Color Grading
D-to-Z Diamonds
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About the Cover

The GIA D-to-Z color grading system revolutionized the manufacture of diamond jewelry because it allowed for tighter control over the quality of the diamonds used in individual pieces. At the same time, an important market developed for the finest, D-color, diamonds, as represented by the rings shown here. The D color, VVS2 round brilliant in the ring on the right weighs 7.12 ct; courtesy of Harry Winston Inc. The D color, VVS2 heart-shaped diamond in the other ring weighs 10 ct; from Mona Lee Nesseth, Custom Estate Jewels (courtesy of a private collector). The diamonds in the pin have colors ranging from F to I, and their total weight is 21.10 carats; courtesy of Louis Glick & Co. The necklace is a diamond sautoir by Van Cleef & Arpels, Paris, circa 1925; courtesy of a private collector.

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EDITORIAL

Challenges, Changes, and New Directions
Alice S. Keller

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The current global economic downturn has affected every industry, and ours is no exception. As one consequence, GIA has decided to postpone the second Gemological Research Conference, originally scheduled for August 21–23, 2009. Our foremost concern is that many people who had hoped to participate are now facing significant budget and travel restrictions, so—rather than risk the success of this important event—the conference will wait until the economy improves. Our heartfelt thanks go to everyone who was involved with the 2009 GRC, and we look forward to the time when preparations can be resumed.

To better continue delivering comprehensive research and exceptional gem photography, Gems & Gemology will be shifting a portion of its content online, starting with this issue. The Book Reviews and Gemological Abstracts sections are now available (in their entirety) only in electronic form, as are our year-end subject and author indexes, traditionally part of the Winter issue. The annual indexes will also be combined with our online cumulative index, which covers the years 1981–2008. Additional content, such as supplemental tables, spectra, and images, will be featured online in the G&G Data Depository. To use these resources, all of them available free of charge, just visit us at www.gia.edu/gemsandgemology.

This issue also marks the long-awaited launch of Gems & Gemology Online, which will offer electronic (PDF format) issues concurrently with the printed issue. Subscribers now have the option of receiving their issues in either print or PDF format, or selecting a combination of the two versions for a nominal additional fee of $25 per year. The online issue will go live the day the printed issue mails (in the case of the present issue, for example, the online version went live Friday, January 16). Those of you, especially overseas, who may have had to wait weeks for your issue to arrive by post will now be able to see it on the Internet the moment it becomes available. Note that we cannot guarantee that the color of the photos seen on your computer monitor will be the same as that verified by art director Karen Myers while the journal is on press. However, the online version uses the same electronic files that are used for the printed version.

G&G Online is hosted by MetaPress, a division of the world's largest subscription agency, EBSCO, at gia.metapress.com. There you will find electronic versions of every article and journal section from 1981 through the present. Editorials, Letters, Book Reviews, and Gemological Abstracts (as well as all the articles on our Free Downloads page) are available without charge. For a limited time, we are also offering a free look at the entire Fall 2008 issue online. Other articles and sections can be purchased for just $10 each, and entire issues for $20 each. (The smaller-format, predominately black-and-white issues from 1934 to 1980 can still be downloaded from the G&G web site at no cost.) If you're interested in upgrading your subscription to include online access, please contact our circulation coordinator.

Gems & Gemology is no stranger to challenges and change. When Robert Shipley published the first issue in 1934, the world was in the throes of the Great Depression. When the current large-format version was launched in 1981, the diamond industry was still recovering from the speculative boom and bust of the late 1970s. Now, as we implement these new initiatives in the midst of another period of economic uncertainty, we're confident we can continue to support GIA's mission to ensure the public trust in gems and jewelry by upholding the highest standards of professionalism in one of our industry's most important sectors: gemological research. G&G looks forward to maintaining excellence in print and online in 2009 and for many years to come.

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Since its introduction in the early 1950s, GIA’s D-to-Z scale has been used to color grade the overwhelming majority of colorless to light yellow gem-quality polished diamonds on which laboratory reports have been issued. While the use of these letter designations for diamond color grades is now virtually universal in the gem and jewelry industry, the use of GIA color grading standards and procedures is not. This article discusses the history and ongoing development of this grading system, and explains how the GIA Laboratory applies it. Important aspects of this system include a specific color grading methodology for judging the absence of color in diamonds, a standard illumination and viewing environment, and the use of color reference diamonds (“master stones”) for the visual comparison of color.

Historically, the evaluation of most gem diamonds focused on the absence of color (Feuchtwanger, 1867; figure 1). Today, this lack of color is expressed virtually worldwide in a grading system introduced by GIA more than 50 years ago that ranges from D (colorless) to Z (light yellow). With the acceptance of this system, color grade has become a critical component in the valuation of diamonds, leading to historic highs at the top end of the scale. In May 2008, the diamond in figure 2 [a 16.04 ct round brilliant that GIA graded D color, VVS2 [potentially Flawless]] sold at Christie’s Hong Kong for a record US$208,500 per carat for a colorless diamond. At the same auction, a 101.27 ct shield-shaped diamond sold for $61,500 per carat, a vivid reminder of the impact of even minor differences in grade; it was F color, VVS1. At auction, and throughout the marketplace, differences between adjacent color grades can result in substantial differences in asking price. For example, on December 5, 2008, the Rapaport Diamond Report noted an approximately 32% (and Idex about 33%) difference between D and E color for a one carat Internally Flawless round-brilliant diamond. Important price distinctions based on color are not limited to high-end colors and clarities. On the same date, the difference between G and H color for a one carat round-brilliant diamond of VS1 clarity was about 16% in both Idex and Rapaport.

As part of its educational program, GIA has taught the basics of color grading D-to-Z diamonds since the early 1950s. And in the more than five decades since the GIA Laboratory issued its first diamond grading report in 1955 (Shuster, 2003), it has issued reports for millions of diamonds using the D-to-Z system. Throughout this period, GIA has experienced increased demand for its diamond grading services over a growing range of diamond sizes, cutting styles, and color appearances. This has required a continual evolution in the equipment and methods used in the GIA Laboratory, while maintaining the integrity of the grading system itself. At the core of the system’s development has been an ongoing assessment of how best to observe a diamond in order to describe its color consistently. At times, the resulting adjustments have appeared to conflict with earlier statements.

This article reviews the history of the system’s development and the various methodologies that have been employed to observe and report on color in diamonds.
development to help clarify the various modifications that have taken place over the years in the color grading equipment and practices used at the GIA Laboratory. We then describe how D-to-Z diamonds are currently graded in the lab, that is, the procedures that have resulted from these years of evolution and refinement. Last, we discuss special considerations in D-to-Z color grading, such as the grading of brown and gray diamonds, the selection process for master color comparison diamonds ("master stones"), and the impact of adopting advanced instrumentation.

BACKGROUND

While there has been evaluation of diamond color (or absence of color) throughout history, the systems and methods used for this purpose were not clearly defined, standardized, or consistently applied before the 1950s. In the late 19th century, color was considered a diamond’s most important value factor, but the naming conventions in use at the time placed color in a variety of categories that were general at best. For example, the color appearance of gem diamonds was often described using metaphoric terms (e.g., “River” or “Water” for the most colorless diamonds), or by association with a geographic location from which similarly colored diamonds were commonly seen (e.g., “Wesselton” and “Top Wesselton” for near-colorless diamonds traditionally associated with the Wesselton mine, “Cape” for pale yellow diamonds from the Cape of Good Hope region, and “Jager” for colorless diamonds with strong fluorescence such as those typically recovered from the “Jagersfontein” mine in South Africa (Shipley, 1950b; Liddicoat, 1993). In the case of blue white, abuse of the term eventually prompted action by the U.S. Federal Trade Commission to ban its misuse in diamond marketing (Shipley, 1938).

Recognizing the importance of objective, consistent color communication, GIA—in conjunction with the American Gem Society (AGS)—began work on color grading standards in the 1930s (“Diamond grading instrument...,” 1934), and by 1941 had developed a color scale for evaluating diamonds (see following section; Barton, 1941; Shipley and Liddicoat, 1941; AGS, 1955; Shuster, 2003). The development of this “color yardstick,” as it was then described, evolved over the next decade and became the basis for what is known today as the D-to-Z color grading scale.

While color grading was introduced in the early 1940s for AGS members, the general trade was not familiar with the GIA/AGS standard. Richard Liddicoat, who became the executive director of GIA in 1952, created a full diamond grading system that he taught for the first time in 1953 to jewelers in classes around the United States (Shuster, 2003). Soon, other GIA instructors such as Bert Krisches and G. Robert Crowingshield became part of the traveling team that taught this new grading system, which greatly expanded interest in—and use of—this approach.
Attracted by the system’s ability to generate a diamond’s market value based on the new quality grades and other concepts regarding diamond proportions and appearance (Gilbertson, 2007), jewelers flocked to GIA to get a better understanding of diamond valuation. GIA’s new system launched the scale of “D to Z” for color grading. Regarding the unusual starting point (the letter D), Richard Liddicoat (Liddicoat, n.d.; Gem Talk, 1981) stated the choice was made to differentiate the GIA system from other less clearly defined ones that used designations such as “A,” “AA,” or metaphoric terms like those noted above.

As jewelers went home to grade their own diamonds, they started to question some of their decisions and sent the diamonds to GIA for checking by their instructors. Over time, this informal practice led to GIA’s diamond grading laboratory service, with the first formal reports issued in 1955.

For more than 50 years, this grading system has been taught by GIA Education and used in the GIA Laboratory. The combination of understandable letter designations for color grades, the availability of a standardized grading environment and sets of diamonds as color references, and GIA’s ability to teach the basics of this system to others provided a new level of stability and confidence in diamond commerce. In the decades since its introduction, the D-to-Z nomenclature has been adopted virtually worldwide for the sale, purchase, and evaluation of polished diamonds. Consequently, other grading laboratories also use this nomenclature, sometimes in combination with their own. While most claim to use the GIA system, however, it is not likely that it is applied as it is at the GIA Laboratory. Using the same color grading terms does not constitute adhering to the conditions or methodology of the GIA system. The reasons for this should be evident by the end of this article.

The Origin of GIA’s Viewing and Comparator Standards. In 17th century India, Tavernier (1676) noted, diamonds were color “graded” at night by lamplight. By the 19th century, however, daylight was the worldwide standard in which gemstones were observed to discern their color (Chester, 1910; Cattelle, 1911; Wade, 1915, 1916; Ferguson, 1927). Unfortunately, the characteristics of daylight vary (throughout the day, in different locations around the globe, and at different times of the year), and these differences in light quality can significantly affect the color appearance of gemstones (Cattelle, 1911; Wade, 1915; Sersen and Hopkins, 1989).

The historical methods used to observe and compare diamond color were just as variable as the type of illumination itself. Observers held diamonds in the palms of their hands or between their fingers, typically examining them against a range of different backgrounds (Tavernier, 1676; Mawe, 1823; Feuchtwanger, 1867; Morton, 1878; Wodiska, 1886; “On diamonds,” 1902; Wade, 1916). For most of history, of course, diamonds were so rare that very fine color distinctions among them were not needed, so the trade could function using such simple evaluation techniques.

With the discovery of large deposits in southern Africa in the late 19th century, more diamonds entered the marketplace than ever before. This influx generated a greater desire (more specifically, a commercial need) for finer color distinctions. By the early 20th century, certain minimal standards for color grading had evolved, as summarized from Wade (1916):

1. Use “good north light unobstructed by buildings or other objects. There must not be any coloured surface nearby to reflect tinted light, as a false estimate might easily result.”
2. Color grade diamonds only between 10 a.m. and 2 p.m.
3. Do not use artificial light.
4. Always use the same location for color grading.
5. Use comparison diamonds.
6. Dim the “fire” (i.e., dispersion) of the diamond, perhaps by breathing on the stone.
7. View the stone on edge as well as face-up (face-up only can yield a false color).
8. Use magnification (aplanatic triplet lens).

Even with these standards, color grading among dealers and retailers was inconsistent; the result in the trade was chaos. Cattelle (1911, p. 134) noted that “Color in diamonds is the opportunity of many dealers, and the despair of others, for it is very deceptive.”

By the mid-20th century, advances in lighting technology and vision science paralleled the jewelry industry’s increasing desire for improvements in diamond color analysis. GIA began one of the earliest efforts to address this desire in the mid-1930s (“Diamond-grading instrument . . . ,” 1934; AGS Research Service, 1936), which culminated in a 1941 article by GIA founder Robert Shipley and Richard Liddicoat. Among their concerns was the need for a grading scale with uniform comparators (i.e., color references) that could be used throughout the industry.

Although jewelers commonly kept a few reference diamonds to use for color comparison, there was no standard for such comparators, so the color grade given to a diamond by one jeweler could easily differ from the grade assigned to it by another. Moreover, different jewelers used different sources of light, both artificial and natural, which further complicated their interactions with one another. To address this situation, Shipley and Liddicoat’s 1941 article announced the development of: (1) a visual color comparison instrument (the GIA Colorimeter), (2) a color scale (the “color yardstick”) that represented grade categories, (3) a standardized light source and viewing environment for use by jewelers (the Diamolite), and (4) a service to grade “reference” or “master” diamonds for AGS jewelers to use as comparators.

The GIA Colorimeter and Color Yardstick. Notes and drawings from Shipley in the 1940s indicate that the GIA Colorimeter was adapted from the Duboscq Colorimeter made by Bausch & Lomb, which was widely used in the medical field up to the early 1960s (Warner, 2006). Shipley and Liddicoat’s colorimeter (figure 3) consisted of a small box with an indirect light source and a split-image magnifier that allowed a grader to compare a diamond placed in a tray to a movable, transparent glass wedge that varied (and was graduated into ranges) from colorless at the thin end to yellow at the thick end (Shipley and Liddicoat, 1941). This first system had 13 grade ranges: The glass wedge had markings for seven categories, which were labeled “0” followed by Roman numerals “I” to “VI,” with these latter six further separated by half-division marks (see figure 3, inset). These 13 grades equated to the D-to-P range in GIA’s later system.

With this colorimeter, the color determination was made by visual comparison, with the diamond placed in three different positions (table-up, girdle-up, culet-up), and 10 observations in each position. The results obtained for each position were then averaged to reach the overall color grade. It is interesting to note that this early form of color grading at GIA was done with magnification (about 4×) and through the comparison of the diamond to a flat transparent wedge of graduated colored glass rather
than other diamonds. The box in which the diamond was viewed against the glass wedge housed a light source for observing the diamond and eliminated influences from outside lighting. The light source used was a blue-coated incandescent bulb that was intended to mimic the color appearance of daylight.

When Shipley’s son, Robert Shipley Jr., felt the colorimeter was ready to use for color grading, he wrote to his father in a letter dated April 2, 1941: “I have run thirty reading checks on four stones on which Dick [Liddicoat] has made thirty reading averages. Neither of us had any reference to the others’ work, and in no case were we off more than .125 of a division!” Nevertheless, to the best of our knowledge, the GIA Colorimeter was never manufactured commercially (apparently only two were made) or used for any purpose other than the grading of AGS master stones by GIA staff.

The GIA Diamolite. While the GIA Colorimeter allowed for the visual comparison of a diamond to a color standard to develop sets of master stones, jewelers still needed a standard viewing environment when comparing other diamonds to these masters. Although some jewelers had stores situated such that they could effectively use north daylight, many did not, so they used whatever light source they had available. Up to this point, artificial light had proved problematic because it was not close enough in color appearance to natural north daylight (the accepted trade standard). And, again, “north daylight” varied depending on the time of year, time of day, weather, and geographic location; the color of the diamond was also influenced by colors in the viewing area. Ultimately, the Diamolite was the first step toward providing a solution to this problem (figure 4).

From the development of their colorimeter, GIA researchers knew that controlling the light source would be critical, but they also realized that an environment to compare diamonds to one another would have different requirements from that of the colorimeter. Shipley Sr. had been working on this issue (“Diamond-grading instrument . . . ,” 1934), but it was Shipley Jr. who carried it to its next step. He and others at GIA examined every new light source on the market; they tried argon bulbs (AGS Research Service, 1936) and filtered incandescent bulbs, as well as the relatively new fluorescent lamps (“Instrument research . . . ,” 1937). Even though GIA was selling the “Da-Grade” fluorescent light source, made by General Electric (GE), for use in displaying diamonds (“At last . . . ,” 1938), they realized an observer could not use it to distinguish fine nuances of faint yellow color, so the search continued for a better light source for grading.

Shipley Jr. also looked to other industries in which fine color distinctions were critical and found a filtered incandescent light source used as a standard for oil colorimetry and cotton grading. The Shipleys and Liddicoat worked with the color technology company Macbeth (Shuster, 2003) to counterbalance the overabundance of the long rays (i.e., “redder” light) of this tungsten bulb with a special blue filter (figure 4, inset). This adjusted the color temperature (i.e., appearance) of the light output to be closer to that of daylight.

Shipley Jr. also recognized that color grading needed an enclosed viewing environment: “Preliminary research shows that the greatest single handicap in accurate color grading is the difficulty of securing a light absolutely free from colored reflections from adjacent objects” (AGS Research Service, 1936, p. 77). The following excerpt from an article about changes to the Gemology textbook (“A note on diamonds,” 1938, p. 174) expresses GIA observations at the time on the effect of external lighting on diamond color appearance and defines “body color” for the trade:

Figure 4. The GIA Diamolite was introduced to AGS members in 1941 and, for the first time, offered jewelers a controlled lighting and viewing environment for comparing “master” diamonds to other diamonds. The Diamolite measured 13 3⁄4 in. high × 12 in. wide × 6 in. deep (34.4 × 30 × 15 cm) and housed an incandescent bulb with a blue filter (inset) designed to make the light output better simulate daylight. Photos by Robert Weldon.
The color of a diamond, as it is seen by the eye, may be affected (1) by the comparative amount of the various spectrum colors which it disperses, and (2) by the color of the light reflected from sky, walls, ceiling or other objects. Upon examination by transmitted light against a white, neutral gray, or black background, the true color of the diamond itself is observable, and the resulting appearance is known as the body color. However, if colored reflections fall upon those surfaces of the diamond which are toward the eye, the true body color may not be observable. (If colored reflections fall upon a white background against which the diamond is being examined, they will also affect the body color.)

Some in the trade already understood this effect of the lighting environment. By the late 1800s, diamond dealers were very aware of how the surrounding environment affected the appearance of color in a diamond. In the New York jewelry district on Maiden Lane, neighboring buildings were painted yellow—and these faced the windows from which dealers judged color in the north daylight. The dealers pooled their funds and offered to repaint the offending buildings. Jewelers' Circular reported: “Several [dealers] stated that it is impossible to sell diamonds in their offices, and unless the colors of [buildings] No. 5 and 7 are changed, they will be forced to vacate their offices” (“Dealers object...,” 1894, p. 16).

GIA researchers ultimately determined that a box made of translucent white matte paper, open on one side for observation, furnished a much more satisfactory environment for judging color (AGS Research Service, 1936). Not only was the direction of the light that fell on the diamond controlled, but the light was also diffused, which subdued surface reflections from facets that might obscure the body color. In 1941, the Institute introduced a commercial version, the GIA Diamolite, for use in color grading (again, see figure 4; Shipley and Liddicoat, 1941). The accompanying brochure stated that this viewing box (small enough for the display counter in a retail store) allowed a jeweler to observe diamonds under a standardized light source in any physical location at any time of the day or night.

Grading Master Stones. Now that it had a light source and an environment in which to visually compare diamond color, as well as an instrument that related the color to an established scale, in the early 1940s GIA began to collect diamonds graded with the colorimeter to use as its own “master” comparators.

After leadership review, the AGS membership voted in 1941 to recommend the use of the GIA Colorimeter for master stones, the scale as the standard for color distinctions in diamonds, the GIA Diamolite, and the new grading service for master stones (see, e.g., Shipley and Liddicoat, 1941). For the AGS master stones, two of the averaged colorimeter grades were reported to the AGS member jeweler—for the table-up and girdle-up positions. The "0–VI" numerical scale of the colorimeter was promoted as the color grade nomenclature to be used for diamonds (Shipley and Liddicoat, 1941) and was strictly for AGS members. Consequently, when GIA chose to develop a diamond grading system available to everyone, it had to use new terms. The choice for color grading colorless to near-colorless diamonds, as previously noted, was the letter grade scale beginning with D.

GIA had offered other technical services, such as pearl identification, to the trade since the 1930s, but the master diamonds service (started in 1941) was the first one related to diamond grading. In fact, this service preceded the 1949 creation of the laboratory as its own division within GIA.

The 1950s to ‘80s: The Continued Evolution of Lighting and Viewing Environments. The last known mention of the name Diamolite was in the January 1950 issue of Jeweler’s Circular Keystone magazine. Gems & Gemology first cited the new name, DiamondLite, in 1949 (Schlossmacher, 1949). This change in name accompanied an updated filter from Macbeth that was still used with an incandescent bulb (Shipley, 1950a). We do not know if the physical proportions of the viewing environment changed at that time.

This period saw the continuation of GIA’s investigation into alternative light sources for the assessment of diamonds. By the 1950s, Liddicoat and his contemporaries had apparently developed a comfort level with the fluorescent lamps available at that time. GIA began teaching that jewelers could use a modified fluorescent lamp for color grading when they didn’t have access to a (still incandescent bulb) DiamondLite: “a reasonable substitute may be secured by adapting a simple fluorescent tube desk lamp...lined with flat-white paper...enclosed on the back and two sides so as to exclude as nearly as possible all reflections from surroundings” (Shipley, 1955, p. 5). A sheet of
white tissue paper between the fluorescent tubes and the diamond further diffused the light.

Depending on the evenness, thickness, and type of phosphor coating, however, the early generations of fluorescent lamps had considerable variation in the wavelengths and intensities of their light output. This included an inconsistent—and, at times, relatively high—amount of ultraviolet (UV) emission, in contrast to the extremely low UV content in incandescent bulbs commonly used at that time. Realizing that some fluorescent diamonds appeared different when observed under lights with UV content (fluorescent lamps as well as the former Diamolite with its UV source, added in 1947, turned on), GIA made a number of statements related to this with each accompanying modification. In 1955 course material, GIA advised that “This factor should not cause too much difficulty, however, since only a very small percentage of stones fluoresce strongly enough to modify the body color under this light source” (Shipley, 1955, p. 5). Nevertheless, contemporary course materials advocated the grading properties of the (low-UV) incandescent bulb grading light, even to the disavowal of the historical standard, (relatively higher UV) daylight. GIA Assignment 2-31 (Shipley, 1957, p. 8) stated that “Fluorescent stones should be graded at their poorer color [as seen] in artificial light devoid of ultraviolet radiation [i.e., the incandescent bulb of the DiamondLite], rather than at their daylight grade [i.e., the grade they would receive if viewed against a comparison stone in daylight].”

In the mid-1960s, GIA introduced the GIA Diamond Color Grader tray to their Mark IV gemological microscope, which allowed color grading under the overhead diffused fluorescent source attached to the microscope. However, Gem Instruments advised that “Highly fluorescent stones cannot be graded in the GIA Color Grader” (GIA, 1966), due to the proximity of the diamond to the lamps.

In 1974, Ken Moore (director of Gem Instruments Corp.) introduced a DiamondLite (figure 5) that used 6-watt fluorescent lamps made by Verilux, with a new coating that minimized UV emission as compared to similar lamps and especially as compared to earlier fluorescent lamps. GIA often promoted the minimized UV emission in these lamps. For example, in 1979 course materials, GIA described these new lamps as “practically devoid of ultraviolet waves” (GIA, 1979, p. 9). Later course material (GIA, 1995, p. 9) claimed that “Filtered, cool white balanced fluorescent light is best: unlike sunlight, it is nearly free of ultraviolet.” As frequent as such statements were, it is important to remember that no fluorescent lamp is truly “UV free.”

From the 1990s to the Present: Lighting and Viewing Refinements. Research in the 1990s and the first decade of the new millennium led to refinements in the equipment and processes used to color grade diamonds that are reflected in the grading methods in place today.

Light Source Testing. Since at least the 1970s, millions of diamonds have been graded using fluorescent lighting, at GIA and throughout the industry. It has become the standard in the diamond industry. To ensure consistency in GIA’s grading, proposed changes in lighting must be thoroughly tested to balance the potential benefits to the grading methodology against the very real damage that would be caused if subsequent color grades were inconsistent with earlier ones. (The success the laboratory has had in this regard can be tracked in very real terms through its update service. Today we occasionally see diamonds graded in the 1970s that have been submitted for updated grading reports; after they have undergone a full grading process using contemporary equipment and procedures, the vast majority are returned with the same grade determinations.) With that as a guideline, GIA began researching options for fluorescent “daylight equivalent” lighting. In part, this research was driv-
by the fact that the Verilux 6-watt lamps used in the DiamondLite were no longer readily available, and other manufacturers’ lamps of the same size did not consistently meet GIA lighting standards.

