for the color components in a picture are stored in 8 bits, and the compression algorithm combines the discrete cosine transform with either Huffman or arithmetic encoding. An \textbf{extended sequential mode} is also defined with more options than the baseline sequential mode and in which color components can be specified using 16 bits. In the JPEG \textbf{progressive mode}, an image file is processed using several passes so that “layers” of the image can be generated at varying resolutions. This mode, generally referred to as \textit{progressive JPEG}, is popular for Internet applications, because a rough approximation of a picture can be viewed quickly before downloading the complete image file. Another collection of procedures for obtaining incrementally improved versions of an image is contained in the JPEG \textbf{hierarchical mode}, which divides an image into a set of subimages. This allows selected sections of a picture to be progressively constructed. Because of its complexity, hierarchical JPEG is not widely used.

Options could be provided in a large-scale JPEG implementation for the selection of both a compression mode and the compression parameters, such as the number of terms to be used in the summation calculations for the inverse, discrete cosine transform. Also, the JPEG compression definitions specify that either Huffman encoding or arithmetic encoding can be combined with the discrete cosine transform. But implementations of JPEG never use the arithmetic-encoding algorithms because these algorithms are patented and require a licensing fee.

Although the JPEG specification does not define a specific structure for the compressed image file, implementations now use the JPEG \textbf{File Interchange Format (JFIF)}, proposed by Eric Hamilton at C-Cube Microsystems and based on suggestions from many JPEG users. In this format, the file header contains a unique JFIF identifier (referred to as the file “signature”), the version of JFIF used to set up the file, the image size (either in pixels per centimeter or pixels per inch), the height and width of an optional RGB preview image of the file (referred to as a “thumbnail” image), and the RGB values for the optional preview image. Pixel values in the compressed file are stored using the $Y, C_b, C_r$ color model, and the color components are stored in the following order: $Y$ first, $C_b$ second, and $C_r$ third. For a gray-scale image, only the $Y$ component is used. Other information in the file includes the tables needed by the compression algorithms. Integers are stored in JPEG files using the big-endian format.

The JPEG/JFIF baseline-sequential encoding of an image file typically consists of the following operations:

1. \textbf{Color Conversion}: Pixel RGB color values in an image file are converted to $Y, C_b, C_r$ color components.

2. \textbf{Color Sampling}: The number of color values in the file can be reduced by using only the values from selected pixels or by averaging the color components for adjacent pixel groups. A simple implementation for this sampling operation might take the color values from every other pixel, every third pixel, or every fourth pixel. Usually, the color components
are sampled at different frequencies, so that more luminance values, \( Y \) components, are selected. This allows greater compression ratios to be achieved because fewer distinct chrominance values, \( C_r \) and \( C_b \) components, are saved.

(3) **Discrete Cosine Transform**: Next, \( 8 \times 8 \) groups of pixel color values are converted to discrete cosine-transform values using Equation 3.

(4) **Reduction of Transformed Values**: To further compress the encoded image file, a reduced set of transform values is stored (Section 3). The number of values in the reduced set can be fixed, or it can be computed using an algorithm to determine the influence of the various transform terms.

(5) **Huffman Encoding**: A final compression operation is performed by converting the discrete cosine-transform values to Huffman codes, as discussed in Section 3.

The **Still-Picture Interchange File Format** (SPIFF), developed by Eric Hamilton and the ISO JPEG committee, is an extension of JFIF. This format has many more features and options than JFIF, and it is expected that SPIFF will eventually replace JFIF in JPEG implementations. However, like JPEG, this extended JFIF format contains many more options than may be practical in one implementation. For example, JFIF uses just one color model \((Y,C_r,C_b)\), but SPIFF provides options for 13 different color models.

For photo-realistic computer-graphics images and digitized photographs, current JPEG implementations provide a greater compression ratio than any other system. But other formats can provide comparable compression ratios without loss of color information for simple pictures that contain large single-color areas.

**CGM: Computer-Graphics Metafile Format**

The CGM format is another standard developed by ISO and ANSI. It is designed for use on any computer system and in any area of computer graphics, including scientific visualization, computer-aided design (CAD), graphic arts, business graphics, electronic publishing, and any application using the GKS or PHIGS graphics library. Thus, CGM supports a variety of features and options.

As the designation “metafile” indicates, CGM allows an image description to be given as a pixmap or as a set of geometric definitions, including attributes such as line size, line type, fill style, and character-string specifications. Various other parameters can be included in an image file, such as the maximum value for color components, the size of a color table, list of fonts used in the file, and the bounds for a clipping window.