Researchers compiled a list of factors to be considered in selecting an alternative lamp: worldwide availability, suitable illumination levels, uniform distribution across the work area, a spectrum that mimics International Commission on Illumination (known as CIE) D55-to-D65 specifications, a color temperature in the 5500–6500 K range, and a color rendering index of at least 90 (for details, see box A). Data collection began in January 1998, with 40 different lamps from various manufacturers. The 6-watt Verilux lamp used in the DiamondLite at that time was included for comparison. For each manufacturer, four different sizes of fluorescent lamps were tested: 4 watt (134 mm/5–6 in.), 6 watt (210 mm/8–9 in.), 15 watt (435 mm/17–18 in.), and 18/20 watt (590 mm/23–24 in.). Spectra were collected when the lamps were first turned on and after “burn in” times of 50, 100, 500, 1,000, and 2,000 hours, so that the evolution of each spectrum could be analyzed (see, e.g., figure 6). Data were collected using a Photo Research PR–704 Spectrascan spectroradiometer, an Ocean Optics SD2000 spectrometer, and a Gossen Mavalux digital lightmeter.

In late 1998, GIA researchers did additional testing on three of these lamps: an 18-watt Osram Biolux, a 20-watt Verilux, and a 20-watt Macbeth. For comparative visual observations, choices were narrowed to the Verilux and Macbeth lamps. The former was selected because researchers felt that continuity with the Verilux lamp characteristics was important, and the latter was chosen because GIA had been using the Macbeth lamp successfully to color grade colored diamonds for a number of years. After weighing all the factors and completing data analysis, GIA decided that the Verilux lamp was the best lighting source for the purpose of color grading D-to-Z diamonds.

Viewing Environment. The decision to use the longer (23–24 in.) 20-watt lamps required the design of a new and much larger viewing environment to house them. From its research on color grading colored diamonds (see, e.g., King et al., 1994), GIA recognized that a larger viewing box would also better shield the observer from distracting visual clutter in the surrounding environment, give a larger neutral background for the field of vision, and be more comfortable for the observer. These factors contributed to our decision to make the height of the box 18.5 in. (47 cm), which is similar to the Judge II used for colored diamonds (20 in.; illustrated in King et al., 1994, p. 230). Recognizing that the depth of environments used for D-to-Z color grading had traditionally been shallow (approximately 8 in. in the DiamondLite to 6 in. in the Diamolite), we decided to make the prototype viewing box 6 in. (15.24 cm) deep. As mentioned above, the width was determined by the length of the two lamps being used. In practice, we found that the wider environment was better for handling the diamonds. To improve the consistency and life cycle of the lamps, we successfully experimented with electronic high-frequency ballasts. Although the walls of the DiamondLite were coated dull white, GIA staff members had found that medium to light gray walls reduced eye fatigue (at the same time, various standards organizations also recommended a neutral gray surround for color grading environments; ASTM, 2003; ISO, 2005).

Subtle modifications were made to the design of the box over the next five years, as a prototype was formalized for use in the laboratory in the early 2000s. At the same time, we also developed a product for commercial release, a viewing environment that incorporates two 15-watt Verilux lamps into a slightly smaller (more retailer friendly) box, which is market-
Proper illumination is critical when performing tasks requiring subtle color comparisons, as is the case with color grading D-to-Z diamonds. There must be enough light to view the subtle differences, but not so much that color perception is affected by surface glare or that the light causes eye fatigue. We have found the acceptable range for light output to be between 2000 and 4500 lux. The light output needs to be stable (which is accomplished with an efficient, high-frequency ballast) so there is no variation in intensity (e.g., flickering), and it should be consistent across the entire viewing surface. It should take very little time for the lamps to become stable once they are turned on. The light emitted also must be diffuse, since point or spot lighting can cause bright surface reflections, more obvious dispersion, and strong contrasts in polished diamonds.

There are a number of characteristics of daylight that are valued in diamond color grading and were considered in choosing the standard light. These include, for example, daylight’s overall spectrum, its color appearance in the northern hemisphere, and its ability to render colors. While the full emission spectrum should be considered when choosing lamps, it is also important to note the output in the areas that can affect the predominant task at hand, in this case the grading of colorless to light yellow diamonds. In particular, there must be enough output in the blue region of the spectrum, as it is the absorption area of wavelengths in this region that allows yellow diamonds to be perceived at optimum visual acuity. Fluorescent lamps have “spikes” in their emission spectra (see again figures 6 and 8): narrow ranges of wavelengths that have much greater intensity. The positions of these spikes and their potential effect on color grading were important to our choice of a light source. In addition, the spectrum of a lamp used for color grading should not emit short- and mid-wave UV, as these emissions can be harmful to the eyes of the observer over extended periods of time. However, the lamp should emit long-wave UV, which is an important characteristic of daylight. The CIE standards for D55 to D65 light also specify a UV component.

Regarding the UV component, we have learned that for some fluorescent diamonds the distance between the lamps and the grading tray can influence the final color grade. For consistency, we use a distance of 8-10 in. (20-25 cm) between the lamps and the diamond. Bringing a fluorescent diamond closer to the lamps may result in a stronger fluorescence impact. For instance, a yellow diamond with strong blue fluorescence could appear less yellow (i.e., to have a higher color grade) as it gets closer to the lamps. Moving the same diamond more than 10 in. from the lamps will have the opposite effect; that is, the color will appear more yellow (a lower color grade).

The relative amount of UV versus visible light

Figure 7. This viewing box, the DiamondDock, provides a good surround for making visual comparisons of diamonds, both for color grading D-to-Z stones and observing round brilliant cuts. When the raised platform is inserted in the box as seen here, a viewing tray of master diamonds is at the appropriate height for color grading. When the platform is removed (and the auxiliary LED lights turned on), cut can be observed with the diamonds placed in a tray on the base of the unit. The box measures about 18 in. high × 21 in. wide × 6½ in. deep (45 × 52.5 × 16.25 cm). Photo by Robert Weldon.
York and Carlsbad laboratories independently color graded the same diamonds in the DiamondLite and the two new viewing environments to verify the consistency of grading results. Our findings showed that overall results were within tolerances recorded in the history of GIA’s D-to-Z color grading.

Spectral analysis of the three lamps showed consistency as well (figure 8).

THE UV CONTENT IN LIGHT SOURCES USED FOR D-TO-Z COLOR GRADING

The potential effect of lighting on diamond fluorescence (and therefore color appearance) has long been a subject of discussion, although the UV component in a light source only impacts color appearance in some obviously fluorescent diamonds. In the 1930s and before, there was general agreement that diamonds should be observed in north daylight, which contains UV. Yet the first widely accepted standard viewing environment, the Diamolite, contained a filtered incandescent bulb, which had low UV content compared to daylight. As noted earlier, fluorescent lamps of that era were neither stable enough nor consistent enough to meet the requirements of diamond color grading.

During the 1940s, the appearance of a highly fluorescent diamond in daylight was considered a positive attribute. Recognizing the inherent limitations of the Diamolite’s incandescent bulb in this regard, GIA introduced a stand-alone long-wave ultraviolet light source in 1945 (Shannon, 1945) and in 1946 updated the Diamolite by adding a UV source. When used alone, this UV lamp revealed the presence and strength of fluorescence in a diamond and, when used in conjunction with the filtered tungsten lamp of the Diamolite, created a condition that was felt to better simulate daylight and “show the
diamond off under the most attractive and favorable conditions” (Collison, 1947, p. 431).

As noted earlier, GIA continued to experiment with different lighting sources throughout the 1950s and ’60s, and began moving toward the use of fluorescent lamps, both in conjunction with the incandescent bulb of the DiamondLite and separately in the overhead light source of their microscopes.

Eventually, the use of different phosphors, phosphor layer thicknesses, and new ignition technologies reduced the amount of UV emitted by fluorescent lamps. Research that GIA began in the early 1970s revealed that the lower UV content helped reduce the extreme appearance differences in diamonds that had been encountered with earlier lamps. To market the new DiamondLite, with its 6-watt Verilux lamps, some GIA literature implied that little to no UV content was preferable for D-to-Z color grading. Actually, the low UV content of these lamps was preferable to the UV content of earlier fluorescent lamps, which was higher or inconsistent. Indeed, the lamps chosen in the ’70s had a small, but not negligible, UV component. And we continue to see this UV component in lamps chosen since then.

We recognize, however, that language used and certain statements made by GIA in the past several decades have led to confusion about the presence and, therefore, perceived desirability of the UV component in lighting used for D-to-Z diamond color grading. As mentioned above, course materials from the late 1970s to mid ’90s described the lamps as “practically devoid” or “nearly free” of UV (GIA, 1979, p. 8; 1995). Even in the GIA Laboratory, an internal manual published in 1989 (p. IH-1) noted “Use a cool white, filtered, ultraviolet free fluorescent light [in the lab, the DiamondLite] in an area of consistent, subdued light.”

“Filtered” referred (incorrectly) to the coatings used on the lamps to control output across the spectrum; this could be confused with using a filter to block UV. Again, these fluorescent lamps were not UV free.

In the late 1990s, referring to lamps with essentially the same UV content as their predecessor, laboratory staff spoke of the appropriateness of that UV component. As one of the authors [MK] commented in an interview (Roskin, 1998, p. 149), “Yes, you can create an environment devoid of UV but it’s a false situation…. It may sound like the ideal, but it steps outside the practical world. It’s not relevant because it doesn’t really exist anywhere. We try to be sensitive to the practical gemological issues.” Tom Moses corroborated this position at GIA by stating, “we found that the Verilux bulbs used in GIA’s diamond-grading units, standard cool-white fluorescent light bulbs, and northern hemisphere daylight (even filtered through a glass window) all have a certain amount of UV radiation. Hence the Verilux sources are similar—in terms of UV exposure—to grading environments throughout the world” (Moses, 1998, p. 21). The light source GIA uses for color grading has continued to be discussed in the trade (Tashey, 2000, 2001; Haske, 2002; Cowing, 2008).

The fact is that since the 1974 implementation
of new coatings on fluorescent lamps, GIA has promoted using a daylight-equivalent fluorescent lamp with a non-negligible amount of emitted UV. GIA will continue to study the subject as new lighting technology and research become available, with the goal of maintaining existing color grading standards. In fact, one of the authors (RG) was recently informed by CIE (Peter Hanselaer, pers. comm., 2008) that the UV content of the proposed new reference illuminant for “Indoor Daylight” (e.g., ID65) that CIE is working on will define the daylight specifications indoors under standardized conditions regarding glass thickness and absorption of the windows and standard angles of incidence of the light. This “Indoor” reference illuminant will have a noticeably reduced UV content compared to the regular CIE Daylight standard D65 because of the typical absorption of glass in the UV region, so its resulting spectrum will be even closer to the Verilux lamps GIA is using in the lab.

THE GIA D-TO-Z COLOR GRADING SYSTEM
The equipment and methods used today at GIA to color grade D-to-Z diamonds have come from the experience gained through the observation of millions of diamonds, as well as from continuous research into advances in lighting technology and vision science. Even though there have been modifications, such as changes to the viewing environment, it is important to recognize that the GIA standard—the spacing of the key historical grade markers—has remained unchanged since the system’s inception more than 65 years ago.

D-to-Z color grading is based on the observations of a trained observer, who compares a diamond to color master stones of known position on the grading scale (see box B for a discussion of the selection and care of master stones for clients and the laboratory). To achieve repeatable results, graders use a standard light source and a controlled viewing environment. Also important are the proper maintenance of equipment and consistency in the set-up of references, viewing geometry, and methodology. In addition, the observer must have been tested and shown to have normal color vision.

Screening, Training, and Monitoring of GIA Color Graders. Controlling all the conditions would be of little value without the proper screening and training of staff. At GIA, eligible staff members must pass tests such as the Dvorine Color Test, the Matchpoint Metameric Color Rule Test, and the Farnsworth-Munsell 100 Hue Test to ensure that they have normal color vision, discrimination, and acuity. Other tests are designed to gauge visual and verbal understanding of the color grading process.

Training sessions with experienced graders allow those staff members who are accepted as new color graders to gain first-hand knowledge over a period of weeks. All staff members are routinely monitored through the data collection of “blind” observations on control stones as well, to help insure color grading consistency.

To control for potential perception differences from individual to individual, GIA’s grading process requires a minimum of two or three random, independent opinions (depending on the size of the stone). A consensus is required before a color grade is finalized. For larger or potentially D-color stones, the laboratory’s computer operating system identifies the need for the most experienced graders. Last, to avoid the potential of reduced accuracy due to eye fatigue, color grading sessions are limited to approximately one hour, at which point a minimum break of one hour must be taken.

Routine Calibration and Maintenance. Since the viewing environment (e.g., the DiamondDock) is the neutral surround for the observer’s field of vision, its care and cleaning is the first priority before grading even begins. If it is soiled it can distract the observer just as objects in the field of vision would. In the laboratory, the viewing environments are cleaned with a mild soap and soft cloth every week. Prior to placing new lamps in a unit, we capture their spectra with an Ocean Optics spectrometer equipped with an integrating sphere to make sure they are within tolerances. The light output at the surface of observation is checked monthly using a Gossen Mavo lux Digital 5032B lux meter. Our testing has shown that the average life of a lamp is around 5,000 hours; from our experience, the “useful” life of the lamp for color grading purposes is half that, 2,500 hours. To avoid any deterioration in the illumination, we replace the lamps even sooner, at approximately 1,800 hours, unless some problem is noted earlier (i.e., discoloration at the ends of the lamps or a prominent drop in the lux meter readings). In the color grading area of the laboratory, ambient lighting is also controlled. Overall, the lighting is subdued, with no influence from natural daylight.

Again, see box B for the routine cleaning and care of master stones.
Locations of Master Stones in the Grade Ranges.

Every individual grade designation on the D-to-Z scale is actually a range of colors within that grade. The GIA master stones are located at the highest boundary of each grade range (figure 9), that is, at that end of their respective grade range that has the least color. Therefore, a diamond with less color than the G master stone (but not less than the F) would receive a grade of F. If the diamond appears to have the same amount of color as the G master, it would receive a grade of G. When a diamond has slightly more color than the G master stone but less than the H master stone, it will be called a G color. Any diamond, no matter how colorless in appearance, receives a D grade if it appears to have less color than the E master stone. Thus, no D master stone is necessary.

Set-Up of References in the Viewing Environment.

At the laboratory, a “working master set” of the 10 master stones needed to grade the most commonly submitted diamonds, D through M range, is typical-
ference between two or three adjacent masters (e.g., the E, F, and G masters or the I and J masters). To overcome this challenge, GIA has multiple graders independently grade diamonds submitted for master stone reports (as is the practice for regular grading, too) and uses instrumentation for support. In assembling a new master set, our goal is to create a group of stones that meet this visual tolerance. Such subtle appearance differences relative to a GIA master have not been found to adversely affect the use of sets within the laboratory or by our clients.

Note that there may be differences in the color grade indicated on a GIA Diamond Grading Report and a Master Color Comparison Report; that is, if the diamond was within visual tolerance to the slightly higher side, it could receive a different grade than its master designation. For example, a diamond accepted as an L on a Master Color Comparison Report might be perceived during random grading to be slightly to the high side in the visual tolerance range of a GIA master. In that case, the diamond could be graded K on a Diamond Grading Report (see, e.g., figure B-2).

As a final note, master stones require special care and maintenance. It is particularly important to clean them regularly. The GIA Laboratory routinely boils master stones in sulfuric acid every two to four weeks, depending on the frequency with which they are used, to minimize the potential influence of foreign surface material. Boiling is an especially critical part of master stone maintenance for diamonds with bruted girdles. When outside sources have returned sets for us to review or supplement, the laboratory has seen up to four grade shifts in appearance for diamonds with bruted girdles that have not been boiled for some time.

Constant handling of diamonds can result in minor damage, so this must also be monitored. If a master stone has noticeable chips or is badly worn, its color appearance may be affected. The laboratory uses rubber-tipped tweezers to greatly reduce the risk of damage (these tweezers also reduce the accumulation of surface material mentioned above).

Figure B-1. For almost 70 years, GIA has been building diamond “master sets” as comparators for use in color grading. Since the mid-1950s, such sets have been at the core of the GIA diamond grading system for colorless to light yellow diamonds. The diamonds in the set above range from E to Z. There is no D master; a D-color diamond is one that has less color than the E master. Photo by Robert Weldon.

Figure 9. GIA master stones are located at the highest point in their respective grade range. A diamond equal to the G master is graded a G. If it has slightly less color, it would receive a grade of F. A diamond with more color than the G master and less than the H master would receive a G grade. A diamond with less color than the E master is graded a D. A diamond with more color than the Y-Z master is graded face-up as a fancy color.
ly kept in the viewing box in a 12 in. (30 cm) long V-shaped nonfluorescent white plastic tray (figure 10), with the E master on the left. The portion of the tray on which the diamonds sit is 1 in. (25 mm) wide and the backing is ¾ in. (19 mm) high. This size creates a consistent background for the diamonds. The length of the tray allows enough room between diamonds for the grader to handle them, as well as to focus on color comparisons between specific pairs of diamonds. With this arrangement, a grader can efficiently move through the grading process with an assigned quantity of diamonds without the added time of taking out and replacing the master stones.

While many diamond dealers and manufacturers take out only one or two reference diamonds at a time to compare to a diamond of unknown color, such a procedure is impractical for the production needs of a grading laboratory. It also may require that the observer rely on color memory in the decision making process. Studies have shown that color memory is not reliable for subtle color comparisons (Epps and Kaya, 2004), and having several master stones in the viewing environment at all times eliminates this problem.

Viewing Geometry. The visual complexity and often extremely subtle color of a polished diamond can make the grading of color very challenging. Therefore, the primary observation direction for color grading a diamond in the D-to-Z range is through the pavilion facets, with the diamond in the table-down position in order to reduce the complex, mosaic-like appearance seen face-up. The grader sits with his or her eyes approximately 12–15 in. (30–37.5 cm) from the diamond, closely adhering to a standard “0/45” geometry between the observer, the light source, and the diamond’s pavilion facets (figure 11). The tray holding the diamonds is positioned 8 in. (20 cm) beneath the fixed light source.
During examination, the tray, while remaining on the upper shelf base of the viewing box, is rocked slightly—such that the line of sight varies from approximately perpendicular to the pavilion facets to near-perpendicular to the girdle (figure 12).

The visual comparison of round-brilliant diamonds to round-brilliant master stones eliminates one significant variable: shape. The situation is more complicated, however, when color grading fancy-shape diamonds (figure 13). Most fancy shapes can display up to three distinct amounts of color depending on their orientation in the table-down observation position. We have found that the most representative set-up for color grading fancy shapes is to orient them with their long and short axes at approximately 45° to the observer (figure 14). In this position, the outline of the fancy shape most closely resembles that of the round brilliant (i.e., reduces shape comparison differences) and functions as the best visual “average” for the amount of color observed. It exhibits neither the most intense color appearance nor the “washed out” areas.

From our experience, it is difficult to distinguish subtle differences between the colors of two diamonds that are touching one another. Therefore, we place a diamond close to (no more than 1⁄5 in. or 5 mm) but not touching a master stone when making color observations. The diamond being graded should also be placed in the same line as the master stones, not in front of or behind them, so that they are all the same distance from the observer. Fancy-shape stones also should not be so close that their

Figure 13. Fancy-shape diamonds have long played an important role in diamond jewelry. The color grading of fancy shapes is challenging due to the variations in proportions. GIA uses a standard viewing position for all fancy shapes to average the appearances encountered. The necklace consists of 44.06 carats of F-to-H oval diamonds; the J-color emerald-cut diamonds in the cuff links weigh 3.04 and 3.01 ct. Courtesy of Louis Glick & Co; photo by Harold & Erica Van Pelt.

Figure 14. When grading fancy shapes, the average appearance is best represented by placing the fancy shape with its long axis approximately 45° to the observer. This series of photos for a 1.20 ct emerald cut next to a GIA master stone illustrate why this “middle” position is used. On the far left, the emerald cut is positioned so the observer views the long side—its weakest color appearance. The photo on the far right illustrates the appearance seen when the grader looks through the end of the fancy cut, where the color appears strongest. The angled position used at the laboratory, which averages the color, is seen in the center. Other fancy shapes also best simulate the outline of the round when set in this position. Photos by Robert Weldon.
angled position causes them to overlap the master stone along the observer’s line of sight (figure 15).

It is important to acknowledge that some years ago (in particular the 1970s and ’80s), laboratory staff experimented with placing diamonds in different positions and at different distances from the light source in the color-assessment process (K. Hurwit, pers. comm., 2007). Whether the diamond was a round brilliant or a fancy shape, there were times when observations were made through the crown with the diamond face-up and through the pavilion with the diamond on its side, in addition to the primary direction: through the pavilion with the diamond table-down. In each of these positions, observations of color appearance were made through a wide range of viewing angles. Mentally averaging the appearances encountered through the combination of directions was used in an effort to ascertain differences between the subtle colors of a diamond and a master stone. Ultimately, it was determined that using multiple positions further complicates decision making and the repeatability of the color determination.

For consistent results across many observers and locations, the laboratory restricts the positions in which diamonds are observed. For round brilliant cuts in the D-to-Z range, color is graded table-down only. Because fancy shapes toward the lower end of the D-to-Z scale typically appear to have more face-up color than their round-brilliant counterparts, at or below Q a combination of table-down and face-up is used to balance the grade and acknowledge the more noticeable face-up color. At Z, face-up color determines whether a diamond is a fancy color.

Just as the diamond being graded and the master stone were put in a number of different positions in the past, the viewing conditions have varied, too. In the 1970s and 1980s, the color tray that fit over the well of the microscope for use with the microscope’s overhead light was recommended to members of the trade who did not have a DiamondLite. Occasionally, staff members also used it for color grading. During that same period, GIA’s Gem Instruments division added a small recessed opening at the top front of the DiamondLite. When the cover to this opening was raised, a “color grader” tray could be placed in the opening (which was in front of the lamps). With this configuration, the light from the lamps was filtered through the plastic tray, thus minimizing the diamond’s internal and surface reflections. Trade members and staff in the laboratory occasionally used this upper recessed tray area to observe color, similar to the way some diamantaires breathe on a diamond to “fog it” to minimize reflections.

The only location currently used for color grading at the GIA Laboratory is a V-shaped tray on the upper shelf in the viewing box (again, see figure 11). We do not alter the appearance of the diamond by filtering the light or breathing on the stone. All observations are made without magnification.

**Determining the Diamond’s Color Grade.** GIA grades the overall color appearance of a diamond. Attention is not focused on specific areas, such as the center of the pavilion of a round brilliant or the long flat side of an emerald cut. By observing the overall appearance, the grader mentally blends all the visual sensations of the diamond.

Instead of trying to match the color of a diamond with a reference color, the GIA system involves placing or bracketing the color between pairs of master stones, which for most observers is an easier task. In general, the grading process is one of progressively narrowing the range until the diamond fits within a single grade (i.e., more color than the master stone on the left, and less color than the master stone on the right).

After the diamond to be graded has been wiped clean with a lint-free cloth, it is initially placed at one end (far left—the colorless end—by laboratory convention) of the tray on which the master stones are set in the viewing box. Using a pair of rubber-tipped tweezers, the grader moves the diamond along the set of master stones until it appears to be
one to two grades past the estimated color grade. It will, at this location, appear to have noticeably less color than the master stone to its left. The grader then moves the diamond back by placing it consistently to the right side of each master for comparison. When the diamond being graded appears to have less or the same amount of color as one master stone, and more color than the next master stone to its left, it has arrived at a single color grade range. Its grade is associated with the least colored of the two diamonds, since each master stone represents the highest (least colored) boundary marker in the range.

Some color grades in the D-to-Z scale may not appear to be different at first glance (for example, D, E, and F diamonds all appear virtually colorless). Therefore, it can be challenging for a grader to clearly place the diamond being graded between two master stones through the bracketing process (it may be located much closer to one of the masters). In this situation, it is common to identify the closest master stone, and then determine to which side of that master the diamond being graded should be placed. In making this determination, the grader places the diamond in one of two grade ranges that are separated by a master stone and then observes the diamond on each side of that master. This will result in one of a number of appearance relationships, the five most common of which are described in table 1 along with the corresponding grading decisions.

Master Eye Effect. The procedures detailed in table 1 were instituted to compensate for the phenomenon known as “master eye effect” (highlighted row in table 1) and the very subtle visual deviations in color assessment it may cause. The effect is described as follows: When two diamonds are very similar in color appearance, the amount of color appears to reverse as the position of the diamonds is switched from left to right and right to left (Liddicoat, 1993). Much has been written about the dominance of one eye over the other in human binocular vision (see, e.g., Kromeier et al., 2006). What has long been described as the “master eye effect” in color grading is related to some degree to the difference in perception between the left and right eye. It is likely there are also psychological influences, such as the starting point used by a grader.

The effect of the master eye can be compounded in color grading by the production requirement of having a set of master stones in the field of vision when grading. The overall appearance of this set is that of a color gradation. If the master stones that form the color gradation are too close together, this can affect the grader’s perception of appearance relationships. To confirm this phenomenon, in the late 1990s we created a gradation of light-toned yellow color chips (simulating the D-to-Z range), and placed the chips in the viewing environment with one of them being a duplicate. Graders found that the undisclosed duplicate appeared to have less color than its twin when placed to the side of its twin next to chips with increasing color, and more when placed to the side of chips with lessening color (figure 16).

This tendency held true whether the gradation was arranged so the chips with less color were to the left or right of the observer. Recognizing this effect further strengthened the laboratory’s desire to

<table>
<thead>
<tr>
<th>Appearance of diamond as compared to closest master stone</th>
<th>Grading decision</th>
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<tbody>
<tr>
<td>Left side of master stone</td>
<td>Right side of master stone</td>
</tr>
<tr>
<td>Slightly more color than master stone</td>
<td>AND noticeably less color than master stone</td>
</tr>
<tr>
<td>Same amount of color as master stone</td>
<td>AND less color than master stone</td>
</tr>
<tr>
<td>Slightly more color than master stone</td>
<td>AND slightly (and to the same degree) less color than master stone</td>
</tr>
<tr>
<td>Same or slightly more color than master stone</td>
<td>AND same or slightly less color than master stone</td>
</tr>
<tr>
<td>Noticeably more color than master stone</td>
<td>AND slightly less color than master stone</td>
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transition to a longer viewing tray. Graders can now create much wider spacing between master stones to minimize the impact of the color gradation.

**Additional Considerations for Color Grading.** Color Grading Diamonds that Differ Significantly in Size from the Master Stones. Overcoming the visual effect of size differences between the diamond being graded and the master stone is an additional challenge even for the most experienced grader (figure 17). To aid in making this determination, graders observe an overall blend of color, similar to that previously described for colored diamonds (King et al., 2005), rather than select visual details. Through experience, the grader also learns how to gaze simultaneously at both the master and the diamond being graded. In so doing, the blend of color in each diamond is easier to relate regardless of size.

Color Grading Fluorescent Diamonds. While some obviously fluorescent diamonds can appear different under differing conditions (Moses et al., 1997), our goal is to report the colors of all D-to-Z diamonds under one standard set of conditions. Therefore, fluorescent diamonds are graded using the same viewing environment and geometry as for other diamonds in the D-to-Z color range. In fact, all diamonds are color graded before they are checked for fluorescence strength, as we have noted that occasionally the color appearance of a diamond will change temporarily when exposed to UV radiation. As a result, color graders do not know the degree of fluorescence in a diamond before they assess its color.

Color Grading Diamonds with Eye-Visible Clarity Characteristics. While not common, the laboratory occasionally encounters diamonds with large, extensive, and/or colored inclusions that affect or obscure the bodycolor when observed under normal color grading conditions. In these instances, the color grade includes the effect of the inclusions. Noticeable inclusions become blended into the overall appearance such that, for example, dense areas of dark inclusions result in the diamond having a gray appearance (figure 18, left). If inclusions are restricted to a small area, their effect is limited (figure 18, right), because the diamond can usually be positioned so as to minimize the visual impact of the inclusions for grading purposes.

There are also times when the lab encounters diamonds with stains in fractures. If the stain is so prominent that it affects overall appearance, the lab will not grade the diamond because all or a portion of the stain might be removed by boiling in sulfuric acid, a procedure commonly used to alter the appearance of diamonds with surface-reaching fractures. Such a diamond will be graded only after the client has boiled it and the stain has been removed, as that is considered its permanent state.

Diamonds with dense clouds of tiny particles or whitish graining may appear translucent in the color grading process. If the transparency is greatly

Figure 16. To better understand the effect of surrounding colors in making comparisons, color experiments were performed using standard color chips. In this photo, the third and fourth chips from the left are the same color. Graders found that the undisclosed duplicate appeared to have less color when placed to the side of its twin with increasing color and more when placed to the side with lessening color. This effect can be minimized in the grading process by spacing the color comparators farther apart on a long viewing tray. Photo by Jian Xin Liao.

Figure 17. When grading diamonds significantly larger or smaller than those in the master set, the observer must look at the overall blend of color rather than the details. Here, a 10+ ct round brilliant is positioned next to an 0.70 ct J master. The best comparison process in this situation is for the experienced grader to observe the overall blend of the two diamonds simultaneously, rather than switching between the two diamonds, so that subtle color differences stand out. Photo by Robert Weldon.
affected, the diamond is graded as a colored diamond and described as Fancy white. At the laboratory, a master diamond is used for this comparison, but a simple method to help understand the approximate color-grade boundary is as follows: If the transparency is so affected that a grader cannot readily observe the pavilion facets through the table of the diamond using a 10× loupe under standard conditions, it is too translucent to grade on the D-to-Z scale and should be described as Fancy white. If it is not too translucent, it follows standard D-to-Z color grading procedures (figure 19). The movement of translucent diamonds off the D-to-Z scale is similar to the movement of light yellow diamonds past Z: In both cases, it is the degree of color, or translucency, seen table-up that determines whether the diamond enters the Fancy range. A translucent D-to-Z diamond may receive virtually any color grade.

Grading Mounted Diamonds. Prior to the 1980s, the laboratory graded mounted D-to-Z diamonds, reporting the grade in a two- to three-grade range. Over time, the laboratory decided to discontinue this practice and issue reports only on unmounted diamonds. Currently, we examine mounted diamonds solely as part of a “confirmation process”—that is, to confirm it is the same diamond as one described on an existing report—within the laboratory’s Verification Service. Because the subtle color appearance of D-to-Z diamonds makes consistent color grading in mountings very challenging (since they can be affected by the color of the surrounding metal), the lab uses grade ranges (e.g., H to J) in these instances. For the Verification Service, the range is described only on “in-house” documents as part of the identification process required to match a diamond to a GIA report.

Color Grading Brown or Gray Diamonds in the D-to-Z System. From its inception, the D-to-Z system included near-colorless to light brown diamonds. Prior to and throughout the 1980s, the use of yellow master stones for brown diamond comparisons was a common procedure. At that time, the brown diamonds typically submitted for grading reports were in the E-to-J range. While there is a noticeable difference in hue, brown diamonds in this range share tone and saturation qualities with their yellow counterparts (for a discussion on the three attributes of color—hue,
tone, and saturation—see King et al., 1994). This made the visual comparison to yellow masters compatible for brown diamonds in these letter grades.

With the influx of stones from Australia’s Argyle mine since the mid-1980s, there has been greater industry awareness and marketing of brown diamonds (Richardson, 1991). As a result, more brown diamonds have been submitted to the GIA Laboratory, not only in the near-colorless region but throughout the color grade scale. Accordingly, the laboratory created a master set of brown diamonds beginning at K (figure 20). As these stones become darker, the differences in hue, tone, and saturation are more pronounced. This contributed early on to the laboratory's decision to begin associating a word description with the letter grades of brown diamonds beginning at K (figure 21). Today, a letter grade plus word descriptions of “Faint brown,” “Very Light brown,” and “Light brown” are used for the grade ranges of K–M, N–R, and S–Z, respectively.

The color transition between brown and yellow diamonds is continuous, and the laboratory occasionally encounters diamonds with color appearances that are “in-between” the two different colors of the master sets (e.g., yellow-brown). It is important to choose the appropriate set of masters (i.e., yellow or brown) for the comparison process. This is usually accomplished by comparing the diamond being graded to both sets and selecting the one closest in appearance.

We recognize that others in the industry do not have D-to-Z scale brown master sets (and grading brown master stones is not a service the GIA Laboratory currently offers). Assessing the color of brown diamonds using only yellow master stones can be challenging. When doing so, the observer must remember to assess the overall depth of color—the combined effect of tone (lightness to darkness) and saturation (strength or weakness) of a color (King et al., 1994). Some observers try to grade just as they would yellow diamonds, and only look for saturation differences (the “amount” of yellow), which can result in an incorrectly high determination compared to laboratory grading. If yellow master stones are the only ones available, the observer should assess the overall depth of color and equate it to the overall depth of the yellow master stone.

The reporting approach for gray diamonds is similar to—but not the same as—that used for browns. In the colorless to near-colorless range (E to J), they are graded using the D-to-Z scale letter grades. Beginning at K, though, gray diamonds receive a word description only of “Faint,” “Very Light,” or “Light” gray for the same letter grade ranges as for brown diamonds (King et al., 1994). Although gray diamonds are reported with only word terms in this range, historically they have not been considered a “fancy” color until they reach a description of “Fancy Light” (as with yellows and browns).

Color Grading at the Lower End of the D-to-Z Range. Color grading at the lower end of the scale (below N or O) can present special challenges for graders. As the color becomes more noticeable, so do the differences between color attributes. In determining the relationship of a diamond to a master stone, an observer must contend with subtle differences in tone (lightness or darkness) and hue (as opposed to the predominance of saturation in the decision making for other areas of the scale).

The difficulty in making grade distinctions between single color grades in this range limits the usefulness of all the individual color grades in the O-to-Z range. More importantly, we have found that such fine distinctions are not in demand among the laboratory’s clients; nor are they significantly useful to the trade for valuing these diamonds. We have informed clients that reporting color grades in this portion of the grading scale by using grade ranges is the best solution. The master stone locations used for laboratory reporting are O, Q, S, U, W, Y, and the Z/Fancy Light boundary. Therefore, GIA grading reports will note a color as “S–T range” or “Y–Z range,” for example.
As mentioned previously, round brilliants are graded table-down up to Z on the color grading scale, but face-up observation increases in importance when we are grading fancy shapes. From our experience, the majority of yellow fancy shapes graded Q or lower table-down appear to be one or more grades lower than this when observed face-up (figure 22). Historically, this led us to assign a final grade that averaged the two appearances when both diamonds fall on the D-to-Z scale. At the transition boundary between the D-to-Z scale and fancy colors, face-up appearance becomes the single factor that determines the color grade; that is, a diamond that has a stronger face-up color appearance than the Z/Fancy Light boundary master stone is considered a fancy color regardless of the color observed table-down.

(For a detailed discussion of the transition of yellow diamonds from the D-to-Z scale to the terminology for colored diamonds, see King et al., 2005.) Over a period of months in the late 1990s, the laboratory researched ways to increase consistency of grading yellow fancy shapes in this part of the scale while acknowledging the relationship of the two observation positions. Working from the known face-up location of the Z/Fancy Light boundary, staff members made table-down and face-up comparisons for hundreds of fancy-shape yellow diamonds. These data were used to establish the relationship between the two observation positions. At that point, the laboratory selected a series of fancy-shape diamonds (figure 23) that would represent the face-up fancy shape boundary for the reported Light yellow grade.

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**COLOR GRADE TERMINOLOGY BOUNDARIES**

<table>
<thead>
<tr>
<th>Yellow</th>
<th>Brown</th>
<th>Gray</th>
<th>Other</th>
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</thead>
<tbody>
<tr>
<td>D</td>
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<td>E</td>
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<tr>
<td>G</td>
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<td>Colored Diamond Color Grade</td>
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<td>H</td>
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<td>I</td>
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<tr>
<td>J</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K</td>
<td>Letter Grade Only</td>
<td></td>
<td></td>
</tr>
<tr>
<td>L</td>
<td>“Faint Gray”</td>
<td></td>
<td></td>
</tr>
<tr>
<td>M</td>
<td>“Faint Brown”</td>
<td>“Very Light Brown”</td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>“Very Light Gray”</td>
<td></td>
<td></td>
</tr>
<tr>
<td>O–P</td>
<td></td>
<td>“Light Gray”</td>
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<tr>
<td>Q–R</td>
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<td>S–T</td>
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<td>W–X</td>
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<tr>
<td>Y–Z</td>
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<tr>
<td>&gt; Z</td>
<td>Colored Diamond Color Grade</td>
<td></td>
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</tbody>
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Figure 21. Shown here are the various boundaries at which colors transition off the D-to-Z scale, as well as terminology associated with the scale. After K, reports note a word description and letter grade for brown diamonds and a word description only for grays. Yellow, brown, and gray diamonds transition to the colored diamond color grading terminology after Z; all other colors transition at G.
Transitioning from the D-to-Z Scale to Fancy Color Grades for Colors other than Yellow, Brown, and Gray. The occurrence of subtle colors other than yellow, brown, or gray is so rare that the presence of even slight tints is acknowledged in their color grading. When colors such as blue, pink, or green are equivalent to G or lower (i.e., the amount of color has moved out of the colorless range and into the near-colorless range), colored diamond color grading terminology is applied to describe that diamond (e.g., Faint pink or Faint blue [figure 24]). Figure 21 notes the location, in relation to the D-to-Z scale, where alternative terminology is applied for these colors as well as for brown and gray.

D-TO-Z INSTRUMENTAL COLOR MEASUREMENT
Colorimetry was introduced into gemology in England in the 1930s (“Measurement . . .,” 1933; “The standardization of colour,” 1933), but it was restricted to use with colored stones. Diamonds, with their often-subtle color differences, were more challenging. Robert Shipley Sr. envisioned the use of instruments for the color measurement of diamonds (Shipley, 1940) and, as mentioned earlier, introduced a visually comparative colorimeter in 1941. It was soon in full use at GIA for the grading of master stones. However, as Shipley Sr. recalled later (Shipley, 1958, p. 136), he was concerned because “the facets of the diamond were still pronounced, with the color varying over the observed portion of the stone. In other words, the facets being observed broke the color into a mosaic of varying intensities and this mixed pattern made it quite difficult to match the other half of the field [the portion of the wedge being viewed] with the diamond, since there was no single block of color to match against.”

In 1949, GIA instructor Joe Phillips developed an electronic colorimeter that employed a selenium photoelectric cell. Because it measured the relative transmission of yellow and blue light by a diamond, it was referred to as a distimulus (i.e., two stimuli) colorimeter. It was fairly effective but too expensive to produce commercially, and Phillips failed to resolve a number of other problems. While it was eventually abandoned, its design became the starting point for a small colorimeter developed by Robert Shipley Jr. several years later (see, e.g., GIA, 1962).

Shipley Jr. demonstrated his colorimeter at the 1956 AGS Conclave. Designed for use by AGS members in their stores, the new distimulus colorimeter had several limitations: It did not accurately grade stones with a greenish or brownish cast or those that were poorly cut. Large diamonds (over 5 ct) were also problematic (“Operating and maintenance instructions . . .,” n.d.; Sloan, 1956; GIA, 1962), as were highly fluorescent diamonds, since this instrument used an incandescent bulb with virtually no UV component. Even so, this electronic colorimeter was soon in use. Shipley Jr.’s colorimeter expanded the AGS color scale from VI to X and encompassed 11 AGS grades (AGS, 1965); however, it was not used for GIA grades (even though the 11 grades spanned the full 23-color D-to-Z color grading scale). The GIA Laboratory only used the electronic colorimeter to grade masters for AGS members (who used a 0 to 10 scale, not D to Z) and to check the calibration of colorimeters being supplied to AGS members by GIA.
By 1975, the costs and difficulties of repairing Shipley’s colorimeters rendered them obsolete. Given the limitations of the instrument, GIA concluded it was not a suitable foundation for further developments in this area. Others in the industry did pursue such instrumentation, and Eickhorst patented a device using more advanced technology in 1974 (Eickhorst and Lenzen, 1974).

Over the course of the next 30 years, GIA researchers evaluated a number of color-measurement devices already in the marketplace that had been developed for various applications, including gemstones. They tested several colorimeters and spectrometers extensively and, for reasons such as lack of reproducibility or efficiency, concluded that none served the laboratory’s purpose. In 1997, GIA made the decision to use its laboratory and research resources to develop a color measurement device of its own design for internal use. It sought a device that would mimic the visual D-to-Z color grading methodology as closely as possible.

Work started on this project in March 1998. In early 1999, the first instrument was constructed and put into use in the laboratory. For approximately one year, measurement data were collected in tandem with the visual grading results on thousands of diamonds. The statistical analysis of these data showed a good correlation between instrumental and visual color grades, and minor modifications to the device and measurement protocols continued to bring results even closer together. In addition, modulating the UV content in the light source allowed the laboratory to obtain reliable color measurement results for diamonds with obvious fluorescence that were very similar to those obtained with visual color grading. By mid-2000, the color grading accuracy of the device was similar to that of the laboratory graders, but with higher repeatability. Around this time, the laboratory began to use several of these devices to support the graders’ opinions. This “instrument” opinion was not influential in the grading decision, but it helped support the visual grade determination and avoid errors. The process was started with stones below 2 ct and eventually expanded to larger sizes.

This approach was followed for the next year until the device’s ability to perform accurate color grading had been validated. In 2001, following its application in the grading of tens of thousands of diamonds, we integrated the device as a “valid” opinion in the grading process, with visual agreement by one or more graders required to finalize the color grade of a particular diamond. Since then, the vast majority of diamonds passing through the laboratory have been graded by combining visual observation with instrumental color measurement. Note that this instrument is for the laboratory’s internal use and is not available commercially.

**SUMMARY AND CONCLUSIONS**

Over the course of more than half a century, the D-to-Z diamond color grading system has become a critical component in the valuation of gem diamonds worldwide (figure 25). At the close of the background section, we noted that “using the same color grading terms does not constitute adhering to the conditions or methodology of the GIA system.” We trust it is now clear that there is much more involved than the D-to-Z scale alone. The GIA system requires the use of standardized viewing conditions, calibrated references, and consistent...
procedures to achieve sound, repeatable results. And it recognizes the importance of having standard policies and procedures not only for the majority of cases that are encountered but also for those seen less frequently. Special approaches must be taken when the diamond being graded is significantly different from the master stones used in the laboratory. Larger diamonds, fancy shapes, those with a hue other than yellow, heavily included diamonds, and borderline fancy-color stones all require specific protocols to ensure the highest level of consistency and accuracy in the color grading process.

Technologies have evolved and will continue to evolve, so it is important to stay abreast of changes that may prove helpful in establishing color grades based on the original historic choice of those grade ranges. Some of the important advances have been the move from incandescent to fluorescent lighting in the viewing environment, the development of a viewing environment that maximizes the efficiency of the observer, and refinements in the system to accommodate increasing numbers of diamonds at the lower end of the scale. As it entered the 21st century, GIA developed color grading instrumentation to support the visual grading process.

Although daylight is the historical and universal standard for diamond observation, in reality no artificial light duplicates natural “daylight,” which itself changes with time and location. Nevertheless, we believe that a standard light source for diamond color grading should have key characteristics of daylight, including a UV component.

For GIA, any future updates in its diamond grading system must show a high correlation to past results in order to have merit. Nevertheless, we recognize that lighting technology and the understanding of human perception are constantly evolving, and believe that research is critical to maintaining the fundamental integrity of the system.

Figure 25. Color is so critical in the valuation of diamonds that diamond manufacturers must estimate the resulting color from the rough when calculating for the best yield. The octahedron pictured at top weighs 15.98 ct, while the macle weighs 22.33 ct. The faceted diamonds, ranging in color from D (the round on the left) to K (the oval), all weigh between 3.00 and 3.50 ct. Photo by Robert Weldon.

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Since late 2007, rubies and sapphires have been mined by hand methods from both eluvial and primary deposits at Winza in central Tanzania. The gem corundum is related to “dikes” of amphibolitic rocks that belong to the Paleoproterozoic Usagaran Belt. Based on crystal morphology, Winza corundum is subdivided into two types: prismatic-tabular-rhombohedral and dipyramidal. In general, medium red and dark (orangy) red top-quality rubies are rhombohedral. Pinkish red and purplish red rubies, as well as pink, purple, and blue (often strongly color zoned) sapphires are, for the most part, dipyramidal. The top-quality rubies are characterized by a distinct assemblage of long tube-, fiber-, needle-, or hair-like inclusions containing an orange-brown material (most likely limonite). The lower-quality material generally contains a larger amount of solid inclusions (mostly amphibole crystals), fissures, and growth features. Unique to corundum from this locality are bluish violet color zones oriented parallel to the prism and basal pinacoid, and occasionally also parallel to rhombohedral and dipyramidal faces. The relatively high Fe content of Winza rubies separates them from most other natural and almost all synthetic counterparts.

In early 2008, several very fine faceted rubies (e.g., figure 1) arrived in the market from a new deposit near the village of Winza in central Tanzania (Dimitri Mantheakis, pers. comm., 2008). These stones created considerable excitement, as the supply of fine unheated rubies had been scarce for years. In the ensuing gem rush, about 6,000 people had moved into the Winza area by July 2008. Although a few exceptional gems were recovered, for the most part the deposit has produced large quantities of ruby and pink-to-blue sapphire of much lower quality; very rarely, “padparadscha” sapphire is also found. Much of the material has been purchased and traded by Thai and Sri Lankan gem dealers who established buying offices in the nearby town of Mpwapwa. This report describes the mining and geology of the deposit, and provides a detailed gemological characterization of the rubies and sapphires.

Additional photos of the Winza mining area and photomicrographs of internal features are available in the G&G Data Depository (www.gia.edu/gemsandgemology). Also, more images and information will be available on author VP’s Winza webpage at www.fieldgemology.org, beginning in mid-February 2009.

HISTORY

According to Druml et al. (1992), ruby was first discovered in Tanzania in the early 1900s near Longido, in the northern part of the country (figure 2). At Longido and other similar Tanzanian ruby-producing localities such as Lossogonoi, the corundum is found in “anyolite,” a rock composed of green zoisite and dark green to black amphibole. In 1960, rubies and sapphires were discovered near the Kenyan border in the Umba River valley (Hänni, 1987), where desilicated pegmatites intrude serpentinite.
During the 1970s, rubies and spinels were found associated with marbles in primary or secondary deposits at two main areas in the Morogoro region of central Tanzania (Hänni and Schmetzer, 1991). The first area, located just east of the town of Morogoro near the village of Matombo, produced mainly during the 1980s. The second area is located to the south of Morogoro, near the mountain city of Mahenge. This area is quite large, and numerous ruby mining operations were known at Lukande, Mayote, Chipa, Epanko, Kitonga, and Kitwaro. The area is still being worked, but it is no longer as active as it was during the 1980s (Pardieu, 2005, 2007).

In the southern part of Tanzania, gem corundum was also found near Songea (in 1993) and Tunduru (in 1994; Milisenda et al., 1997), and these areas are still active today. However, mining declined after the discovery of the large gem placers at Ilakaka in Madagascar in 1999.

With the exception of some remarkable stones primarily from the Morogoro deposits, Tanzania was historically known to produce mostly cabochon-quality rubies or stones that required heat treatment (Dirlam et al., 1992). Therefore, the recent discovery of high-quality rubies at Winza is a welcome event for the Tanzanian gem trade.

Prior to the ruby rush, the Winza area was sparsely inhabited by maize farmers. The original discovery of ruby and sapphire there is shrouded in mystery, but some local residents report that gems were actually mined for years by a farmer who kept his activities secret. The farmer reportedly visited Dar es Salaam regularly to sell his stones. After the farmer died, his young son continued his business but was not able to keep the secret. In November 2007, Tanzanian traders learned about Winza rubies and came to the area to mine. Local reports aside, we do know that mining activities have apparently taken place in the Winza area since at least the 1950s, as evident from a Mpwapwa district map published in that decade (and seen by one of the authors [VP] in Tanzania); it showed mine symbols near the location of the present corundum deposits. Another author (JMS) recalls that in the late 1960s, large quantities of well-crystallized opaque brown-gray corundum crystals measuring up to ~10 cm were produced from near Mvomero, which is located ~100 km east of Mpwapwa, but none of the material showed potential for gem use.

By December 2007, about 600 diggers and brokers from various mining areas and gem markets in Tanzania had moved to Winza to work the deposit for ruby and sapphire. In January 2008, one of the gem brokers (Abdul Msellem) told author VP about the new deposit. At the February 2008 Tucson gem show, Tanzanian dealer Dimitri Mantheakis informed author BML that more than 1,000 miners were active in a “gem rush” to the area.