A character-encoding scheme is used in CGM to minimize file size, and a numerical, binary code is optimized for fast encoding and decoding of the image file. Pixel values can be given using various color schemes, such as RGB, CMYK, \( YC_rC_b \), CIE models, and color tables. In addition, pixmap files can be compressed using variations of run-length encoding and Huffman encoding.

**TIFF: Tag Image-File Format**

A consortium of computer companies chaired by the Aldus Corporation developed TIFF as an efficient format for transferring raster images between different applications and computer systems. Although it is highly complex, TIFF is one of the most versatile formats and it can be customized for individual applications. It is widely used in such diverse applications as medical imaging, desktop publishing, graphical user interfaces, satellite image storage, and fax transmissions.
The TIFF format can be used with bi-level, gray-scale, and full-color images, and TIFF files are designed to store multiple raster images. Pixel color information can be provided as RGB components or as color tables. More compression alternatives are provided in TIFF than in any other system. These compression schemes include combinations of run-length encoding, LZW encoding, Huffman encoding, and the suite of JPEG methods.

**PNG: Portable Network-Graphics Format**

Designed by an independent group of developers, PNG provides a highly efficient lossless compression scheme for storing images. Compression algorithms in PNG include Huffman encoding and variations of LZ encoding. This format is very popular on the Internet for image storage and transmission. It is also useful for temporarily storing images for repeated editing. For simple computer-graphics pictures, PNG generates files with very high compression ratios, comparable to those of compressed JPEG files.

Integer values are stored in big-endian order, and color components can be specified in a precision of up to 16 bits per pixel. A number of options are supported in PNG, including RGB color components, XYZ color components, grayscale, color tables, and an alpha value for transparency information.

**XBM: X Window System Bitmap Format and XPM: X Window System Pixmap Format**

Unlike other formats, XBM and XPM store picture information as C or C++ code that is to be processed on workstations using the X Window system. Thus, pixel values are represented in arrays, stored in scan-line order, left to right. As the names imply, XBM is a format for bitmaps (1 bit per pixel) and XPM is a format for pixmaps (multiple bits per pixel). These formats are supported by most Web browsers.

The XBM and XPM formats contain no compression algorithms, but the size of the files can be reduced using specially designed compression programs. Instead of header files, these formats use `#define` preprocessor directives to specify information such as the number of pixels per scan line and the number of scan lines. In the XBM format, bit values equal to 1 represent the current foreground color and bit values equal to 0 represent the current background color. In the XPM format, pixel values can be stored in color tables using RGB or hue, saturation, and value (HSV) components.

**Adobe Photoshop Format**

Widely used in image-processing applications, the Adobe Photoshop format is optimized for the fast accessing of large, full-color raster images. In contrast, very little compression is achieved with the run-length encoding scheme used in Photoshop, and earlier versions of Photoshop contained no compression methods.

Pixel values are stored in big-endian order, and Photoshop provides a number of options. Photoshop supports pixmaps, bitmaps (monochrome images), and gray-scale images. Colors can be stored using RGB color components, CMYK color components, or color tables. In addition, various schemes are provided for representing multiple colors per pixel and halftone images, as well as transparency parameters.
MacPaint: Macintosh Paint Format

A product of the Apple Corporation, MacPaint is a standard format for all Macintosh applications. Image files for this format are bitmaps, with a 0 value indicating white and a 1 value indicating black. The MacPaint format is typically used for text, line drawings, and clip art.

Pixel values are stored in big-endian order, and MacPaint files always contain 576 pixels per scan line and 720 scan lines. A run-length encoding scheme is used to compress image files.

PICT: Picture Data Format

PICT, a hybrid format, is another product for Macintosh applications from the Apple Corporation. It supports images that are specified as bitmaps, pixmaps, or geometric representations. A PICT file in the geometric-representation format contains a list of Macintosh QuickDraw functions that define a picture as a set of line segments, polygons, arcs, bitmaps, other objects, clipping parameters, attributes, and other state parameters.

Images can be specified using a monochrome form (bitmap), RGB color components, or a color table. Raster files can be compressed using a run-length encoding algorithm.

BMP: Bitmap Format

Although it is called a bitmap format, BMP actually supports image files that contain multiple bits per pixel. This format was developed by the Microsoft Corporation for Windows operating-system applications. Another similar pixmap format (also called BMP) is used by the IBM OS/2 operating system.