In April 2008, author VP visited Winza with gemologist Jean Baptiste Senoble, and in June author BML visited the mining area with Dr. James Shigley of GIA Research. At the time of these two visits, about 5,000 miners were working the deposit. There were numerous brokers on-site, and several buying offices had been set up in the town of Mpwapwa. By July 2008, there were more than 100
foreign buyers (mostly Thai and Sri Lankan) in Mpwapwa and about 6,000 miners at Winza (A. Msellem, pers. comm., 2008). As of December 2008, the Tanzanian government was processing applications for 600 small-scale mining licenses (8 × 10 hectares each) in the Winza area (V. Komu, pers. comm., 2008).

LOCATION AND ACCESS

The Winza mining area is located 120 km (by air) from the capital city of Dodoma (again, see figure 2). From Dodoma, the small town of Mpwapwa is reached in about 2 hours by following a paved road for 48 km and a dirt road for 60 km. From Mpwapwa, a dirt road leads 85 km to Winza village (07°00’58” S, 36°21’54” E) and another 10 km to the mining area (07°05’03” S, 36°19’11” E). The travel time from Mpwapwa is 2.5–3 hours in a four-wheel-drive vehicle during the dry season (from approximately May to October). During the rainy season, deep mud makes the road difficult to impassable, and it may take several days to reach Winza.

The mining area is sometimes referred to as “Mtakanini” (meaning “what do you want?” in Swahili) after the name of a nearby hill. Foreigners are not allowed to visit Tanzanian gem mining areas without proper authorization from government and local authorities.

GEOLOGIC SETTING

The Winza area belongs to the Paleoproterozoic Usagaran Belt (figure 3), a rock unit composed of highly metamorphosed basement rocks, metasediments that have undergone a lower grade of metamorphism, and felsic magmatic intrusives and volcaniclastic sediments that are nearly unmetamorphosed and undeformed. Structurally, the Usagaran Belt constitutes the eastern border of the Archean Tanzania Craton (see, e.g., Fritz et al., 2005).

The eastern parts of the Usagaran Belt were reworked during the Neoproterozoic East African Orogeny and have been designated the Western Granulites, while further to the east, a unit of younger Mesoproterozoic rocks composed of enderbite gneisses (metamorphosed igneous rocks of the charnockite series), schists, and marbles has been named the Eastern Granulites (Fritz et al., 2005). The Eastern and Western Granulites were metamorphosed simultaneously, and together they constitute the Neoproterozoic Mozambique Belt of Tanzania and Kenya.

Most of the known gem corundum occurrences in Tanzania and southeastern Kenya—including Longido, Umba, the Mangari area, Morogoro, Mahenge, Tunduru, and the Umba River valley.
corundum crystals embedded within dark-colored amphibolite. Weathering of the primary deposits resulted in an overlying soil horizon (eluvial deposit) that also contains gem corundum.

MINING AND PRODUCTION
As of July 2008, mining in the Winza area was performed by several hundred small groups organized around a local miners’ association. The miners constructed a settlement adjacent to the mining area, near a seasonal river (Mtindiri) that supplies water to wash the soil and for personal use. Both the eluvial and primary deposits have been worked (figure 4). The eluvial soil was excavated with picks and shovels, and taken to the river for washing. The soil was transported in bags by hand or on bicycles; a few operations also employed hand carts or pickup trucks. Although no mechanized mining was taking place, one of us (BML) saw a jigging apparatus in the mining area that was awaiting installation.

The miners use small pumps and simple screens to wash the soil (figure 5), and then pick out the gems by hand. There is little or no water in the river during the dry season, so the miners dig pits in the river bed and build dams to create small pools for their washing activities.

Work on the primary deposits was initiated in March 2008. At the time of the authors’ visits, tunnels up to 30 m deep had been excavated. Material was transported to the surface using buckets that were raised by windlasses or simply by pulling ropes...
Local miners informed author VP that the best-quality rubies were recovered from the eluvial deposits, although author BML was told by a similar source that at least one exceptional stone was produced from one of the tunnels in the primary deposit. During both visits, the production from the primary deposits was reported to be lower compared to the eluvial soil. As of October 2008, however, the eluvial deposits appeared to be mostly exhausted, but sapphires continued to be mined from a few tunnels in the primary deposit (A. Msellem, pers. comm., 2008).

Most of the Winza production is purchased at the mines by Tanzanian brokers, who typically sell the stones in Mpwapwa to foreign buyers. The buyers have constructed several offices and their advertisements are seen throughout Mpwapwa, making it reminiscent of the gem rush towns of Tunduru in southern Tanzania and Ilakaka in southern Madagascar.

Most of the Winza rough is brought to Bangkok and Colombo for distribution into the world market. Initially, heat treatment of Winza ruby was not very successful: Thai dealers reported that the stones turned orangy red without a significant improvement in color or clarity. Nevertheless, during the Gübelin Lab’s off-premises testing activities at the Hong Kong Jewellery & Watch Fair in September 2008, author DS identified the first heat-treated gem-quality rubies from Winza that were submitted by customers for certification by GGL. These stones (up to ~5 ct) displayed an orange modifying hue.

Although not abundant, some very clean and highly transparent rubies—which do not need heat treatment—have been discovered at Winza (again, see figure 1). Rough material of this quality has been mistaken as synthetic by some buyers because of its strong color and transparency. Synthetics are the main problem local buyers encounter when purchasing Winza stones in Tanzania (e.g., figure 7). Also of concern is the presence of other gem materials, such as pink-red spinel, in parcels of Winza ruby.

**MATERIALS AND METHODS**

For this study, we characterized 289 rubies and sapphires from Winza by a variety of techniques. A description of the samples and the methods by which they were tested is given in table 1.

At GGL, we used a Topcon refractometer, with a near-sodium equivalent light source, to measure refractive indices and birefringence. Specific gravity was determined by the hydrostatic method. The fluorescence behavior to standard 365 nm long-wave and 254 nm short-wave UV radiation was observed in a darkened room. Internal features were observed with standard gemological microscopes (Bausch & Lomb and Schneider Steini 2000 with Zeiss optics). Mineral inclusions were analyzed using a Renishaw Raman 1000 spectrometer with an Ar+ (or green) laser at an

**Figure 5.** The gem-bearing soil is washed using water from Mtindiri River. The miners dig pits in the riverbed to make small reservoirs, and the water is pumped through a network of hoses to the washing screens. Photo by V. Pardieu/GGL.

**Figure 6.** Ruby and sapphire mining in Winza is now concentrating more on the primary deposit located under the gem-rich soil. Mining takes place in pits that follow near-vertical amphibolite “dikes” (one is visible here between the two workers). Photo by V. Pardieu/GGL.
### TABLE 1. Materials and methods used in this study of Winza gem corundum.

<table>
<thead>
<tr>
<th>Sample groups</th>
<th>Heated?</th>
<th>Color Description</th>
<th>Morphology and growth structures</th>
<th>Standard gem properties</th>
<th>Internal features</th>
<th>Associated inclusions</th>
<th>UV-Vis-NIR</th>
<th>FTIR</th>
<th>EDXRF</th>
<th>LA-ICP-MS</th>
<th>Isotope analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1 No</td>
<td></td>
<td>1 matrix specimen</td>
<td>-</td>
<td>-</td>
<td>1 (EMPA), 1 (LA-ICP-MS)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>A2 No</td>
<td></td>
<td>25 rough</td>
<td>25</td>
<td>-</td>
<td>3 (EMPA), 1 (XRD)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>A3 No</td>
<td></td>
<td>70 (56+14) windowed</td>
<td>10</td>
<td>12</td>
<td>70 (Raman), 6 (LA-ICP-MS)</td>
<td>35 48 70 26 3</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>B No</td>
<td></td>
<td>60 faceted</td>
<td>20</td>
<td>8</td>
<td>60 (Raman)</td>
<td>11 11 60 20 -</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>C Yes</td>
<td></td>
<td>33 faceted</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>33 33 -</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>D No</td>
<td></td>
<td>5 faceted</td>
<td>-</td>
<td>5</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>E No/yes</td>
<td></td>
<td>3 pairs windowed</td>
<td>-</td>
<td>6</td>
<td>-</td>
<td>6</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>F Yes</td>
<td></td>
<td>3 faceted</td>
<td>-</td>
<td>3</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>G1 No</td>
<td></td>
<td>25 matrix specimens</td>
<td>25</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>G2 No</td>
<td></td>
<td>50 rough</td>
<td>50</td>
<td>-</td>
<td>5</td>
<td>4 (SEM) 1 (XRD)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>H No</td>
<td></td>
<td>14 faceted</td>
<td>-</td>
<td>14</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

*Notes:
- A3: 56 samples were selected (based on color and transparency) from about 500 pieces of reportedly unheated rough obtained from miners or brokers in Mpyawpa and Anusha by one of the authors (VP) in April 2008. Fourteen samples were selected from rough material donated by various companies located in Bangkok, Colombo, and Idar-Oberstein. All samples were transparent and suitable for cutting. After two parallel windows were polished on each sample, they ranged from 0.32 to 8.71 ct.
- B: Most are from various companies in Thailand (~0.5–13.3 ct); includes >10 top-quality rubies submitted by GGL customers during and after the 2008 Basel Fair.
- C: Acquired from various companies in Thailand from April to June 2008 (~0.4–1.3 ct). All were represented as being from Winza, and their properties were consistent with the rough samples that were obtained in Tanzania by the authors.
- D: Examined microscopically by author DS in Bangkok in May 2008 (~0.5–13.3 ct). All were represented as being from Winza, and their properties were consistent with the rough samples that were obtained in Tanzania by the authors.
- E: Three untreated samples (~1.5–2.5 ct, orange red and purplish pink) were each cut into two pieces, and then one piece from each sample was submitted for heat treatment in Bangkok in June 2008.
- F: Submitted to the GIA Laboratory in Bangkok for reports (1.56–11.51 ct).
- Analytical technique used for identification is shown in parentheses. Abbreviations: EMPA = electron microprobe analysis, LA-ICP-MS = laser ablation-inductively coupled plasma-mass spectrometry, Raman = Raman microspectroscopy, SEM = scanning electron microscopy (with energy-dispersive spectroscopy), XRD = X-ray diffraction.
excitation wavelength of 514 nm. Polarized ultraviolet-visible-near infrared (UV-Vis-NIR) absorption spectra were taken with a Perkin Elmer Lambda 19 spectrometer, in the range of 280–880 nm. Unpolarized mid-IR spectra (5000–1500 cm$^{-1}$) were collected using a Philips PU9624 Fourier-transform infrared (FTIR) spectrometer and a DRIFTS beam condenser, at a resolution of 4 cm$^{-1}$ and with 200 scans.

Also at GGL, semiquantitative energy-dispersive X-ray fluorescence (EDXRF) chemical analysis was performed with a QuanX EC instrument (compare to Schwarz et al., 2000). It was operated using a special set of parameters optimized for the analysis of corundum with various conditions for voltage (six steps from 5–30 kV), lifetime (200–300 seconds), and filter type (no filter, cellulose, aluminum, palladium of different thickness). Laser ablation–inductively coupled plasma–mass spectrometry (LA-ICP-MS) data—approximately four spot analyses per sample—were measured with a Perkin Elmer ELAN DRC-e single collector quadrupole mass spectrometer and a CETAC LSX-213 Nd:YAG laser ablation system. The Q-switched 213 nm laser was set to a 10 Hz pulse rate with an energy of ~30 mJ per pulse of 5-nanosecond duration. All analyses were performed as single-spot (100 μm) depth profiles. In addition to the 53 faceted and 26 windowed corundum samples analyzed by this technique, the minerals in one petrographic thin section (two pale pink corundum crystals, as well as garnet and amphibole from the matrix) were analyzed by both LA-ICP-MS and electron microprobe. Electron microprobe analyses were performed at the Eugen Stumpf Electron Microprobe Laboratory, Universities of Graz and Leoben, Austria, with a JEOL JXA 8200 instrument. Analytical conditions consisted of an accelerating voltage of 15 kV and a 10 nA sample current; matrix corrections were made following the ZAF procedure.

At GIA in Bangkok, gemological properties were measured on 14 rubies (e.g., figure 8) using standard techniques.

Crystal morphology was studied by author KS by taking angle measurements of crystals/fragments using a contact goniometer and by angle measurement in the immersion microscope.

Growth structures were observed in rough, windowed, and faceted samples by microscopic techniques with the samples immersed in methylene iodide. Five black minerals that were attached to or exposed at the surface of these samples were analyzed by scanning electron microscopy with energy-dispersive spectroscopy (SEM-EDS) and X-ray diffraction.

Figure 8. The Winza rubies studied for this report include these samples (2.07–6.09 ct), which were examined at GIA’s Bangkok lab in early 2008. All the stones proved to be unheated. Photo by Suchada Kittayachaiwattana.

Figure 9. These slabs were cut from a Winza rock sample. The slab on the left consists of uniformly fine-grained amphibolite (amph) margins flanking orangy brown garnet (gt), ruby (ru), and sapphire (sa). The slab on the right also contains plagioclase feldspar (fsp). Photos by Patrick Lagrange.
**TABLE 2.** Crystal faces observed in rubies and sapphires from Winza, Tanzania.

<table>
<thead>
<tr>
<th>Habit</th>
<th>Observed crystal form</th>
<th>Dominant face</th>
<th>Subordinate face</th>
<th>Miller index hkl</th>
<th>Angle between c-axis and crystal face</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rhombohedral to prismatic</td>
<td>Basal pinacoid</td>
<td>c</td>
<td>—</td>
<td>(0001)</td>
<td>90°*</td>
</tr>
<tr>
<td></td>
<td>Second-order</td>
<td>a</td>
<td>—</td>
<td>(112)</td>
<td>0°*</td>
</tr>
<tr>
<td></td>
<td>hexagonal prism</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Positive rhombohedron</td>
<td>r</td>
<td>—</td>
<td>(1011)</td>
<td>32.4°*</td>
</tr>
<tr>
<td></td>
<td>Negative rhombohedron</td>
<td>s</td>
<td>—</td>
<td>(0221)</td>
<td>17.6°*</td>
</tr>
<tr>
<td></td>
<td>Hexagonal dipyramid</td>
<td>—</td>
<td>n</td>
<td>(2243)</td>
<td>28.8°*</td>
</tr>
<tr>
<td>Tabular</td>
<td>Basal pinacoid</td>
<td>c</td>
<td>—</td>
<td>(0001)</td>
<td>90°*</td>
</tr>
<tr>
<td></td>
<td>Negative rhombohedron</td>
<td>—</td>
<td>s</td>
<td>(0221)</td>
<td>17.6°*</td>
</tr>
<tr>
<td></td>
<td>Hexagonal dipyramid</td>
<td>n</td>
<td>—</td>
<td>(2243)</td>
<td>28.8°*</td>
</tr>
<tr>
<td>Dipyrudal</td>
<td>Basal pinacoid</td>
<td>c</td>
<td>—</td>
<td>(0001)</td>
<td>90°*</td>
</tr>
<tr>
<td></td>
<td>Positive rhombohedron</td>
<td>—</td>
<td>r</td>
<td>(1011)</td>
<td>32.4°*</td>
</tr>
<tr>
<td></td>
<td>Negative rhombohedron</td>
<td>—</td>
<td>s</td>
<td>(0221)</td>
<td>17.6°*</td>
</tr>
<tr>
<td></td>
<td>Hexagonal dipyramid</td>
<td>—</td>
<td>v</td>
<td>(4483)</td>
<td>15.4°*</td>
</tr>
<tr>
<td></td>
<td>Hexagonal dipyramid</td>
<td>—</td>
<td>n</td>
<td>(2243)</td>
<td>28.8°*</td>
</tr>
</tbody>
</table>

* Based on morphological cell with a : c = 1 : 1.365.

Similar nontransparent-to-semi transparent violet areas in nine rough and windowed samples from GGL were analyzed by electron microprobe (three samples) or LA-ICP-MS (six samples).

Oxygen isotope analyses of portions of three win- dowed GGL rubies (pink to red, deep red with blue banding, and deep red) were performed by author AEF in Scotland using a modification of the laser-fluorination technique described by Sharp (1990). Most analyses were run two or three times to check for isotopic heterogeneity and analytical artifacts. The method involved complete reaction of ~1 mg of powdered corundum, heated by a CO2 laser, with ClF3 as the fluorine reagent. The released oxygen was passed through an in-line Hg-diffusion pump before conversion to CO2 on platinized graphite. The yield was measured by a capacitance manometer, and the gas-vacuum line was connected to a dedicated VG Prism 3 dual inlet isotope-ratio mass spectrometer.

Three untreated samples (~1.5–2.5 ct, orangy red and purplish pink) were each cut into two pieces, and then one piece from each sample was submitted for standard heat treatment in Bangkok in June 2008.

### MINERALOGICAL AND GEMOLOGICAL PROPERTIES

#### Host Rock Constituents.

The matrix samples consisted of corundum crystals embedded within areas of coarse-grained orangy brown garnet that are hosted by granular dark green-black amphibolite (figure 9). This assemblage locally contains areas of plagioclase, with accessory spinel, mica, kyanite, and allanite.

Electron microprobe analyses of the amphibole identified it as alumino-pargasite with a significant amount of chlorine (0.8–1.0 wt.% Cl). The chemical composition varied slightly between the core and rim and within different areas of the sample. A representative chemical formula of an amphibole core is 

\[
\text{Na}_{0.39}\text{K}_{0.12}\text{M}_{1.24}\text{Al}_{2.03}\text{Fe}^{2+}_{0.59}\text{Fe}^{3+}_{0.48}\text{Si}_{5.97}\text{O}_{22}\text{(OH)}_{1.72}\text{F}_{0.06}\text{Cl}_{0.22}.\]

The host-rock garnet was found to consist of 34% grossular, 32% pyrope, 32% almandine, and 2% spessartine, with the chemical formula (Ca1.02Mg0.96Fe0.99Mn0.1)(Al2.0)(SiO4)3.

### Corundum Crystal Morphology.

All the crystals studied showed some highly reflective planar faces, but most also had less reflective, not exactly planar, and somewhat inclined and rounded faces. The latter planes were apparently contact surfaces with other corundum crystals or associated minerals. This was carefully considered when the morphology of the sometimes extremely distorted crystals was determined. Some of the crystals had distinct growth striations on their crystal faces that were oriented perpendicular to the c-axis.

The corundum crystals showed a variety of habits. Some were elongated and mostly broken, whereas others were more equidimensional, essentially complete crystals. Occasionally, elongated forms also had end faces, but complete, well-terminated crystals were extremely rare. Four habits were recognized: rhombohedral (negative and positive), prismatic (long and short), tabular, and dipyramidal. The crystal forms observed for all habits are summarized in table 2. The top-quality rubies were rhombohedral, while the lower-quality corundum was characterized by different morphology.

The rhombohedral and prismatic types were gradational into one another. No intermediate samples,
however, were observed between the rhombohedral-to-prismatic type and samples with dipyramidal habit. We saw only one tabular crystal, probably related to the positive rhombohedral type.

Surprisingly, most rhombohedral crystals consisted of the basal pinacoid $c$ and the negative rhombohedron $s$ as dominant forms (figure 10), rather than $c$ and the positive rhombohedron $r$. The latter habit, with $c$ and $r$, is commonly observed in rhombohedral corundum from Mogok, Myanmar (Bauer, 1896; Melczer, 1902), and from Morogoro, Tanzania (Hänni and Schmetzer, 1991).

The negative rhombohedron $s$ is extremely rare in corundum, though it was mentioned by Goldschmidt (1918) as a subordinate crystal form. To the best of our knowledge, $s$ was not described previously as a dominant crystal form of corundum. The $r$ and $s$ forms can be distinguished by their different inclinations to the $c$-axis, which are $32.4^\circ$ and $17.6^\circ$, respectively.

The angles formed by the basal pinacoid and the two rhombohedral forms were calculated as follows:

- Basal pinacoid $c$ and positive rhombohedron $r$: $c^\prime r = 122.4^\circ$, $r^\prime r = 94.0^\circ$
- Basal pinacoid $c$ and negative rhombohedron $s$: $c^\prime s = 107.6^\circ$, $s^\prime s = 111.3^\circ$

The two angles in corundum crystals with $c$ and $s$ faces are closely related to the characteristic angle of a spinel octahedron (109.5°). Therefore, this type of Winza corundum can also be described as pseudo-octahedral.

We observed only one Winza sample that showed just the $c$ and $r$ forms, but several intermediate samples with both $r$ and $s$ (figure 11). These crystals were often somewhat thick tabular (platy) in habit. Occasionally, the rhombohedral crystals also showed small hexagonal dipyramids $n$. In the one tabular sample, these faces were dominant (along with basal pinacoids) and the rhombohedral faces were subordinate.
In all the prismatic Winza crystals, in addition to the basal pinacoid $c$, the hexagonal prism $a$ was dominant, while $s$, $r$, and $n$ faces were subordinate. According to the relative size of the $c$ and $a$ faces, we observed long-prismatic and short-prismatic crystals (figure 12). We also saw samples with a habit intermediate between rhombohedral and short-prismatic, some with a complicated morphology (again, see figure 12).

In contrast, samples with dipyramidal habit had simple forms. Most broken crystal fragments showed only one dipyramidal face, either $\vartheta$ or less frequently $v$. We also commonly observed $c$, occasionally in combination with small $r$, $s$, and $n$ faces (figure 13). Interestingly, the dipyramidal Winza material also showed the rare negative rhombohedron $s$, but no sample with dipyramidal $\vartheta$ and $v$ faces in combination with the prism $a$ was observed.

The forms described above can be recognized in faceted stones by examining them in immersion and performing growth structure analysis (see below).

**Figure 13.** The dipyramidal Winza ruby and sapphire crystals, such as those shown here (lower right is $7 \times 13$ mm), are dominated by $\vartheta$ faces, which are accompanied by various combinations of $c$, $r$, $s$, $v$, and $n$ faces. Photo by K. Schmetzer.

**Visual Appearance and Gemological Properties.** On the basis of our faceted samples (as well as those seen in the trade), we know that Winza rubies and sapphires range from blue to red. Although some of the rubies show exceptional transparency, most Winza stones have some milkiness due to fissures and mineral inclusions. Both rubies and sapphires commonly have distinct color zoning, often with blue (or, more rarely, yellow or colorless) domains. Fine, evenly colored, saturated red or blue stones are quite rare. Top-quality vivid red rubies are generally very homogeneous in color (figure 14), but they may contain small areas of narrow blue zones. Most of the rubies are pinkish to purplish red to dark (orangy) red with moderate saturation. The sapphires are pink, purple, and blue, typically with moderate saturation (e.g., figure 15). Only a very few blue sapphires (<5 ct) of good quality from Winza have been reported in the Thai market. Orange-pink padparadscha Winza sapphires are even rarer. Our heat-treated specimens were orangy red to orange-red. Some stones cut from Winza material (unheated and heated) show more than one color (e.g., figure 16).

The physical properties of some unheated samples are reported in table 3; there were no variations in their characteristics according to color.

**Growth Features and Color Zoning.** Growth planes were observed parallel to the dominant crystal faces in all morphological types. However, in most of the faceted samples examined, growth zoning was only seen parallel to the basal pinacoid $c$; rarely, it was parallel to several dominant faces (figure 17). Also uncommon was color zoning in completely red or pink samples (i.e., consisting of red or pink layers of various intensity; figure 18).

**Figure 14.** Note the homogeneity of color in this fine unheated 9.15 ct Winza ruby. Courtesy of Hakimi & Sons; photo by Robert Weldon.
The samples typically showed characteristic growth and color zoning consisting of bluish violet lamellae in otherwise uniformly colored red, pink, orange-pink, or pinkish violet corundum. These lamellae were observed occasionally in rhombohedral and prismatic samples, and frequently in dipyramidal rubies and sapphires. Distinct lamellar zoning was easily seen—even with the naked eye—only in prismatic samples. Viewing perpendicular or oblique to the planes of these lamellae caused them to become transparent bluish violet. In contrast, when viewed parallel to the lamellar direction, the lamellae typically appeared black and nontransparent. Only in immersion, with proper orientation of the samples, did the fine lamellar nature of the color zoning become sharp in all samples (e.g., figure 19). Thus, such viewing conditions were necessary to resolve nonstructured color patches into fine lamellar patterns. In samples with bluish violet lamellae oriented parallel to several forms, the appearance of the color zones was rather complex (e.g., figure 20).

In prismatic crystals, the sharp bluish violet layers were typically oriented parallel to the dominant prism (again, see figure 19). The pattern in rhombohedral and dipyramidal samples, however, was more complex. Viewed parallel to the c-axis, the dark bluish violet zone appeared confined to the outer rim of the crys-

---

**Figure 15.** The faceted cushion cut (1.50 ct, courtesy of Mark Kaufman) and dipyramidal crystal (2.3 cm tall, courtesy of Werner Radl) provide attractive examples of sapphires from Winza; photo by Robert Weldon. The inset photos show color variations within a dipyramidal Winza sapphire crystal (1.3 cm tall, photo by Werner Radl) and in various faceted rubies and sapphires (0.61–1.69 ct; photo by K. Schmetzer).

**Figure 16.** Some Winza sapphires have been cut to show more than one color. These heat-treated samples weigh 0.88 and 1.03 ct. Courtesy of Michael Nemeth; photo by Robert Weldon.

---

**TABLE 3.** Physical properties of Winza corundum.

<table>
<thead>
<tr>
<th>Property</th>
<th>Unheated ruby and sapphire</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color</td>
<td>Red, purplish red, pinkish red, orangy red Blue, pink, padparadscha</td>
</tr>
<tr>
<td>Dichroism (ruby)</td>
<td>Orange to orangy red Purple-red to violet-red</td>
</tr>
<tr>
<td>e-ray (or E II c)</td>
<td>1.767–1.771</td>
</tr>
<tr>
<td>o-ray (or E I c)</td>
<td>1.758–1.762</td>
</tr>
<tr>
<td>RI</td>
<td>0.008–0.010</td>
</tr>
<tr>
<td>Birefringence</td>
<td>4.00–4.03 (average 4.02)</td>
</tr>
<tr>
<td>SG</td>
<td>Mostly weak to moderate; top-quality rubies strong to moderately strong Very weak to weak; rarely, inert</td>
</tr>
<tr>
<td>UV fluorescence</td>
<td>Typical chromium spectrum; in part, also Fe³⁺-related features</td>
</tr>
<tr>
<td>Long-wave</td>
<td></td>
</tr>
<tr>
<td>Short-wave</td>
<td></td>
</tr>
<tr>
<td>Spectroscope</td>
<td></td>
</tr>
<tr>
<td>Internal features</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Long (typically curved) tube-, fiber-, needle- or hair-like inclusions of orange-brown color; common in top-quality red material, but not seen in pink, purple, or blue stones</td>
</tr>
<tr>
<td></td>
<td>• Inclusion association: amphibole + garnet + apatite (rare)</td>
</tr>
<tr>
<td></td>
<td>• Partially healed fissures composed of cavities containing mostly a polyphase filling of solid material</td>
</tr>
<tr>
<td></td>
<td>• Fissures containing a grayish white to pale yellow substance resembling flux residues in synthetic rubies</td>
</tr>
<tr>
<td></td>
<td>• Color zoning with thin bluish violet lamellae in both ruby and sapphire; various types of bluish violet layers parallel to the prism and/or the basal pinacoid</td>
</tr>
<tr>
<td></td>
<td>• Rhombohedral twin planes, possibly with intersection tubules in one or more directions)</td>
</tr>
<tr>
<td></td>
<td>• Minute (exsolved?) particles</td>
</tr>
</tbody>
</table>

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RUBIES AND SAPPHIRES FROM WINZA, TANZANIA

GEMS & GEMOLOGY

WINTER 2008
Distinct bluish violet, almost black color zoning is common in Winza rubies and pink sapphires (Hänni and Krzemnicki, 2008; Krzemnicki and Hänni, 2008; Peretti 2008; Senoble, 2008). Abduriyim and Kitawaki (2008) mentioned that this color zoning is similar to the pattern observed in rubies from Mong Hsu, Myanmar. To investigate whether such color zoning might be helpful for origin determination, as well as for separating faceted rubies and sapphires from their synthetic counterparts, the authors examined several samples of ruby and pink, pinkish orange, or pinkish...
violet sapphire with dark bluish violet- to black-appearing color variations (e.g., figure 22). Three different phases were identified:

1. Sapphire, which was black in reflected and transmitted light, but appeared bluish violet in thin slices or fragments (as described above).

2. Amphibole inclusions, which were black in reflected and transmitted light, but appeared green in thin areas. Such inclusions never revealed a fine lamellar or oriented pattern.

3. Black spinel rims on rubies with dipyramidal habit, but no black spinel was seen within the corundum crystals. Three chemical analyses gave a compositional range of $\text{Mg}_{0.5-0.6}\text{Fe}_{0.4-0.5}\text{Al}_2\text{O}_4$, which is intermediate between magnesium spinel and hercynite. Such rims are normally removed during the faceting process.

It was not always possible to distinguish between these three black-appearing phases without a detailed phase determination.

**Twinning and Milky Domains.** Rhombohedral twin planes (figure 23) were common. The samples typically contained one system of “intersection tubules,” but three-dimensional arrangements of these tubules were also observed (figure 24).

Very common in the sapphires were slightly milky domains in the form of clouds, bands, or “growth sectors” (figure 25). In general, their milky appearance was caused by the presence of very tiny gray pinpoint inclusions.

**Inclusions.** Elongate Needles. The most distinctive features observed in Winza rubies were long tube-, fiber-, needle-, or hair-like inclusions (figure 26). These inclusions were especially common in the top-quality Winza rubies. The curved tubes were not observed in pink, purple, or blue samples. They were straight, slightly curved, bent, or even (rarely) spiral-like. They were filled with an orange-brown (probably polycrystalline) solid material, which generally did not give any useful signals when analyzed by Raman spectroscopy. A very large, slightly flattened tube exposed at the surface of a 0.60 ct faceted stone, however, showed an interesting mineral association. Several reflective (probably “unaltered”) sections of the tube provided a clear amphibole
Raman spectrum, while brownish material between the reflective areas was identified as hematite.

Mineral Inclusions. Mineral inclusions were quite common. Compared to rubies/sapphires from many other localities, however, we identified (by Raman analysis or electron microprobe) only a small variety of mineral species in our samples: amphibole, garnet, apatite, and spinel.

By far the most common inclusion mineral observed was amphibole. It was very typical in the sapphires, but observed in only one (top-quality) ruby. The amphibole inclusions showed large variations in color, size, and shape. Many amphibole crystals were nearly colorless, while others were slightly green to dark green or brownish green to nearly black, and often showed strong pleochroism. For the most part, the amphiboles displayed irregular, slightly rounded forms, but well-developed prismatic crystals also were common (figure 27). They occurred as single crystals or were clustered in groups or compact agglomerations. In some cases, they were accompanied by stress fissures in the host corundum.

The garnets were intense orange-yellow (figure 28); they were typically transparent and had well-developed or resorbed forms. Apatite was a very rare inclusion mineral (identified only in two of the top-quality rubies). It formed as colorless, transparent, euhedral crystals that were not oriented crystallographically (figure 29). Spinel (Fe- and Mg-rich) was identified by electron microprobe as small inclusions in corundum, but it was not seen during

Figure 23. Rhombohedral twin planes are common in Winza sapphires. Photomicrograph by V. Pardieu/GGL; magnified 40×, crossed polarizers.

Figure 24. Intersections of the twin lamellae resulted in the formation of linear tubules—typically one system, but also three-dimensional networks as seen here. Photomicrograph by V. Pardieu/GGL; magnified 30×.

Figure 25. Winza sapphires commonly contain slightly milky domains caused by the presence of minute gray pinpoint inclusions. Photomicrograph by V. Pardieu/GGL; magnified 20×, fiber-optic illumination.

Figure 26. Probably the most characteristic features observed in some of the Winza rubies are long tube-, fiber-, needle-, or hair-like inclusions. These inclusions were especially common in the top-quality stones. Photomicrograph by V. Pardieu/GGL; magnified 40×.
the microscopic examination of the samples. It also formed dark rims surrounding dipyramidal rubies (see above). Opaque mineral grains displaying a grayish black metallic luster were tentatively identified as chalcocite \((\text{Cu}_2\text{S})\) by Raman analysis.

Uncommon in the sapphires were reflective particles that were generally accompanied by small, delicate, disk-like inclusions resembling the “thin films” more typical of basalt-related rubies (figure 30, left). Somewhat coarser particles were present in diffuse clouds or in stringer-like formations (figure 30, right).

Very rare were so-called comet tails associated with mineral inclusions (figure 31).

FTIR spectroscopy revealed the presence of various OH-bearing minerals, such as kaolinite, “limonite,” chlorite, and (very rarely) boehmite. These were observed on fissure planes and as components of the
tube fillings. They also may be present in the form of submicroscopic “particles.”

Fractures and Fluid Inclusions. Unhealed fissures were quite common in the commercial-quality material. These fissures were either strongly reflective (mirror-like) or they displayed a slightly frosted appearance caused by the presence of a gray or brown substance (e.g., oxihydrates, identified by FTIR).

Partially healed fissures were often present in medium- to low-quality rubies/sapphires. In general, they showed relatively coarse textures with highly reflective inclusions that were rounded or irregularly shaped; well-developed networks or fingerprint-like patterns were rare. Less common were healed fissures composed of cavities that for the most part were developed as negative crystals containing polyphase fillings with black (opaque), brown, and colorless (transparent, singly and doubly refractive) constituents (figure 32).

That could not be identified by Raman analysis.

Many fissures appeared to be filled with a grayish white or pale yellow solid material (figure 33) that could not be identified. These inclusions may resemble the flux material common in some synthetic rubies.

CHEMICAL COMPOSITION

EDXRF analyses revealed that Winza rubies and sapphires showed a relatively uniform chemical composition. The chromophores Cr and Fe were present in significant concentrations. Cr concentrations fell into the common range for rubies (including the pinkish red and purplish red material) originating from most occurrences: ~0.10–0.60 wt.% Cr2O3. For the blue to purplish blue sapphires, Cr2O3 ranged from ~0.10 to 0.30 wt.%.

The Fe concentration in the rubies was relatively high, with more than 95% of the samples in the range of ~0.30–0.80 wt.% Fe2O3. In very few samples, Fe2O3 reached up to ~1 wt.%. Top-color Winza rubies were characterized by a combination of relatively low Fe2O3 (~0.30–0.40 wt.%) and relatively high Cr2O3 (~0.40–0.60 wt.%). Blue to purplish blue Winza sapphires had Fe2O3 concentrations of ~0.60–0.95 wt.%.

Winza rubies of the best color contained very little or no Ti (typically below the detection limit). For the other samples, TiO2 concentrations were ~0.005–0.020 wt.%. The blue to purplish blue Winza sapphires typically contained ~0.01–0.03 wt.% TiO2; the highest value found was ~0.045 wt.% TiO2.

The V concentration of the rubies and sapphires was generally low, and in many samples it was below the detection limit of ~0.005 wt.% V2O3. Most common were V2O3 contents in the range of ~0.005–0.015 wt.%.
TABLE 5. Chemical composition by LA-ICP-MS of different colors of gem corundum from Winza, Tanzania (in ppmw).a

<table>
<thead>
<tr>
<th>Element</th>
<th>Red</th>
<th>Purple</th>
<th>Orange</th>
<th>Blue</th>
<th>Colorless</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>1.6 ± 1.7</td>
<td>26</td>
<td>1.9 ± 2.2</td>
<td>3.8 ± 3.4</td>
<td>1.5 ± 0.8</td>
<td>34</td>
</tr>
<tr>
<td>Na</td>
<td>3.1 ± 8.1</td>
<td>40</td>
<td>5.0 ± 19</td>
<td>2.5 ± 5.5</td>
<td>0.8 ± 2.4</td>
<td>23</td>
</tr>
<tr>
<td>Mg</td>
<td>39 ± 36</td>
<td>54</td>
<td>36 ± 20</td>
<td>45 ± 36</td>
<td>26 ± 25</td>
<td>55</td>
</tr>
<tr>
<td>Al</td>
<td>519900 ± 2220</td>
<td>54</td>
<td>521500 ± 2190</td>
<td>519500 ± 1640</td>
<td>521900 ± 1660</td>
<td>55</td>
</tr>
<tr>
<td>Si</td>
<td>4500 ± 1210</td>
<td>54</td>
<td>4060 ± 1650</td>
<td>3910 ± 1450</td>
<td>4710 ± 1180</td>
<td>55</td>
</tr>
<tr>
<td>P</td>
<td>21 ± 13</td>
<td>51</td>
<td>17 ± 12</td>
<td>27 ± 14</td>
<td>14 ± 6.6</td>
<td>50</td>
</tr>
<tr>
<td>K</td>
<td>9.3 ± 8.8</td>
<td>12</td>
<td>12 ± 16</td>
<td>7.8 ± 7.2</td>
<td>11 ± 7.5</td>
<td>3</td>
</tr>
<tr>
<td>Ca</td>
<td>38 ± 7</td>
<td>3</td>
<td>42 ± 22</td>
<td>&lt;40</td>
<td>0</td>
<td>55</td>
</tr>
<tr>
<td>Sc</td>
<td>0.060 ± 0.0025</td>
<td>2</td>
<td>0.071 ± 0.025</td>
<td>0.060 ± 0.0025</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Ti</td>
<td>63 ± 63</td>
<td>51</td>
<td>59</td>
<td>30</td>
<td>15</td>
<td>55</td>
</tr>
<tr>
<td>V</td>
<td>2.5 ± 2</td>
<td>54</td>
<td>2.1 ± 1.4</td>
<td>3.1 ± 1.1</td>
<td>1.8 ± 1.3</td>
<td>55</td>
</tr>
<tr>
<td>Cr</td>
<td>2350 ± 960</td>
<td>54</td>
<td>1310 ± 700</td>
<td>820 ± 270</td>
<td>520 ± 270</td>
<td>55</td>
</tr>
<tr>
<td>Mn</td>
<td>1.8 ± 2.6</td>
<td>51</td>
<td>17</td>
<td>7</td>
<td>3 ± 0.2</td>
<td>2</td>
</tr>
<tr>
<td>Fe</td>
<td>2370 ± 670</td>
<td>54</td>
<td>2430 ± 670</td>
<td>3890 ± 990</td>
<td>3070 ± 890</td>
<td>55</td>
</tr>
<tr>
<td>Co</td>
<td>0.072 ± 0.053</td>
<td>19</td>
<td>0.035 ± 0.023</td>
<td>0.056 ± 0.013</td>
<td>0.036 ± 0.018</td>
<td>15</td>
</tr>
<tr>
<td>Ni</td>
<td>3.3 ± 3.8</td>
<td>27</td>
<td>2.4 ± 1.8</td>
<td>4.3 ± 1.8</td>
<td>1.8 ± 1.4</td>
<td>31</td>
</tr>
<tr>
<td>Cu</td>
<td>0.71 ± 0.98</td>
<td>20</td>
<td>1.5 ± 2.7</td>
<td>7.4 ± 13</td>
<td>0.8 ± 2.2</td>
<td>20</td>
</tr>
<tr>
<td>Zn</td>
<td>28 ± 8.7</td>
<td>3</td>
<td>17</td>
<td>&lt;1.7</td>
<td>1.5 ± 0.7</td>
<td>1</td>
</tr>
<tr>
<td>Ga</td>
<td>21.6 ± 5.5</td>
<td>54</td>
<td>23 ± 6.6</td>
<td>28 ± 5</td>
<td>21 ± 7.7</td>
<td>55</td>
</tr>
<tr>
<td>Rb</td>
<td>0.034 ± 0.033</td>
<td>10</td>
<td>0.094 ± 0.13</td>
<td>0.304 ± 0.048</td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td>Sr</td>
<td>0.94 ± 2.4</td>
<td>13</td>
<td>2.8 ± 6.6</td>
<td>1.0 ± 2.1</td>
<td>0.017 ± 0.0085</td>
<td>3</td>
</tr>
<tr>
<td>Zr</td>
<td>0.019 ± 0.004</td>
<td>6</td>
<td>0.023 ± 0.011</td>
<td>0.17 ± 0.016</td>
<td>0.018</td>
<td>1</td>
</tr>
<tr>
<td>Nb</td>
<td>0.11 ± 0.16</td>
<td>10</td>
<td>0.025 ± 0.022</td>
<td>0.034 ± 0.018</td>
<td>0.046 ± 0.034</td>
<td>5</td>
</tr>
<tr>
<td>Sn</td>
<td>0.22 ± 0.13</td>
<td>41</td>
<td>0.25 ± 0.15</td>
<td>0.38 ± 0.12</td>
<td>0.21 ± 0.14</td>
<td>34</td>
</tr>
<tr>
<td>Cs</td>
<td>0.016 ± 0.019</td>
<td>3</td>
<td>0.03 ± 0.016</td>
<td>0.020 ± 0.008</td>
<td>0.005 ± 0.001</td>
<td>2</td>
</tr>
<tr>
<td>Ba</td>
<td>1.4 ± 3.5</td>
<td>11</td>
<td>4.1 ± 6.1</td>
<td>0.86 ± 1.3</td>
<td>0.097 ± 0.051</td>
<td>2</td>
</tr>
<tr>
<td>Ce</td>
<td>0.026 ± 0.017</td>
<td>8</td>
<td>0.084 ± 0.076</td>
<td>0.12 ± 0.0066</td>
<td>0.013</td>
<td>1</td>
</tr>
<tr>
<td>Ta</td>
<td>0.078 ± 0.13</td>
<td>15</td>
<td>0.046 ± 0.032</td>
<td>0.085 ± 0.063</td>
<td>0.18 ± 0.013</td>
<td>6</td>
</tr>
<tr>
<td>W</td>
<td>3.6 ± 7.8</td>
<td>28</td>
<td>1.7 ± 3.4</td>
<td>0.35 ± 0.43</td>
<td>0.47 ± 0.89</td>
<td>7</td>
</tr>
<tr>
<td>Pb</td>
<td>0.20 ± 0.61</td>
<td>30</td>
<td>0.16 ± 0.34</td>
<td>0.30 ± 0.38</td>
<td>0.10 ± 0.14</td>
<td>35</td>
</tr>
</tbody>
</table>

*Notes: ppmw = parts per million by weight; < = value below the detection limit; N = number of values above the detection limit. Ablated material was carried to the ICP by He (5.0) carrier gas at a rate of 0.8 liters/minute (l/min). The plasma conditions of the ICP-MS were optimized to maximum intensity at U/Th ratio ~1 and Th/ThO ratio <0.5. This was achieved using the following parameters: plasma gas flow (Ar) 14.0 l/min, nebulizer gas flow (Ar) 0.85–0.9 l/min, auxiliary gas flow (He) 0.70–0.75 l/min, and RF power 1400 W. In the corundum matrices, the elements Be, B, Na, Mg, Al, Si, P, K, Sc, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, Zn, Ga, Rb, Sr, Zr, Nb, Mo, Sn, Cs, Ba, Ce, Ta, W, Pb, Th, and U were measured quasi-simultaneously in each individual analysis.

TABLE 4. Electron microprobe analyses of two samples of pale pink Winza corundum.

<table>
<thead>
<tr>
<th>Oxide (wt.%)</th>
<th>Wz1_Cor*</th>
<th>Wz2_Corb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al₂O₃</td>
<td>98.80 ± 0.44</td>
<td>98.50 ± 0.18</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.07 ± 0.02</td>
<td>0.05 ± 0.02</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>0.36 ± 0.06</td>
<td>0.39 ± 0.06</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.02 ± 0.02</td>
<td>0.09 ± 0.08</td>
</tr>
<tr>
<td>Total</td>
<td>99.25</td>
<td>99.03</td>
</tr>
</tbody>
</table>

*15 point analyses.  b 6 point analyses.

wt.%; rarely, values up to ~0.02 wt.% were measured.

Ga content, for the most part, was below the detection limit (~0.005 wt. % Ga₂O₃). The highest Ga concentration found was ~0.01 wt. % Ga₂O₃.

Electron microprobe analyses of two pale pink corundum crystals in a rock thin section showed appreciable Fe and significant Cr, along with traces of Ti (table 4).

Figure 33. These irregularly shaped cavities are filled with a grayish white to pale yellow solid substance. Such inclusions may resemble the flux material observed in synthetic gems. Photomicrograph by V. Pardieu/GGL; magnified 60x.
LA-ICP-MS data for Winza corundum are reported in Table 5. Generally, Si, Cr, and Fe were abundant, with concentrations above 500 ppm (0.05 wt.%). Traces of B, Na, Mg, P, Ti, V, Co, Ni, Cu, Ga, Sn, Ta, W, and Pb were commonly present. Na, Mg, Ti, Cr, V, and Fe were variable, while all other elements occurred in quite constant concentrations. A weak positive correlation was observed for the elements V and Ti.

**SPECTROSCOPY**

UV-Vis-NIR spectroscopy of Winza rubies revealed the well-known Cr$^{3+}$ absorption bands at ~405–410 and 560 nm [figure 34, top]. In addition, the spectra generally displayed a strong “background absorption” starting around 600 nm and increasing toward the UV edge. The absorption spectra of blue to purple-blue and padparadscha sapphires were, in general, combination spectra showing the Cr$^{3+}$ absorption features together with a pronounced contribution of Fe$^{3+}$ bands at 377/388 nm and 450 nm (figure 34, middle). More rarely seen was a combination of the Cr$^{3+}$ features with the Fe$^{2+}$-Ti$^{4+}$ charge-transfer band of sapphire at around 700 nm (figure 34, bottom). The Cr “doublet” at 694 nm was visible in all spectra.

The FTIR spectra in the mid-infrared region between 5000 and 1500 cm$^{-1}$ allow for the distinction of four main features (figure 35):

1. A more-or-less pronounced broad band with a maximum at ~3450 cm$^{-1}$ was recorded in 41 of the 59 samples tested (spectrum A).

2. A prominent peak at ~3160 cm$^{-1}$—with accompanying peaks at ~3350, 3240, and 2420 cm$^{-1}$—was recognized in 38 samples (spectrum B). Most of the remaining samples showed a shoulder (sometimes very weak) at 3160 cm$^{-1}$. The 3160 cm$^{-1}$ absorption was completely absent from only a very few samples. It is interesting to note that the “3160 group” (also named “3161-series”; Smith and Van der Bogert, 2006) was most prominent in the high-quality Winza rubies [i.e., in more than 90% of these stones].

3. An additional group of absorption peaks at ~3695, 3670, 3650, and 3620 cm$^{-1}$ was rarely exhibited by high-quality rubies (spectrum C).

4. Distinct band groups in the 3560–3420 cm$^{-1}$ region were detected in only three samples (spectrum D).

The spectral features described above may be present in combination, with varying intensities of the different components [see, e.g., the combination of the band groups in the 3560–3420 cm$^{-1}$ region and the 3160 cm$^{-1}$ absorption in spectrum E, observed in a top-quality ruby]. In two samples, weak peaks related to boehmite were detected at around 2100 and 1980 cm$^{-1}$ [e.g., spectrum A].
The oxygen isotope compositions obtained for three Winza rubies were δ₁⁸O = 4.6‰, 4.7‰, and 4.9‰ (figure 36), giving a mean δ₁⁸O of 4.7 ± 0.15‰. For information about the use of oxygen isotopes to characterize corundum from different deposits, see Giuliani et al. (2007).

HEAT-TREATED SAMPLES

Heat-treated Winza rubies examined by one of the authors (DS) in Bangkok in May 2008 showed a distinct orangy red hue (as seen in the sample on the left in figure 16). The most striking internal feature was the presence of partially healed fissures displaying strong alteration patterns, with a drop-like or network-like melting appearance that is typical of intensely heat-treated rubies/sapphires. In part, these partially healed fissures were associated with altered mineral inclusions. Identical features were observed in the heat-treated rubies submitted to GGL for testing in Hong Kong in September 2008. The infrared spectra recorded for these samples did not show any features related to the presence of OH/H₂O (see Discussion below).

Comparison of the three pairs of unheated/heated corundum showed that the orangy red sample did not change color. However, one of the purplish pink samples became intense red-orange after heat treatment. The second purplish pink sample developed an inhomogeneous color distribution, with orange and violet areas. All the unheated pieces showed the 3160 cm⁻¹ absorption in varying intensity; these bands disappeared from the heat-treated samples.

DISCUSSION

Physical Properties. The measured refractive indices and specific gravity values from Winza (table 3) are consistent with those known for rubies and sapphires from all other localities. The morphology of the crystals is quite variable, and some of the samples have habits not seen previously in material from other ruby/sapphire deposits.

Microscopic Characteristics. Growth Features and Color Zoning. Although some Winza material may bear a superficial resemblance to Mong Hsu rubies, the orientation of the thin bluish violet lamellae within Winza corundum is completely different. Mong Hsu rubies typically have dark cores that are formed by bluish violet layers oriented parallel to rhombohedral and dipyrmaidal faces (Peretti et al., 1995). In the Winza samples, the bluish violet color zoning was oriented parallel to the prism and basal pinacoid, occasionally in combination with bluish violet layers oriented parallel to rhombohedral and dipyrmaidal faces. Such zoning has not been
observed in gem corundum from other localities, to the best of the authors' knowledge.

Inclusions. Our findings are consistent with previous reports of inclusions in Winza corundum (Abduriyim and Kitawaki, 2008; GIT Gem Testing Laboratory, 2008; Hänni and Krzemnicki, 2008; Krzemnicki and Hänni, 2008; Pardieu and Schwarz, 2008; Smith et al., 2008).