Pixel values in a BMP file are stored in little-endian order using 1, 2, 4, 8, 16, 24, or 32 bits per pixel. The pixel color values can be specified with RGB color components or with color tables. And the pixel scan lines are stored from bottom to top, with the coordinate origin at the lower-left position of the pixmap. A BMP file is usually not compressed, but a run-length encoding algorithm can be applied to pixmaps with 4 or 8 bits per pixel.

PCX: PC Paintbrush File Format

Developed by the ZSoft Corporation, PCX is another pixmap format used by Windows operating systems. Image files in the PCX format can contain from 1 to 24 bits per pixel, and pixel values can be specified using RGB components or color tables. Values are stored in little-endian order, with the scan-line ordering from the top of the image to the bottom. In addition, the raster files can be compressed using run-length encoding.

TGA: Truevision Graphics-Adapter Format

Developed by the Truevision Corporation for use with the Targa and Vista graphics adapters, the TGA pixmap format is also known as the Targa format. This format is popularly used for video editing.

In the TGA format, pixel values are stored in little-endian order, and image files can contain 8, 16, 24, or 32 bits per pixel. Pixel colors can be specified as RGB components or in tables, with two possible table formats. A single RGB color table can be used or the R, G, and B components can be given in separate tables. Typically, TGA files are not compressed, but run-length encoding algorithms can be applied to larger image files.
GIF: Graphics Interchange Format

GIF, designed for efficient telephone-line transmission of raster image files, is a product of the CompuServe Corporation. Using an LZW algorithm, GIF provides reasonably good compression ratios for simple computer-graphics pictures. But the compression ratios generated by GIF for photo-realistic images are not as good as those produced by JPEG or PNG. Although GIF has been used in many applications, its popularity has drastically declined because of the patent issues associated with the LZW compression algorithms.

Either monochrome or multicolor pictures can be processed by GIF, but pixel values can be specified only in the range from 1 to 8 bits, allowing a maximum of 256 colors. Pixel values are stored in little-endian order using RGB color tables.

5 Summary

For a raster-graphics system, an image file is typically an RGB pixmap, which is often referred to as a raw raster file. The RGB pixel values are stored as integers in the range from 0 up to a maximum value that is determined by the number of bits available to each pixel. A picture can also be stored using a representation that contains geometric descriptions of the picture components, such as line segments, fill areas, and splines.

When raster image files are to be transferred between systems or stored in a particular form, it may be necessary to reduce the number of color values represented in the image. We can uniformly reduce the number of colors by combining color levels in various ways, such as averaging the levels. The popularity method for reducing colors selects the most frequently occurring color values. And the median-cut method subdivides the color space into a set of blocks, with all colors within each block replaced by the average block color.

Various formats have been developed for storing image files in a convenient form for particular applications or particular systems. These formats differ in the structure of the header file, the byte ordering (big endian or little endian) for integer values, and the methods used (if any) to reduce the file size for storage. The effectiveness of a file-reduction method is measured by the compression ratio, which is the ratio of the original file size to the compressed file size. File-reduction algorithms that alter the color values in an image file are described as lossy, and those that can exactly restore the color values are described as lossless. Some file formats also employ color-reduction schemes.

A common compression method for image files is run-length encoding, which replaces a sequence of repeated pixel values with the value and the run length. The LZW file-compression scheme is a variation of run-length encoding that replaces repeated patterns of pixels with a code. Other pattern-recognition compression methods include scan-line comparisons and fractal procedures for identifying self-similar sets of pixel values. In Huffman encoding, a variable-length code is assigned to color values so that the most frequently occurring values have the shortest codes. Arithmetic encoding uses the frequency count for color values in an image file to create subdivisions of the unit interval from 0.0 to 1.0. The bounds on each subinterval are then used to encode the sequences of color values represented by that subinterval. The discrete cosine transform multiplies pixel color values by cosine terms with increasing frequency, and then sums these products. This summation process converts a set of pixel color values to a transformed set of values. File compression is then achieved by eliminating some of the transformed values, which produces a lossy compression of the image.
Many file formats are available for various graphics applications and for different computer systems. Some formats were developed by the standards organizations ISO and ANSI, some came from computer software or hardware companies, and some are the products of independent groups. A few of the widely used formats are JPEG, TIFF, PNG, and those for the X Window system, Apple Macintosh computers, and the Windows operating systems.

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