Microscopic examination often can be used to distinguish Winza rubies/sapphires from those originating in other genetic environments or geographic localities. The following internal characteristics can be considered locality-specific for rubies/sapphires from Winza: (1) long tube-, fiber-, needle-, or hair-like inclusions of orange-brown color that are straight, slightly curved, bent, or rarely show a spiral-like appearance (restricted to medium red and vivid [orangy] red Winza rubies); and (2) a suite of mineral inclusions composed of amphibole + garnet + apatite + an opaque mineral (tentatively identified as chalcosite, based on Raman data); (3) any of various types of color zoning; (4) partially healed fissures composed of cavities displaying varying shapes (irregular, but also regular and developed as negative crystals) that contain a polyphase filling; (5) fissures containing a grayish white to pale yellow substance that often resembles flux residues in synthetic rubies; and (6) rhombohedral twin planes. The polyphase filling material of (4) shows various colors (black, brown, or colorless) and could not be identified. No Raman signal could be obtained for these fillers or those of (5).

The variety of mineral inclusions observed in Winza corundum is rather small compared to that seen in rubies and sapphires from other localities. The internal mineral association of a ruby or sapphire always reflects the nature of the host rock in which it crystallized. The association amphibole + garnet + plagioclase (+kyanite identified as an accessory mineral) in the host rock indicates that the corundum formed in a metamorphic environment. Rubies from other localities reveal, in general, quite different

Figure 36. Oxygen isotope values for three rubies from Winza are shown with those for corundum from various types of primary and secondary deposits in Kenya and Tanzania. The data are reported in the conventional delta notation relative to V-SMOW (Vienna Standard Mean Ocean Water) equal to 0‰ and are compared with the oxygen isotopic ranges defined for various deposit types, after Giuliani et al. (2005, 2007). The symbol colors correspond to sample colors; white symbols represent colorless sapphires.
inclusion mineral associations (e.g., Henn et al., 1990; Hughes, 1997; Mercier et al., 1999; Simonet, 2000; Schwarz, 2001; and GGL database). Marble-hosted rubies from Myanmar’s Mogok stone tract, for example, contain mostly rutile (needles and/or irregularly rounded opaque crystals of varying size), carbonates, sphene, zircon, apatite, garnet, graphite, spinel, sphalerite, pyrite, pyrophyllite, mica, olivine, pargasite, and anhydrite (the association “rutile + sphene + zircon” is probably locality-specific for rubies from Mogok).

Chemical Properties. SSEF (2008) reported Cr and Fe as the main trace elements, with “little” Ga, and Ti and V below the detection limit of their EDXRF instrument. Semiquantitative EDXRF analyses performed by GIT Gem Testing Laboratory (2008) indicated moderate contents of Cr (0.35–0.68 wt.% Cr₂O₃) and Fe (0.25–0.41 wt.% Fe₂O₃), very low to low amounts of Ti (55–192 ppm TiO₂) and V (from not detectable to 164 ppm V₂O₃), and low-to-moderate Ga contents (64–146 ppm Ga₂O₃). These values are in fairly good agreement with our data.

Comparison with Rubies from Different Genetic Environments. The chemical fingerprint of Winza rubies is quite different from that of rubies originating from marble-type deposits such as Mogok (figure 37A). The main difference is in the iron concentration: Rubies from marble host rocks typically have low iron contents. In Mogok and Mong Hsu rubies, for example, GGL data show that the Fe₂O₃ concentration is normally below ~0.05 wt.%. However, the lower Fe₂O₃ limit we found for Winza rubies is ~0.3 wt.%, and GIT Gem Testing Laboratory (2008) reported 0.25 wt.%. When comparing the Cr and Fe contents of Winza rubies with those of basalt-related rubies from the Thai/Cambodian border region, we found an almost complete overlap of the chemical data (see again figure 37A).

Comparison with Rubies Originating from Other African Deposits. Rubies from Songea, Tanzania, generally have low-to-moderate contents of Ti, V, and Ga—often below the EDXRF detection limit. The highest concentrations of these elements (from the analysis of ~30 samples; Schwarz, 2001) are ~0.03 wt.% TiO₂, ~0.05 wt.% V₂O₅, and ~0.05 wt.% Ga₂O₃. The Cr content of Songea rubies normally varies between ~0.2 and 0.7 wt.% Cr₂O₃ (as compared to ~0.1 wt.% to ~0.6 wt.% [occasionally ~0.8 wt.% for top-quality] for Winza rubies). Songea rubies distinguish themselves by very high Fe contents (this is

valid for the entire Songea corundum production, independent of the bodycolor)—~0.8–2.0 wt.% Fe₂O₃—as compared to ~0.3–0.8 wt.% Fe₂O₃ for Winza rubies. As evident in figure 37B, there is minimal overlap between Songea and Winza in Cr and Fe.

Rubies from Kenya’s Mangari region most commonly show ~0.005–0.15 wt.% Fe₂O₃ (very rarely up
to ~0.2 wt.%). This range, too, only very slightly overlaps the Fe content of Winza rubies (again, see figure 37B). The Cr contents of Mangari rubies fall mostly into the range of ~0.3–1 wt.% Cr2O3, which does overlap those from Winza.

The Fe and Cr contents of Malawi rubies are also quite different from those of Winza (again, see figure 37B), despite the fact that there are some similarities in their host-rock composition (Fe/Cr-rich amphibolite; Dill, 2005). Rubies from the Chimwadzulu Hill deposit have somewhat higher Fe contents (~0.7–1.1 wt.% Fe2O3) and a distinctly higher Cr concentration (~0.75–2 wt.% Cr2O3). GGL data indicate that rubies from both deposits have low Ti (generally <0.03 wt.% TiO2), V (normally <0.02 wt.% V2O3), and Ga (often below detection limit; highest concentration found was ~0.01 wt.% Ga2O3).

There is very little LA-ICP-MS data for gem corundum available in the literature for comparison. Rankin et al. (2003) reported concentration ranges of Mg, Ca, V, Cr, Fe, Cu, and Ga for rubies originating from Longido (Tanzania) and Chimwadzulu Hill. Winza rubies and sapphires have Mg, Cr, Fe, and Ga concentrations that are equal to or slightly lower than the element concentration ranges reported by Rankin et al. (2003). Calcium, V, and Cu are much lower in Winza rubies and sapphires compared to stones from Longido and Chimwadzulu Hill.

**Spectroscopy.** The GIT Gem Testing Laboratory (2008) reported that the UV-Vis absorption spectra of Winza rubies typically show the Cr3+ bands and lines together with Fe3+ absorptions (377/387 and 450 nm) that would be expected for a ruby with high iron content.

The dominant absorption feature in chromium-colored corundum is the presence of intense bands related to Cr3+ at 405 and 560 nm. Additional spectral features observed for rubies are: (1) the so-called “background absorption,” which represents an increasing absorption toward the UV edge; and (2) the presence of Fe3+-related groups at ~375 nm and 450 nm. Rubies with low Fe contents (e.g., marble-hosted Mogok rubies and most rubies from Kenya’s Mangari region) typically show relatively “pure” Cr3+ spectra with variable, but low, UV absorption. The other “extreme” is provided by corundum from Songea. These rubies (and also sapphires of various colors) are characterized by very strong Fe3+-related features (Schwarz, 2001). Rubies from Malawi show, in general, spectra with combined Cr3+/Fe3+ absorption components.

Winza rubies and pink/purple sapphires, as well as rubies from the Thai/Cambodian border region, show the chromium bands and a generally strong “background absorption,” which is influenced by several factors (figure 34A). Features related to Fe3+ are only weakly developed. Blue to purplish blue Winza sapphires display distinct Fe3+-related features (figure 34B), or they show the combination of Cr3+ bands and the Fe2+/Ti4+ charge-transfer absorption around 700 nm (figure 34C).

The padparadscha sapphire that was examined for this report owed its color to Cr3+/Fe3+ absorptions, rather than the combination of Cr3+ and color centers that are seen in “classic” padparadscha sapphires (i.e., from Sri Lanka). GIT Gem Testing Laboratory (2008) described three diagnostic patterns for OH-related peaks in the mid-IR absorption spectra of Winza rubies: Pattern A showed a broad absorption band from 3735 to 3000 cm−1, centered around 3450 cm−1, assigned to goethite. Pattern B showed absorption bands with approximate peaks at 3335, 3242, 3160, 3075, 2459, and 2420 cm−1. Pattern C showed mixed absorption bands from patterns A and B.

Smith and Van der Bogert (2006) commented on the 3161 cm−1 spectral feature (formerly described mainly in natural-color yellow sapphire from Sri Lanka). They indicated that, when present in high intensity, it appears to consist of at least six bands. This so-called 3161 series was attributed by past researchers to OH groups involved in charge-compensation with Si4+. However, Smith and Van der Bogert (2006) suggested that the 3161 series is actually due to structurally bonded OH associated with Mg2+. The average Mg contents measured in our samples were in the range of 30–65 ppm (table 5). The evaluation of whether this concentration is sufficient to support the “Mg-OH model” is beyond the scope of this article.

Balmer et al. (2006) reported that peaks at 3353 and 3242 cm−1 are probably associated with the 3160 cm−1 absorption. They could not confirm whether this group of peaks is related to goethite or to silanol groups (OH group attached to Si4+).

The FTIR spectra of the present study allow the following correlations:

1. All top-quality rubies containing the orange-brown tube/hair-like inclusions had the 3160 cm−1 absorption as the dominant feature.
2. Only one sapphire showed a comparably prominent 3160 cm−1 feature.
3. The 3160 cm−1 absorption was present in almost all of the lower-quality samples, but it
appeared very weak or as a low shoulder when in combination with other bands.

4. Very few samples did not have the 3160 cm\(^{-1}\) feature.

These results imply that, although there is a strong correlation between the orange-brown tube/hair-like inclusions and a prominent 3160 cm\(^{-1}\) absorption, other factors must be taken into consideration. Since there is no clear correlation between other microscopic features and the presence/intensity of the 3160 cm\(^{-1}\) absorption band, the influence of submicroscopic inclusions should be considered.

The characteristic group of four peaks with maxima at 3695, 3670, 3650, and 3620 cm\(^{-1}\) (spectrum A in figure 35) is typically attributed to kaolinite minerals (Beran and Rossman, 2006). This group was found in 17 samples (in four top-quality rubies, this group was weak, while in other samples it was, in part, quite strong).

According to A. Beran (pers. comm., 2008), the broad 3450 cm\(^{-1}\) absorption band (spectrum A in figure 35) is related to H\(_2\)O (submicroscopic inclusions or adsorptive humidity).

OH bands associated with the presence of chlorite-group minerals show typical absorption features in the range between 3560 and 3420 cm\(^{-1}\) (spectrum D in figure 35).

**Geologic Origin.** Observations by two of the authors (VP and BML) of primary deposits at Winza, as well as the examination of corundum-bearing host-rock specimens in the laboratory, showed a close association between corundum and cross-cutting layers or “dikes” of a garnet-bearing amphibolitic rock. Macroscopic observations by authors GG and DO of a sample obtained by JMS (figure 9) clearly showed that, according to its mineralogy, the dike-like body was mafic in composition before it was metamorphosed. Possible lithologies for the protolith include high-alumina gabbrón (layered gabbrons) or leucogabbros. The central part of the “dike” is composed of a metamorphic garnet + hornblende ± plagioclase ± corundum association. Metamorphic conditions estimated by using garnet- amphibole-corundum equilibria (winTWO; Berman, 2007) showed that the metamorphic overprint was 800 ± 50°C and 8–10 kbar.

The color/growth zones of the corundum are commonly complex, which indicates crystal development during a number of stages characterized by variable Ti, Cr, and Fe availability.

Spinel that (rarely) overgrows or is included in the corundum is Fe- and Mg-rich, whereas spinel from the host amphibole is Cr-rich, with up to 32 wt.% Cr\(_2\)O\(_3\) (unpublished electron microprobe data of the authors). The presence of two chemically distinct spinel compositions indicates that two different generations are present. The Cr-spinel is probably a relict of the initial magmatic rock before the metamorphism, while the Fe- and Mg-rich spinel crystallized during the metamorphic stage that formed the corundum.

The amphiboles did not show any preferred orientation in the host corundum; they can be considered protogenetic inclusions reflecting the nature of the host rock. Amphibole inclusions showing various morphologies were commonly present. This is a strong indication that all the corundum morphologies originated from an amphibolite environment.

In the ternary Mg-Fe-Ti diagram that was established by Peucat et al. (2007) to distinguish magmatic from metamorphic sapphires, the chemical fingerprint of the Winza sapphires overlaps the border separating the metamorphic and magmatic population fields (figure 38).

Oxygen isotope values for three Winza corundum samples defined a consistent and restricted δ\(^{18}\)O range of 4.7 ± 0.15‰ indicating that they formed under comparable genetic conditions. This composition suggests two possible origins for Winza rubies (Giuliani et al., 2005, 2007): either in metamorphosed mafic-ultramafic rocks (worldwide δ\(^{18}\)O range of 3.2–6.8‰ n =19), or in desilicated pegmatites within mafic-ultramafic rocks such as plumesites (4.2–6.7‰ n =16). We therefore conclude that the Winza rubies are of metamorphic origin, with a high-alumina meta-(leuco)gabbro as the protolith for the host rock.

Formation of the rhombohedral top-quality rubies with their exceptional combination of large size, intense color, and high transparency requires growth conditions different from those of the lower-quality corundum (characterized by a prismatic-tabular or dipyramidal morphology and the general presence of strong color zoning and abundant inclusions that reduce their transparency). An explanation for the (co-)existence of these two corundum types in the Winza area could be that they represent separate corundum “generations” formed at different geologic times or under locally different pressure-temperature (PT) conditions.

**Separation from Synthetics and Heat-treated Natural Corundum from Other Sources.** When the first Winza rubies appeared on the market, they were praised for
their exceptional color and clarity. At the same time, they caused suspicion and doubt, because the top-quality stones possessed such a high transparency and bright color that many dealers and gemologists who saw them for the first time thought they were synthetic. The identification and separation of Winza rubies from synthetics is generally quite easy—as long as the typical internal features described above are present. These are mineral inclusions such as amphibole, garnet, and apatite; orange-brown “tubes/needles/hairs”; and color zoning in various patterns. Only one inclusion feature could potentially cause confusion: Partially healed fissures consisting of irregularly shaped cavities filled with a grayish white to pale yellow solid material may resemble the flux components seen in some synthetics.

Another powerful tool for the separation of Winza rubies from synthetics (especially if characteristic inclusions are not present) is chemical fingerprinting, notably the Cr/Fe correlation. Elements such as Ti, V, and Ga are of little help because their contents are normally quite low in synthetic rubies/sapphires, as they are in the stones from Winza. The key element for the separation is Fe:

Contents in the range of ~0.3 to 0.8 wt.% Fe$_3$O$_4$—which are typical of rubies/sapphires from Winza—are only rarely found in synthetic rubies. “High-iron” synthetic rubies are known from Douros (flux grown), Ramaura (flux grown), Gilson (flux grown), and Tairus (hydrothermally grown). In a detailed study by Schwarz et al. (2000), however, only very few synthetic rubies were found with Fe$_3$O$_4$ contents reaching the lower limit of the Winza rubies. These included some hydrothermal synthetic rubies and a very few flux synthetics grown by Douros (2 out of 35) and from the (experimental) production of Gilson (2 out of 27). Gilson synthetic rubies have not been produced and marketed commercially. In addition, all the flux synthetic samples examined by one of the authors (DS) showed healed fissures with typical flux patterns. Douros samples also distinguish themselves by high Ga concentrations (~0.04 to 0.1 wt.% Ga$_2$O$_3$). Practically all production from Tairus shows diagnostic internal features (especially pronounced growth structures). Therefore, chemical fingerprinting, when applied with other standard geological tests, is an effective method for the separation of Winza rubies from synthetics.

The same is valid for the separation of heat-treated Winza rubies/sapphires from stones of other sources. Since heat treatment results in the alteration of fissure textures and solid inclusions, it tends to remove locality-specific internal features. Chemical fingerprinting is not affected by traditional heat treatment and can, therefore, be used in the same way as described for unheated stones. We believe that probably less than 1% of the faceted gems from Winza now on the market have been heat treated. This percentage may increase in the future if treaters are successful in applying heating techniques to pink, purple, or pinkish and purplish red starting material.

SUMMARY AND CONCLUSION

The arrival on the market of very fine cut rubies (e.g., figure 39) from a new deposit near Winza in central Tanzania was welcomed by the international gem trade, as the supply of top-quality rubies is scarce. It is difficult to predict if Winza will become a major source of fine rubies. Only time will tell. But if the area consistently produces fine stones and if a market is found for the lower-quality material, this mining region could become more than a gem rush.

Winza rubies and sapphires are hosted by metamorphic rocks of the Usagaran Belt. Observations of the associated mineral assemblages indicate that

Figure 38. This ternary Fe-Mg-Ti diagram shows LA-ICP-MS data for blue sapphires originating from Winza and other deposits (all data points from GGL database). Peucat et al. (2007) defined two population fields for magmatic- and metamorphic-type blue sapphires. Note that the blue sapphires from Winza plot in a range overlapping both fields (this could indicate that melting processes might be involved, related to high-PT metamorphism). Modified from Peucat et al. (2007).
the gem corundum is metamorphic in origin. The corundum is contained in "dikes" consisting mainly of the assemblage amphibole-garnet ± plagioclase. Oxygen isotope data are consistent with an origin in metamorphosed mafic-ultramafic rocks.

The physical properties of Winza rubies and sapphires are quite constant, with RIs and SGs in the same range known for rubies and sapphires originating from other localities. However, Winza rubies and sapphires can be characterized by inclusion features that, in part, can be considered locality specific: long tube-, fiber-, needle-, or hair-like inclusions, containing an orange-brown, solid polycrystalline material (most likely limonite); amphibole crystals showing large variations in shape, along with garnet and possibly apatite and/or an opaque mineral; partially healed fissures containing cavities filled with a polyphase material; and fissures containing primarily a grayish white to pale yellow substance. These internal features provide confirmation of natural origin and are a very strong indication of the host stone's locality of origin. Only the fissures with the grayish white to pale yellow substance may cause confusion, because they can resemble flux material in synthetic corundum. The specific type of bluish violet color zoning seen in rhombohedral and dipyramidal samples has not been previously found in natural ruby or pink sapphire, to the best of our knowledge, so it is also useful for determining a Winza origin.

The chemical composition of Winza rubies and sapphires, which formed in association with amphibolites, is characterized by a relatively high Fe content. This separates them from most natural and almost all synthetic counterparts. Although there is extensive chemical overlap with rubies originating from the Thai/Cambodian border region, the latter display internal features that are totally different from the inclusions in Winza stones.

Figure 39. The Winza area of central Tanzania is a new source of fine untreated ruby, such as the 3.18 ct center stone in this ring. Courtesy of The Collector Fine Jewelry, Fallbrook, California; photo by Robert Weldon.
The Wittelsbach Blue

Rudolf Dröschel, Jürgen Evers, and Hans Ottomeyer

The 35.56 ct Wittelsbach Blue is one of the largest historic blue diamonds ever fashioned. It belonged to the Bavarian House of Wittelsbach and was displayed in the Treasury of the Munich Residence until it disappeared in 1931. It was secretly sold in 1951, “rediscovered” in 1961, and then sold again in 1964 to an undisclosed private buyer. In December 2008, the Wittelsbach Blue was sold at Christie’s London to jeweler Lawrence Graff for just over $24.3 million, a record price for any diamond at auction. This article describes what is known about the Wittelsbach Blue since it was first reported in 1666, and the gemological information released to date on this diamond, which was recently graded Fancy Deep grayish blue. Investigations in the historical archives of Bavaria, Austria, and Spain revealed that there is no archival evidence to support many previous statements about this stone.

Due to their extreme rarity, blue diamonds, even more than colorless ones, have historically epitomized rank and wealth. Two of the largest known blue diamonds were once part of the crown jewels of European monarchies. The Hope diamond (now 45.52 ct) once belonged to the French royal family (Kurin, 2006), and the Wittelsbach Blue (now 35.56 ct, figure 1) was owned by the Bavarian royal family, the House of Wittelsbach.

Like the Hope, the Wittelsbach Blue is an intense steely blue. It is widely accepted that both originated from the Kollur mine in India’s Golconda District (Bauer, 1932; Balfour, 1987). According to its previously published history (e.g., Gaal, 1977; Bruton, 1981, Balfour, 1997; Morel, 2001; Christie’s, 2008a,b), the Wittelsbach Blue arrived in Vienna in 1666 as part of a dowry for a marriage into the House of Hapsburg. In 1722, it passed to the House of Wittelsbach in Munich, again as part of a dowry. It was mounted as a “symbol of dominion and power” (de Smet, 1963, p. 48) on the globe above the Bavarian Royal Crown around 1806 or 1807 (figure 2). During the 20th century, the Wittelsbach Blue was involved in a series of unusual events, which began in 1931 when the Wittelsbach Equity Foundation (German acronym WAF, for Wittelsbacher Ausgleichsfonds) tried in vain to sell it at the Christie, Manson & Woods auction house in London (Christie, Manson & Woods, 1931). From the day of the auction through the next 30 years, the whereabouts of the Wittelsbach Blue were kept secret from the public. In 1951, the WAF secretly sold the Wittelsbach Blue, which was then “rediscovered” in 1961 by Antwerp diamond dealer Jozef Komkommer. In 1964, it was purchased by a private German collector whose identity was not revealed until recently.

During the authors’ investigations in the Bavarian State Archive in Munich, in the Austrian State Archives in Vienna, and in three Spanish historical archives in Madrid, Simancas, and Valladolid, it became evident that many of the accepted “facts” about the Wittelsbach Blue and its history that were published by Antwerp diamond specialist K. de Smet in his book, The Great Blue Diamond, The Wittelsbacher, Crown Witness to Three Centuries of European History (1963), and cited afterwards by many others, had no archival basis. Based on our comprehensive review of the available documents, we can now correct the historical record and report on the true, exciting history of the Wittelsbach Blue as it moved through Europe.
INDIAN ORIGINS

India was the world’s only source of diamonds (Untracht, 1997) until about 1725 (Balfour, 1987), when the first Brazilian mines began to open (Legrand, 1981). Thus, all diamonds in Europe before that date, including the Wittelsbach Blue, must have had their origin in India (Harlow, 1998; Webster, 1994). Most of India’s diamonds, and its largest, were found in a vast area on the eastern side of the Deccan plateau (Bharadwaj, 2002). Many of these diamond sources were located in the former kingdom of Golconda, which lay between the rivers Godavari in the north and Pennar in the south.

Jean Baptiste Tavernier (1605–89), the famed French diamond dealer, visited three mines in the Golconda region between 1630 and 1668, among them Coulour (also spelled Kollur). This famous mine was the source of several historic diamonds, including the Koh-i-Noor, the Hope, the Dresden Green, the Orlov, the Regent, and the Sancy (Kurin, 2006). According to Bauer (1932), it was also the source of “nice blue diamonds.” Balfour (1987, p. 112) wrote that the “Kollur mine, then, appears to be the only . . . known source of Type IIb natural blue diamonds” in India. The identity of the party who brought the Wittelsbach Blue (perhaps still in rough form) to Europe has been lost to the mists of history, but Tavernier is a possibility, given the timing of its arrival.
After investigations in the state archives of Bavaria and Austria during September and October of 1961, Schneider traveled to Spain early in 1962 to begin research in an unidentified Madrid archive (de Smet, 1963). Here, he said, he found the first mention of the Wittelsbach Blue “as early as 1664,” when King Philip IV (1605–1665) of the Spanish line of the House of Hapsburg “ordered the treasurer to gather a dowry for his daughter,” Infanta Margarita Teresa (1651–1673; figure 3), “from new acquisitions of precious stones from India and Portugal... It included a large blue diamond” (de Smet, 1963, p. 16). According to Schneider, he could find no addi-

(tional information on the Wittelsbach Blue in Madrid, allegedly because “in the bloody years of the Spanish Civil War (1936–1939) the last existing documents were undoubtedly lost with the destruction of the Madrid archives” (de Smet, 1963, p. 17).

One of the authors (JE) visited Spain in November and December 2007 to seek some archival basis for Schneider’s statements. These investigations were performed with the support of five Spanish archivists in the General Archive of the Royal Palace in Madrid (which had no record of Schneider’s visit in 1962), in the Archivo General (General Archive) of Simancas, and in the Archivo de la Real Chancilleria (Archive of the Royal Chancery) of Valladolid. No documents could be found to support the contention that Philip’s treasurer had purchased a large blue diamond—or other stones from India and Portugal—“as early as 1664.” In addition, the archivists were unaware of any records being lost during the Spanish Civil War.

Our 2007 research also reviewed the invoices of the silver- and goldsmiths of the Spanish Court in the General Archive of the Royal Palace in Madrid from 1660 to 1669 for any mention of a large blue diamond, without success. Similarly, the invoices of court treasurers Baltasar Molinet, Antonio de León, and Agustín Spinola for 1660–1669, stored in the Valladolid archive, have no mention of a great blue diamond. One document in particular (found in file number 1816-19) seems to rebut de Smet’s contentions. This record dealt with Margarita Teresa’s trip to Vienna in 1666 to prepare for her marriage to Emperor Leopold I (Widorn, 1960). The document includes several quotes from goldsmith Luis de Sausalca, who had set diamonds for Margarita Teresa’s mother, Queen Maria Anna (1634–96), and who would likely have been commissioned to set a stone such as the Wittelsbach Blue (J. Menéndez Trigos, pers. comm., 2007). Again, however, there is no mention whatsoever of such a diamond. Therefore, Schneider’s statements (de Smet, 1963) dealing with Spanish origins of the Wittelsbach Blue must be considered unsubstantiated. The only mention we could find of such a diamond as part of the dowry was in statements made by Margarita Teresa after she had reached Austria.

Vienna. Margarita Teresa married her uncle, Emperor Leopold I (1640–1705), in Vienna in 1667 (Widorn, 1960). Figure 3 shows the young Empress with her daughter Maria Antonia three years after the marriage. The marriage contract, dated December 18,
1663, and currently stored in the Austrian State Archive in Vienna (Marriage contract . . . , 1663), contains no mention of a large blue diamond, so the stone likely was not acquired before this date. In 1673, six years after the marriage, Margarita Teresa died. In her testamentary bequest (figure 4), she stated that her daughter, Maria Antonia, was her sole heiress [Widorn, 1960], with one exception: a precious ornament, which she had brought from Spain, was left to Leopold: “a great breast ornament with a great diamond in the midst.” Though the blue color is not mentioned, this diamond was most likely the future Wittelsbach Blue: No other large diamond is known to have been possessed by Margarita Teresa, and the court officials at the time would not necessarily have recorded the diamond’s color [I. Aguirre, pers. comm., 2008; G. Gonsa, pers. comm., 2008].

Schneider claimed that, on Leopold’s marriage to his third wife, Eleonora Magdalena, in 1676 (after the death of his second wife, Archduchess Claudia Felicitas, heiress of Tirol, that same year), he “was so enchanted by her beauty” that he gave the Empress “all the jewelry which he had inherited” from Margarita Teresa [de Smet, 1963, p. 18]. Schneider also suggested [de Smet, 1963] that Empress Eleonora Magdalena gave the Wittelsbach Blue by testamentary bequest to her granddaughter, Archduchess Maria Amalia (1701–1756); he based this on documents relating to the “Trousseau of Archduchess Maria Amalia.” However, investigations in the Austrian State Archive between August 2006 and July 2008 failed to uncover any records supporting Schneider’s claims.

Nor does Empress Eleonora Magdalena’s testamentary bequest, written between 1711 and 1720, make any mention of leaving the Wittelsbach Blue to Maria Amalia. During her lifetime, Eleonora Magdalena distributed her jewels to her daughters and also one piece to her son Joseph (Maria Amalia’s father), who had succeeded Leopold as Emperor in 1705, but she did not itemize gifts for her granddaughters. She left such decisions to her heirs. However, other documents in the Austrian State Archives may lead to a new explanation for the transfer of the Wittelsbach Blue to Maria Amalia.

Archduchess Maria Antonia (figures 3 and 5),
daughter of Margarita Teresa and Leopold, married Bavarian Elector Maximilian II Emanuel Wittelsbach (1662–1726) in July 1685. But it was not a happy marriage, and in early 1692, her husband left Munich to become governor of the Spanish Netherlands in the midst of the Nine Years War (1688–1697) and she returned to Vienna. According to Maria Antonia’s “General Inventarium” (General Inventory . . ., 1685; figure 6), prepared at the time of her marriage, she then possessed all the jewelry that her mother had brought from Spain in 1667 as her dowry. This 1685 document listed the large ornament with “a thick stone in the midst,” using nearly identical language as in her mother’s 1673 bequest (figure 4), though still with no mention of the blue color. Although this ornament, again certainly the future Wittelsbach Blue, had been left to Leopold by Margarita Teresa, it seems clear that he passed it to his daughter for her dowry rather than giving it to Eleonora Magdalena as Schneider suggested.

This is further supported by Maria Antonia’s testamentary bequest of 1692 (figure 7), which stated that the gold jewelry her mother brought from Spain should stay at the Bavarian court. However, by the time of her death in December of that year, the “Bavarian court” no longer existed as such—Maximilian II Emanuel had by then moved his household to Brussels. Though he returned briefly in 1701, the War of the Spanish Succession, which began that year (and in which Bavaria and Austria fought on opposing sides), would largely keep him away from Munich until the Treaty of Utrecht in 1713. During all this time, Maria Antonia’s jewelry remained at the Bavarian court.
Hapsburg court in Vienna, where it had been at her death, effectively under Emperor Joseph’s control. Thus, these documents may explain why the future Wittelsbach Blue became part of Maria Amalia’s dowry in 1722, when she, Emperor Joseph’s second daughter, married Bavarian Crown Prince Charles Albert (1697–1745).

CROWN JEWEL OF BAVARIA

Munich. Charles Albert was the son of Maximilian II Emanuel and his second wife, Princess Therese Kunigunde Sobieska. In the inventory of the marriage contract (figure 8), it was agreed that Maria Amalia would bring from Vienna to Munich “gems, jewels, and ornaments,” and that inventory documented for the first time “a large blue brilliant, encircled with small brilliants.” The official bridal portrait of Maria Amalia, painted in 1722 by Frans van Stampart (Glaser, 1976) and currently exhibited at Castle Hämelschenburg in Emmerthal, Lower-Saxony, shows Maria Amalia wearing a hair ornament set with a blue diamond that is clearly identifiable as the Wittelsbach Blue (figure 9). This painting is the oldest visual record of the diamond.
After the 1742 coronation of her husband, Emperor Charles VII, now-Empress Maria Amalia wore the Wittelsbach Blue in a crown made to resemble the Ottonian imperial crown (Ottomeyer, 1979; figure 10; the Ottonian dynasty was a line of Holy Roman emperors during the 10th and 11th centuries).

Maria Amalia died in 1756 and in 1761 her son, Elector Maximilian III Joseph (1727–1777), had the Wittelsbach Blue mounted in a badge of the Order of the Golden Fleece, surrounded by large white and yellow diamonds (Schatzkammer der Münchner Residenz, 1937; Brunner, 1970, 1977; figure 11). This badge is currently displayed in the Treasury of the Munich Residence, though with the Wittelsbach Blue replaced by a glass imitation (Ottomeyer, 1979). A January 1774 inventory of all jewels stored in the Munich Treasury described the “Carat 36” Wittelsbach Blue as the most precious gem in the collection, with a value of 300,000 guilders (see the G&G Data Depository at www.gia.edu/gemsandgemology).

In January 1806, the kingdom of Bavaria was founded with Maximilian I Joseph as its first king. The new royal Bavarian crown (figure 12) prominently featured the Wittelsbach Blue in the orb under the cross, representing the heraldic blue color of the House of Wittelsbach. It occupied this unique position from 1807 to 1931.

European crowns typically have a main stone that exceeds all other personal possessions of the ruler in size, color, and value. The Bavarian crown was a state symbol of a constitutional monarchy, and its legal possession was an outward sign of the legitimacy of the king. In Bavaria, the crown and other regalia, such as the sceptre, sword, orb, and the like, served as symbols of the sovereignty of the kingdom.
new kingdom during the opening of parliament, the oath of each new King of Bavaria, and other state ceremonies (unlike the practice in other monarchies, the Bavarian crown was not used in coronations or worn by the king). Thus, the jewel on the top of the crown had a meaning beyond its material existence (Puhle, 2006).

The crown was designed in 1806 by Charles Percier, a famous Paris architect and designer for Emperor Napoleon I. Actual construction was by goldsmith Martin-Guillaume Biennais and his craftsmen. The jeweler Borgnis in Frankfurt served as an intermediary to match existing gems in the treasury with new acquisitions to complete the ring of pearls and the settings of various secondary stones (Ottomeyer, 1979; Erichsen and Heinemann, 2006).

The Munich Residence was the city palace of the Bavarian dukes, electors, kings, and emperors from 1508 to 1918. The Treasury of the Munich Residence, located on the ground floor in the eastern part of the Royal Palace (figure 13), is among the foremost of such collections in Europe (Heym, 1999). Until 1931, the most precious gem stored here was the Wittelsbach Blue. In a general bill (“Generalrechnung”) from 1807, the Wittelsbach Blue was appraised at 300,000 florins, as much as all other royal ornaments combined (Ottomeyer, 1979).

DISAPPEARANCE AND REDISCOVERY

London. In 1918, at the end of the First World War, the Kingdom of Bavaria was replaced by the democratic Free State of Bavaria. After years of difficult negotiations, a contract was arranged in 1923 between the House of Wittelsbach and the Wittelsbach Equity Foundation (WAF). All former properties of the House of Wittelsbach, including the inventory of the Treasury of the Munich Residence, were transferred to the WAF to be displayed in public museums. In an appendix, it was stated that the sale of any property belonging to the WAF required the approval of the Bavarian state government. In 1931, the House of Wittelsbach and its head, Crown Prince Rupprecht, were faced with grave financial problems (“Verkauf . . .,” 1931), so much so that the WAF decided to sell the Wittelsbach Blue and various other jewels through the auction house of Christie, Manson & Woods (later Christie’s) in London (Christie, Manson & Woods, 1931). The Bavarian State Government, under minister president Dr. Heinrich Held, gave the required export permission for the Wittelsbach Blue. However, at the London auction on December 21, 1931 (figure 14), bidding failed to reach the reserve price, and the Wittelsbach Blue was not sold (Bruton, 1981). From the day of the auction until 1961, the whereabouts of the diamond were not publicly known.

Figure 12. From 1807 to 1931, following the establishment of the Kingdom of Bavaria by Maximilian I Joseph in 1806, the Wittelsbach Blue sat at the top of the royal Bavarian crown. As with the badge in figure 11, it has since been replaced by an imitation made of blue glass. Photo by J. Evers.

Figure 13. The Royal Palace of the Munich Residence was built over 10 years, from 1823 to 1832. The Treasury of the Munich Residence is on the first/ground floor, to the right of the entrance as one goes into the building. It was home to the Wittelsbach Blue during most of the 19th and early 20th centuries. Photo by J. Evers.
Antwerp, Brussels, and Bruges. Acting in secret, the WAF sold the Wittelsbach Blue in Antwerp in 1951 to a merchant specializing in jewels and ornaments (Kuballa, 1964; Biehn, 1965). Baron Teuchert, speaking for the administration of the WAF, later stated that they did not know the name of this buyer because he was represented by an agent (“Der Blaue Wittelsbacher . . .,” 1964). Later in 1951, the Wittelsbach Blue was resold to Antwerp diamantaire Romi Goldmuntz (Kuballa, 1964; figure 15), one of the world’s most prominent diamond dealers (de Smet, 1963; “Er rührte . . .,” 1972; Laureys, 2006).

Recently, Dr. Gerhard Immler, director of the Bavarian Secret House Archives, shared with one of the authors (JE) his research into the sales of precious art and gems by the WAF in the early 1950s. At the time, the WAF needed funds to repair damage to their properties that was sustained during World War II. The most direct method for obtaining these funds was by selling art in the WAF’s collection. The responsible ministries and their leading secretaries, as well as the Bavarian minister president, supported the decision (G. Immler, pers. comm., 2006).

However, the financial situation of the WAF in the 1950s, as judged by statements made by members of the House of Wittelsbach during a court case in 1962 (“Wer hat denn . . .,” 1962), do not seem to bear out a dire need to sell the diamond. Then, the funds of the WAF were estimated to be DM 300–500 million. The value of the Wittelsbach Blue in 1951, based on events during the 1960s (see below) was probably about DM 1 million. Compared to the estimate of the WAF’s total funds, an additional DM 1 million gained by selling the Wittelsbach Blue—the most precious piece of the Munich crown jewels—seems to be a very small improvement, even in a strained financial situation.

From April 17 to October 19, 1958, the Universal and International Exhibition (EXPO’58) was held in Brussels. Forty-seven Belgian jewelers, including Romi Goldmuntz and J. Komkommer & Son, displayed their gems at the Belgian Diamond

Figure 14. In 1931, the Wittelsbachs were forced to put the Wittelsbach Blue (apparently once again mounted in the Golden Fleece ornament in figure 11) and numerous other items of jewelry up for auction. Shown here is the diamond’s entry in the auction catalogue; it did not sell. Note that the carat weight as given here is incorrect.

Figure 15. Romi Benjamin Goldmuntz (1882–1960) was one of the greatest diamond dealers in the world. In 1951, he purchased the Wittelsbach Blue in Antwerp, after it was sold in secret that year by the WAF. Photo by Beurs van Diamanthandel.
Industry pavilion (Exposition Universelle et Internationale de Bruxelles, 1958, 1961; Official Guide..., 1958). There, among other gems, Goldmuntz displayed the Wittelsbach Blue—without giving its name or origin. Nor was it known at the time that Goldmuntz was the owner of the large blue diamond. Further, though the Wittelsbach Blue was one of the largest and most historic blue diamonds in the world, not one of the millions of EXPO visitors—including diamond expert and later purchaser Jozef Komkommer—appeared to recognize it (de Smet, 1963).

After Goldmuntz’s death in 1960, his heirs discovered the diamond in his gem collection, thinking it nothing but a large old-mine-cut stone. Not knowing the origin of the blue gem, in August 1961 they asked Antwerp jeweler Jozef Komkommer to recut it into a “pear-shape, emerald-cut or oval” (de Smet, 1963, p.7). However, contrary to de Smet’s claims, Komkommer may not at first have realized what they had brought him. In a January 2008 interview with Jan Walgrave, former director of the Provincial Diamond Museum Antwerp, Komkommer’s son Jacques claimed that he initially suggested a probable historic origin for the blue gem (J. Walgrave, pers. comm., 2008). Father and son compared characteristic data for historic diamonds in The Diamond Dictionary (Copeland et al., 1960), and in doing so quickly identified the unknown blue diamond as the missing Wittelsbach Blue. Together with some closely associated jewelers, Jozef Komkommer purchased the Wittelsbach Blue on August 28, 1961 (Burgerwelzijn, 1962; V. De Boi, pers. comm., 2007).

Soon thereafter, Komkommer contacted the Treasury of the Munich Residence—including Duke Albrecht of Bavaria, then head of the House of Wittelsbach, and Baron Teuchert of the WAF—offering to sell back the diamond. We do not know the price he quoted, but it has been reported as either DM 1.5 million (~$375,000; “Der Blaue Wittelsbacher . . . ,” 1964) or DM 2 million (“Wer hat denn . . . ,” 1962). Whatever the price, Duke Albrecht refused the offer. Baron Teuchert, for his part, called the Wittelsbach Blue an “unproductive asset” not worth buying back (“Der Blaue Wittelsbacher . . . ,” 1964). He also then revealed where the diamond had been during 1931–1951: back in the WAF’s safe in Munich.

Lucerne, Hamburg, and Düsseldorf. Spurned by the Wittelsbachs, Komkommer sought other buyers. On August 14, 1963, during the International Lucerne Music Festival, he exhibited the Wittelsbach Blue at the Gübelin jewelry store, then overseen by famed gemologist Dr. Edward Gübelin. At the time, the Wittelsbach Blue was valued at about 2 million Swiss francs or around $500,000. Publicity was high: Three local newspapers reported on the event (“Ein berühmter Steinerner . . . ,” 1963; “De Grote Blauwe Diamant . . . ,” 1963; “Der ‘Wittelsbacher’ in Luzern,” 1963). Komkommer had hoped to sell the Wittelsbach Blue to one of the festival guests, but by mid-September had seen no success. However, while the stone was in Lucerne, Dr. Gübelin was able to conduct the first gemological examination of the diamond (see below).

In 1964, Hamburg jeweler Renatus Wilm recognized a chance to sell the Wittelsbach Blue to one of his countrymen (Kuballa, 1964, Biehn, 1965). Some “residual patriotism” led him to fear that this most important historic German gem could “drift to America,” as had happened earlier with the Hope diamond. Komkommer and Wilm entered into a contract: Wilm would earn a $50,000 finder’s fee if he sold the Wittelsbach Blue by January 31, 1965, but Komkommer would get the same amount of money from Wilm should Wilm be unsuccessful (Kuballa, 1964; Biehn, 1965). To this end, Wilm exhibited the Wittelsbach Blue in his jewelry shops in Düsseldorf and Hamburg during October.

Wilm’s efforts saw more success than those of Komkommer. At the end of 1964, he sold the Wittelsbach Blue and earned his $50,000. The name of the buyer was not disclosed.

SOLD TO A PRIVATE PARTY
Antibes and Zurich. Here, the previously published history of the Wittelsbach Blue (e.g., Gaal, 1977; Bruton, 1981; Balfour, 1997; Morel, 2001) comes to an end, with the fate of the diamond after 1964 largely a mystery. In March 2006, we began our investigations to determine the name of the private purchaser in the digital archives of German newspapers and magazines. Our attention soon focused on Helmut Horten (1909–1987) of Düsseldorf, at one time the owner of one of Germany’s largest department-store chains. One of the authors (RD) had first speculated on Horten’s identity as the buyer of the Wittelsbach Blue in an article published in a small journal on Palatine topography in 1982 (Dröschel, 1982).
The first reports connecting Horten with an unnamed 35 ct blue diamond were published in 1966. Horten had married in secret that year; his bride was a young secretary from Vienna named Heidi Jelinek, 32 years his junior (Bissinger and Lebeck, 1971a). Reports of the wedding appeared in the Rheinische Post (Diebäcker, 1966) and other publications (Adabei, 1966a,b) on August 3, 1966. Among other salacious details about the party at which Horten celebrated his second marriage, the article reported that Horten had presented his wife with a 35 ct blue diamond as a wedding gift. For Rheinische Post journalist Diebäcker, the article had severe consequences (J. Diebäcker, pers. comm., 2006). Horten withdrew all advertisements for his stores (in the order of DM1.5 million), demanding that Diebäcker be fired. Fortunately for Diebäcker, the publishers of Rheinische Post refused. (The two other articles—Adabei, 1966a,b—were published pseudonymously.)

A two-part 1971 article in Stern magazine (Bissinger and Lebeck, 1971a,b) provided more details of the Hortens' post-wedding party at Cap d’Antibes, France, which according to Diebäcker featured performances by the famed BlueBell Girls dancing troupe from Las Vegas, a ballet troupe from Tokyo, a dancing group from Oslo, and musicians from Greece (Diebäcker, 1966). Reportedly, Helmut Horten presented the diamond to Heidi by simply pulling it from his trouser pocket in the midst of this gala. The 240 guests in attendance likely did not realize that the blue gem at the center of this scene was irreplaceable, once part of the Treasury of the Munich Residence.

Other popular journalists were also seeking the diamond. Academy Award-winning Swiss film producer Ernst Albrecht Heiniger (1909–1993; L. Piccolin, pers. comm., 2004) and his wife Jeanne were also avid gem photographers. The Great Book of Jewels (Heiniger and Heiniger, 1974) includes the story of their years-long search for the Wittelsbach Blue in the early 1970s.

The Heinigers had traveled thousands of miles to locate the gem without success. The diamond was nowhere to be found, and the name of the 1964 purchaser “was guarded with utmost secrecy” (Heiniger and Heiniger, 1974). Finally, after three years, they learned by chance that the Wittelsbach Blue was stored in a vault not far from their photo studio in the Zurich Bahnhofstrasse (G. Kling, pers. comm., 2007). Photographing the Wittelsbach Blue required lengthy negotiations and a costly insurance policy (Heiniger and Heiniger, 1974). In addition, they were required to sign documents stating that they would never reveal any information about the owner or the stone’s location. The photo of the Wittelsbach Blue taken by the Heinigers is shown in figure 16.

A variety of other reports of the blue diamond appeared in German publications over the ensuing decades (e.g., Neuhauser, 1971; Zipser, 1991). Although no other blue diamond of this particular weight has ever been reported, none of these articles identified the stone as the missing Wittelsbach Blue. However, a brief 1979 article discussing the 70th birthday celebration of Helmut Horten (“70 Jahre H. Horten,”) finally confirmed the Hortens’ ownership of the historic diamond. The article mentioned that, in 1966, Horten had presented Heidi with one of the greatest diamonds that ever adorned a woman: the famous “Wittelsbach Blue” (figure 17).

In March 2006, 27 years after this article appeared, one of the authors (JE) asked the Welt am Sonntag editor responsible for the 1979 report about its factual basis. He replied that the paper would not have published information on a stone like the Wittelsbach Blue from the private life of a person like Helmut Horten without his specific agreement.

Figure 16. This photo of the Wittelsbach Blue was taken by Ernst Albrecht Heiniger and his wife Jeanne after years spent tracking down the diamond. The mounting was created in the 1960s by Harry Winston and was designed to reduce the visibility of the extremely large culet, and thus improve the overall appearance of the historic stone. Reprinted by permission of J. Heiniger.
The authors also contacted Jeanne Heiniger in September 2007 asking her to review the sections of this article dealing with the Hortens’ ownership and the Heinigers’ photograph of it. She answered by e-mail that “no addendum and no correction” were required (J. Heiniger, pers. comm., 2007). This is the second independent “source” for the Hortens’ ownership.

**St. Moritz and London.** In November 2006, the authors received a report that the elusive Wittelsbach Blue had recently been displayed in a private exhibition at the Bulgari store in St. Moritz, Switzerland. This exhibition and a later one in Vienna were probably the first indications that Heidi Horten planned to sell the Wittelsbach Blue.

In November 2008, Christie’s announced that the Wittelsbach Blue would be sold at auction in London on December 10, 2008 (Christie’s, 2008a,b; Kratzer and Evers, 2008). This auction was conducted at Christie’s headquarters—the same building where the Wittelsbach Blue was unsuccessfully offered nearly 80 years earlier. The winning bid was placed by London jeweler Lawrence Graff, who paid a hammer price of £16,393,250, or just over $24.3 million and $683,000 per carat (Christie’s, 2008c). This is a record price for any diamond or piece of jewelry at auction. The WAF did not participate in the auction, but the authors have been told that it intends to negotiate with Graff to purchase the diamond (R. Borchard, pers. comm., 2008). However, no decisions in that respect had been made when this article went to press.

**GEMOLOGICAL INVESTIGATIONS**

During the short period in 1963 and 1964 between the rediscovery of the Wittelsbach Blue by Jozef Komkommer and its sale to Horten, a few diamond specialists had the opportunity to examine it. In addition to Dr. Gübelin, these included Finnish jeweler and diamond historian Herbert Tillander, who measured the cut, weight, and dimensions (Tillander, 1965). Hamburg jeweler Renatus Wilm, who had sold the Wittelsbach Blue to Horten in 1964, supported Tillander’s findings. Most recently, the Christie’s catalogue in which the diamond appears published a copy of the September 24, 2008, GIA Colored Diamond Grading Report on this stone (Christie’s, 2008b, p. 126).

The rough diamond from which the Wittelsbach Blue was cut was probably a flat slab about 9 mm thick. This can be deduced from the broad proportions of the cut and the extremely large culet. Where the Wittelsbach Blue was cut and polished cannot be stated with certainty, but Paris (Morel, 1988), Lisbon and Venice (Tillander, 1995), and Bruges, Antwerp, and London (Bruton, 1981) have all been proposed in the literature. Jozef Komkommer reported the polish of the Wittelsbach Blue as being “uncommonly smooth, smoother than the work of the very best polishers of today” (de Smet, 1963, p. 8). Tillander also reported that the “appearance of this stone is particularly striking because of its unusually fine
polish and the absolute flatness of the facet surfaces” (Tillander, 1965).

In the GIA report, the cut of the Wittelsbach Blue is described as a “cushion modified brilliant.” Like a modern brilliant cut, it shows excellent eightfold symmetry (figure 18). According to Tillander, it is basically a star cut (figure 18, top left) with its apex replaced by a table facet (figure 18, top center), so that the radially bisected girdle facets remained unchanged. Then, a brilliant cut was superimposed to complete the cut (figure 18, top right; Tillander, 1965). In addition to the main and girdle facets, the Wittelsbach Blue has double pavilion facets—perhaps unique for a historic diamond cut—with eight precisely developed facets around the culet (figure 18, bottom). According to Tillander (1995), it is the earliest known brilliant.

Photos of the stone and the GIA report indicate that over the years the knife-edge girdle was damaged in several places (figure 19). One can only hope that a future owner of the Wittelsbach Blue does not repolish it in order to remove this minor damage, as it would result in a loss of the stone’s subtle original substance, identity, and historic form. However, a statement from Graff Diamonds shortly after the December 10 auction indicated that they intended to do just that, with the aim of making the diamond “flawless and a deeper color” (Reyburn, 2008).

As noted, the first gemological characterization of the Wittelsbach Blue as a type IIb diamond was performed in 1963 by Dr. Gübelin (“De Grote Blauwe Diamant . . . ,” 1963). Balfour (1987) later reported that Gübelin observed red phosphorescence after exposure of the stone to short-wave ultraviolet radiation. Most blue diamonds display a chalky blue to green phosphorescence; only very rarely, as with the Hope diamond, do they exhibit red or orange-red (King et al., 2003; King et al., 1998). Gübelin also reported “strong semiconductivity” (“Ein berühmter Steinenmer . . . ,” 1963; “De Grote Blauwe Diamant . . . ,” 1963). In a letter that accompanied the diamond grading report (Christie’s, 2008b, p. 124), GIA confirmed that the 35.56 ct diamond was a type IIb with a moderate concentration of boron and that it had “bright and persistent red phosphorescence,” similar to that present in the Hope diamond. It is also interesting to note that, like the Hope, the Wittelsbach Blue was color graded Fancy Deep grayish blue. The GIA report gave the clarity grade as VS2, and the color as even. Polish and symmetry were “good,” and—as noted by others—the girdle was extremely thin. Measurements revealed a 64% table, extremely large culet, and 38.8% total depth.
Helmut Horten died in Switzerland in November 1987. Heidi Horten, who has no children, is now the richest woman in Austria, with assets of €3.7 billion (“Special report: The world’s billionaires,” 2008). Although she can use and increase the possessions of her former husband, she is not allowed to dispose of them in her will (Stern, 1971a; “Der Lotto-Gewinn des Jahres . . . .”, 1994). This may be the reason why she decided to sell the Wittelsbach Blue, which was her personal property, at the Christie’s auction.

During the 30 years between 1931 and 1964, the list of poor decisions dealing with the fate of the Wittelsbach Blue is remarkable. However, one man made the right decision for the Bavarian blue diamond, at the right moment: Jozef Komkommer, who refused to recut the Wittelsbach Blue into a more modern shape and thereby preserved its subtle essence, historical identity, and importance. The photo of Jozef Komkommer examining the Wittelsbach Blue in figure 20 serves as a small monument to a conscientious man.

Figure 20. Jozef Komkommer (1911–1980), whose recognition of the Wittelsbach Blue in 1961 (with the help of his son) saved it from recutting, is shown here examining the diamond with a loupe.
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Fancy Dark Brown-Yellow Zoned Type IIa/IIb DIAMOND

Type IIb diamonds are very rare; type IIb diamonds in brown hues are even rarer. While zoned type IIa/IIb diamonds have been reported (see, e.g., Lab Notes: Fall 1993, p. 199; Summer 2005, pp. 167–168), they are also far from common. Therefore, the combination of properties exhibited by the diamond in figure 1, which was sent to the Carlsbad laboratory for a Colored Diamond Grading Report, is remarkable.

The 3+ ct pear shape was color graded Fancy Dark brown-yellow. No color zoning was visible with magnification, even when the stone was immersed in methylene iodide. Internal graining was colorless and weak. The diamond was inert to both long- and short-wave ultraviolet (UV) radiation from a standard gemological UV lamp, and exhibited no phosphorescence. When examined between crossed polarizers, it exhibited a dense “tatami” pattern, with strong localized strain surrounding a crystal (the only solid inclusion observed). The stone also exhibited feathers and indented naturals.

The mid-infrared (IR) spectra (figure 2) clearly showed the presence of two diamond types: IIa and IIb. When the pear shape was oriented such that the shoulder end was sampled, the resulting spectrum was typical of type IIa diamond with no absorption from boron or nitrogen impurities. However, when the point end was sampled, the spectrum exhibited bands at 2930 and 2801 cm$^{-1}$, which are characteristic of boron in type IIb diamond. This IIb spectrum was similar to that of a Fancy Dark greenish yellow-brown type IIb diamond examined at the New York laboratory in 2005 (again, see figure 2), though the boron peaks in the 2008 sample were less intense.

In the DiamondView, the type IIa portions showed blue fluorescence and no phosphorescence, while the IIb regions exhibited much weaker blue fluorescence (figure 3) and short-lived weak blue/white phosphorescence. Boundaries between zones were irregular—in some places linear and in others jagged (figure 4, left). The IIa portions of the stone exhibited “mosaic” dislocations (figure 4, right), which are typical in natural IIa and IIb diamonds (e.g., P. M. Martineau et al., “Identification of synthetic diamond grown using chemical vapor deposition [CVD],” Spring 2004 Gems & Gemology, pp. 2–25). Additionally, a “flame” or “wave” structure observed in the fluorescence of the IIb region was replicated in the phosphorescence for that region.

Since boron acts as an electron acceptor, type IIb diamond is classified as a semiconductor. When the electrical conductivity of the zoned diamond was

![Figure 1. This 3+ ct Fancy Dark brown-yellow diamond was found to be distinctly type zoned (IIa and IIb).](image)
tested, the shoulder end (IIa) did not conduct, while the point end (IIb) was weakly conductive. This observation was not, in itself, conclusive, since type IIb diamonds are not homogeneously conductive, but it did correlate with the mid-IR and DiamondView results.

Photoluminescence (PL) spectra collected with a 488 nm laser revealed the presence of the 3H peak (503.5 nm), which has been reported before in type IIb diamonds (e.g., Fall 2004 Lab Notes, pp. 241–242). This peak occurred in spectra from both regions of the stone, but it was stronger in the IIb region. Spectra collected using three other laser wavelengths indicated that the stone was of natural color. Raman spectroscopy (514.5 nm excitation) was used to try to identify the crystal inclusion, but no match could be found. This was unfortunate, since knowledge of the mineral might have provided valuable information pertaining to the origin of this exceptional diamond.

Karen M. Chadwick

Figure 2. The mid-IR absorption spectra of the brown-yellow diamond in figure 1 indicate the presence of type IIa and IIb zones. Also shown for comparison is the spectrum of a greenish yellow-brown type IIb stone.

Figure 3. This DiamondView image of the pavilion of the pear shape clearly shows the different type zones: moderate blue fluorescence for the type IIa region, and weak blue for the IIb zone.

Figure 4. The boundaries between the type zones are linear in some places and jagged in others (left); the type IIa regions in the diamond also show the subtle mottled appearance of “mosaic” dislocations (right).

HPHT-Treated CVD SYNTHETIC DIAMOND Submitted for Dossier Grading

Gems & Gemology has reported on synthetic diamonds grown by the chemical vapor deposition (CVD) method for several years (e.g., W. Wang et al., “Gem-quality synthetic diamonds grown by a chemical vapor deposition [CVD] method,” Winter 2003, pp. 268–283; P. M. Martineau et al., “Identification of synthetic diamond grown using chemical vapor deposition [CVD],” Spring 2004, pp. 2–25; and W. Wang et al., “Latest-generation CVD-grown synthetic diamonds from Apollo Diamond Inc.,” Winter 2007, pp. 294–312). CVD synthetic diamonds have been submitted to the GIA lab (e.g., Lab Notes: Spring 2008, pp. 67–69, and Summer 2008, pp. 158–159), but those samples were as-grown, even though high-pressure, high-temperature (HPHT) treatment of CVD synthetics is well known.

The 0.21 ct round brilliant in figure 5 was recently submitted for Diamond Dossier grading. Standard testing identified it as a CVD synthetic. It was graded near colorless (GIA does not use
letter grades on synthetic diamond reports) and VVS, with the clarity grade based on pinpoint inclusions. Between crossed polarizers, it exhibited weak birefringence in shades of gray (figure 6). In comparison, the as-grown sample described in the Spring 2008 Lab Note showed strong birefringence with high-order interference colors, consistent with the samples examined by Wang et al. (2007).

The sample was inert to long-wave UV radiation but fluoresced weak yellow to short-wave UV. In the DiamondView, it fluoresced blue-green, with yellow-green striations (figure 7), and exhibited weak blue phosphorescence. The fluorescence color was similar to the green luminescence of HPHT-treated nitrogen-doped CVD synthetic diamonds reported by Martineau et al. (2004), and it contrasted distinctly with the orangy pink to orangy red hues of the as-grown CVD synthetics described in the previous Lab Notes. The striations are growth phenomena, and are typical of CVD synthetic diamonds (see, e.g., Martineau et al., 2004).

The mid-IR spectra established that the sample was type IIa. Neither of the hydrogen-related peaks at 3123 and 3107 cm\(^{-1}\) was detected. Martineau et al. (2004) stated that the former would be removed by HPHT treatment, while the latter could appear after HPHT treatment. A weak peak was observed at 1332 cm\(^{-1}\), consistent with the near-colorless samples described by Wang et al. (2007).

Photoluminescence spectra collected at liquid-nitrogen temperature (~77 K) with 488 and 514.5 nm laser excitation (figure 8) exhibited a very large peak doublet at 736.6/736.9 nm, due to the silicon-vacancy (Si-V) defect. Correspondingly, a very small doublet was recorded in the UV-visible spectrum at ~737 nm; the spectrum was otherwise featureless but with absorption rising toward the
blue region, as is typical of a near-colorless type IIa diamond.

The Si-V defect was originally considered indicative of CVD synthetic diamonds; however, recent work has shown that the defect also exists in some natural colorless and near-colorless diamonds (C. M. Breeding and W. Wang, “Occurrence of the Si-V defect center in natural colorless gem diamonds,” Diamond and Related Materials, Vol. 17, 2008, pp. 1335–1344). Large peaks were observed at 575.0 and 637.0 nm (the zero-phonon lines [ZPLs] of the nitrogen-vacancy centers [N-V]^0 and [N-V]^−, respectively), consistent with CVD synthetic diamond (see, e.g., Wang et al., 2003). In contrast to the as-grown CVD synthetic diamonds documented in the two earlier Lab Notes, the PL spectra for this diamond did not exhibit a 596.5/597.0 nm doublet. Again, this is consistent with the HPHT-treated CVD synthetic diamonds discussed by Martineau et al. (2004). The 488 nm spectrum did display a large peak at 503.1 nm—the H3 ZPL—associated with nitrogen. This is also consistent with the results of Martineau et al. (2004), and contrasts with the relative lack of H3 peaks in the near-colorless as-grown CVD synthetic diamond samples investigated by Wang et al. (2007). It appears that the H3 defect is introduced during HPHT annealing.

We have seen very few HPHT-treated CVD-grown synthetic diamonds in the laboratory, but the criteria discussed above allowed us to successfully identify this sample.

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QUARTZ with Secondary Covellite Dendrites

The discovery of some unusual dendritic inclusions in two transparent faceted quartz gems sent to the laboratory for examination helped us make a connection between two separate notes previously published in Gems & Gemology. The first of these (Spring 2005 Lab Notes, pp. 47–48) described thin hexagonal platelets of the copper sulfide covellite in colorless and smoky quartz, said to be from Minas Gerais, Brazil. The material was being marketed as “pink fire” quartz due to the intense pink reflectivity of the numerous tiny covellite inclusions. These appeared to be primary to the formation of the host, since they were not associated in any way with surface-reaching cracks. As is characteristic for thin crystals of covellite, these inclusions also appeared grayish green in transmitted light.

The second report (Fall 2005 Gem News International, pp. 271–272) dealt with inclusions of a blue-to-green mineral in quartz, which occurred as jellyfish-like radiating clusters. This mineral was identified as gilalite, a hydrated copper silicate. The quartz was stated to have come from the Brazilian state of Paraíba, and is often sold as “Paraíba” quartz.

Our recent examination of the two faceted quartz gems, which are reportedly from the same locality that produced the gilalite inclusions, proved interesting, as both contained obvious black-appearing dendrites that were green in transmitted light (figure 9, left), with a bright pink reflectance color (figure 9, right). This was the same reaction previously reported for hexagonal platy covellite inclusions, which suggested that these dendrites were also covellite. Moreover, one of the gems contained inclusions of gilalite (identified by their appearance) as well, indicating the presence of copper in the system. The dendrites were situated along surface-reaching fracture planes, proof that they formed after the quartz had crystallized.

Since the broad edges of the inclusions had been polished through and were exposed at the surface, these dendrites made ideal targets for Raman microanalysis, which confirmed that they were covellite. Covellite is known to be a secondary copper mineral in copper deposits, so the discovery of these dendrites as fillings in quartz was not surprising even though such secondary inclusions in quartz have not been reported before.

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Induced Copper Contamination of Tourmaline

In recent years, Gem Identification has received numerous requests to analyze tourmalines for the presence of copper,
since (justifiably or not) copper-bearing tourmalines generally command significantly higher prices in the marketplace than their non-cuprian counterparts. Such analyses are typically conducted on surface and near-surface areas of the faceted gems.

Since we are dealing with the detection of trace elements (in parts per million), we invariably face the problem of accidental or deliberate surface contamination. As the experiments described below indicate, this can be accomplished either by polishing a tourmaline on a copper lap, or by soaking the stone (faceted or in its rough state) in a concentrated solution of a copper salt such as copper sulfate. Accidental contamination from a copper lap is a very real possibility, since copper laps are often used to polish gems such as tourmalines.

To demonstrate the potential for this contamination, we started with two small (1–2 ct) faceted "watermelon" tourmalines and analyzed them by energy-dispersive X-ray fluorescence (EDXRF). Neither stone showed any copper peaks. We then rubbed the tourmalines 10 times in a circular motion against a flat copper lap, using moderate thumb pressure. When we repeated the EDXRF analyses, a clear signal for copper appeared.

At this point, we examined the tourmalines with a gemological microscope. We did not see any surface-reaching pits or cracks in the first tourmaline, only fine polishing lines. Nor did we see any evidence of copper on its surface.

The other stone had no eye-visible surface defects, but it did have a surface-reaching crack across the table that picked up minute amounts of microscopic copper when it was rubbed against the lap. In standard darkfield illumination, this contaminant was seen as thin black smudges that resembled the typical surface dirt often seen in such cracks (figure 10, left), but fiber-optic illumination revealed the metallic nature and copper color of the contaminant (figure 10, right). In light of these results, it seems clear that if copper laps are used to polish tourmalines, there is a possibility of accidental (or—though difficult to prove—deliberate) contamination.

This brings us to intentional contamination of tourmalines using concentrated copper salt solutions. The salt we chose for the experiment was copper sulfate, since chalcanthite (the mineral name for naturally occurring copper sulfate) was readily available. We have no reason to believe that any other copper salt in concentrated solution would not have worked just as well.

For this experiment, we selected another small (3 ct) faceted tourmaline with a few surface-reaching features—growth tubes and cracks—that did not detract significantly from its outward appearance. We also treated an ~7.5 ct piece of rough gem-quality tourmaline that had a nice greenish blue color that was not copper related.

Again, we first analyzed both samples using EDXRF; no copper was detected in either of them. Next, we warmed both samples in the light well of a gemological microscope and dropped them into the concentrated copper sulfate solution, which had been cooled in a refrigerator. This created a mild vacuum, which allowed the copper sulfate solution to enter any surface openings in the test subjects. After the solution had reached room temperature, we removed the tourmalines and hand-dried them with paper towels. Again, we repeated the EDXRF analyses.

Both tourmalines had absorbed enough copper solution to produce a visible copper signal on their EDXRF spectra. The copper peaks in the rough sample were not nearly as strong as the signal from the faceted stone, as expected. Although copper sulfate solution has an obvious blue color, no color from the solution was observed in either sample.

In analyzing tourmaline for copper, GIA uses a cleaning protocol that eliminates the potential for errors resulting from such contamination. These experiments underscore the importance of cleaning test samples thoroughly before analysis, and being aware of the very real possibility of surface contamination if unexpected elements are detected during analysis.

John I. Koivula, Kevin G. Nagle, and Philip A. Owens

**Figure 10.** The crack across the table of this tourmaline readily picked up minute traces of copper when the stone was rubbed against a flat copper lap. In standard darkfield illumination (left), the copper looks like typical smudges of black dirt. However, with surface-incident fiber-optic illumination (right), the metallic nature and copper color of the contaminant are clearly seen. Field of view 0.9 mm.

**PHOTO CREDITS**
Robison McMurtry—1 and 5; Karen M. Chadwick—3, 4, 6, and 7; John I. Koivula—9 and 10.
COLORED STONES AND ORGANIC MATERIALS

Visit to andesine mines in Tibet and Inner Mongolia. Gem-quality plagioclase feldspar (labradorite) has been recovered for years from the U.S. state of Oregon (e.g., A. M. Hofmeister and G. R. Rossman, “Exsolution of metallic copper from Lake County labradorite,” Geology, Vol. 13, 1985, pp. 644–647; C. L. Johnston et al., “Sunstone labradorite from the Ponderosa mine, Oregon,” Winter 1991 Gems & Gemology, pp. 220–233). In 2002, red andesine-labradorite appeared in the gem market that was reportedly sourced from an unspecified locality in the Democratic Republic of the Congo (Spring 2002 GNI, pp. 94–95), but some believe that this material actually came from China. In late 2005, a red andesine-labradorite called “Tibetan sunstone” was supplied by Do Win Development Co. Ltd. of Tianjin, China, reportedly from Nyima (actually Nyemo) in central Tibet (Winter 2005 GNI, pp. 356–357). Then, at the February 2007 Tucson gem shows, King Star Jewellery Co. (Hong Kong) and M. P. Gem Corp. (Kofu, Japan) introduced a similar red andesine from Tibet called “Lazasine.” A large supply of red andesine allegedly from China was offered for sale as an official gemstone of the 2008 Summer Olympic Games in Beijing. Despite claims to the contrary, there has been widespread suspicion that the red Chinese andesines are diffusion treated (e.g., G. Roskin, “JCK web exclusive: The andesine report,” posted November 12, 2008, www.jckonline.com/article/CA6613857.html).

In October-November 2008, this contributor visited two andesine deposits in the Chinese autonomous regions of Tibet and Inner Mongolia. The investigation was made possible by the cooperation of mine owners Li Tong of Tibet and Wang Gou Ping of Inner Mongolia, as well as trip organizers Wong Ming (King Star Jewellery Co.) and Christina Iu (M. P. Gem Corp.), who are partners in the Tibetan andesine mine. Also participating in the expedition were Masaki Furuya (Japan Germany Gemmological Laboratory, Kofu, Japan), David Chiang (BBJ Bangkok Ltd., Bangkok), and Marco Cheung (Litto Gems Co. Ltd., Hong Kong).

The Tibetan andesine mine we visited is located 70 km south of the region’s second largest city, Xigazê (or Shigatse), in southern Tibet. This area is well south of the Nyima/Nyemo area (Lhasa region), and our guides were not aware of an andesine mine in that part of Tibet. We drove seven hours from the capital city of Lhasa to the mine, which lies at an elevation of more than 4,000 m. The site is divided into north and south areas with a total coverage spanning 3–4 km east-west and 5–7 km north-south. During our visit, fewer than 10 miners were digging pits in the south area, near a piedmont riverbed (located at the base of a mountain). Organized mining began there in January 2006 under the supervision of Li Tong. The work is done by hand, from April to November. According to the miners, red andesine was originally found in this area in the 1970s, and beads of this material first appeared in Lhasa’s largest bazaar (Bakuo Street) in 2003.

The surface layer at the site consists of humic soil that is 0.5–3 m thick. The andesine is mined from an underlying layer consisting of greenish gray or dark gray sand/gravel in the south area (figures 1 and 2), and yellowish red or greenish gray soil in the north area. The andesine-bearing layers are apparently derived from Tertiary volcano-sedimentary
deposits (Qin Zang Gao Yuan [Tibet Highland] area geologic map, Chengdu Institute of Multipurpose Utilization of Mineral Resources, China Geological Survey, Chengdu, Sichuan Province, 2005). In the south mining area, a few tunnels penetrate several meters horizontally into the andesine-bearing horizons. In addition, a shaft was sunk several meters deep in the north area, but mining there was discontinued after the devastating Chengdu earthquake in May 2008. The andesine is concentrated in patches consisting of several to more than a dozen pieces (100–200 g total) mixed with sand/gravel or soil (again, see figure 2). These accumulations appear to have been concentrated across a wide area by water from seasonal snowmelt.

Alluvial transport has rounded the crystals, and most were found as translucent to transparent pebbles that were <1 cm in diameter (figure 3), though the largest pieces reached 4 cm. Most were orangy red; deep red material was less common. Some had areas that were green or colorless, but we did not see any pieces that were completely brown, yellow, or colorless. The annual production from the region is estimated to be 700–800 kg, of which 30–50 kg are gemstones.

Figure 1. The andesine-bearing deposits in Tibet are exploited in a series of tunnels, with the miners using simple hand tools. Photo by A. Abduriyim.

Figure 2. Rounded crystals of Tibetan andesine are found in concentrations mixed with sand/gravel or soil. Photo by A. Abduriyim.

Figure 3. The Tibetan andesine consists mainly of orangy red pebbles that are <1 cm in diameter. Photo by A. Abduriyim.

Figure 4. This horizon has produced andesines in the Guyang area of Inner Mongolia. Photo by Wong Ming.
quality. A visit to mountain peaks in the mining area revealed Jurassic volcanic rocks and detrital deposits; a volcanic origin is also the case for similar feldspar from Oregon.

The andesine from Inner Mongolia is mined from an alluvial deposit of sand/gravel in the Guyang area, north of Baotou city. The mine is situated in the Yinshan tectonic belt of Mesozoic-Cenozoic age (Inner Mongolia Guyang-Xiaoyutai area geologic map, Inner Mongolia Autonomous Region Geological Survey, Hohhot, 1982). Andesine has been recovered from a region measuring 20 km east-west and 5 km north-south. Humic topsoil overlies Tertiary (Pliocene) and Cretaceous sand/gravel; some areas also show layers of tuff or basaltic rock. The andesine is restricted to a light gray layer (locally iron stained) that is 1–3 m thick and lies several meters beneath the surface—down to more than 10 m—within the sand/gravel (figure 4). Organized mining has taken place near Shuiquan and Haibouzi villages, producing up to 100 tonnes annually. The andesine seen by this contributor commonly had high transparency and was somewhat rounded, except for broken pieces that showed well-developed cleavage surfaces. The stones were typically 0.3–5.5 cm in diameter, with 70–80% in the 1–2 cm range (figure 5). Most of the andesine was pale yellow. Colorless or deep yellow stones were uncommon, while other colors have not been reported from this area.

This field investigation confirmed that the Xigazê region of Tibet does indeed produce natural red andesine, while the Guyang area of Inner Mongolia is a source of pale yellow andesine that may be used as the starting material for diffusion treatment.

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Gemological properties of andesine collected in Tibet and Inner Mongolia. While visiting andesine mines in Tibet and Inner Mongolia (see previous GNI entry), one of these contributors (AA) obtained several samples for gemological study that he witnessed being gathered by the miners. Ten pieces (up to 26.0 g; see, e.g., figure 6) from each region were polished with two parallel windows, and all were characterized for this report. It is currently impossible to unequivocally determine in the laboratory whether red andesine in the gem trade has been diffusion treated. This preliminary characterization was done to gather data on red samples that are known to be untreated, as well as pale yellow material that may be used as a starting material for diffusion treatment.

Figure 5. These andesine pebbles (2–5 cm) were mined from the Inner Mongolian village of Shuiquan. Photo by A. Abduriyim.

Figure 6. These are some of the samples of andesine obtained in Tibet (top, up to 5.4 g) and Inner Mongolia (bottom, up to 26.0 g, some partially polished) that were examined for this report. Photos by M. Kobayashi.
All the samples had a waterworn appearance, and some of the Tibetan pebbles also had embayed areas that appeared to have been created by chemical etching (figure 7). The pebbles from Inner Mongolia also were abraded, but they were more angular than the Tibetan samples and exhibited conchoidal fractures.

The Tibetan samples had the following gemological properties: color—brownish red to orange-red to red; pleochroism—weak; RI—1.550–1.561; birefringence—0.009; optic sign—biaxial positive; SG—2.69–2.72; fluorescence—orange to long-wave, and dark red to short-wave, UV radiation; and Chelsea filter reaction—red. The Inner Mongolian samples were pale yellow, but otherwise they exhibited almost identical properties except that they were inert to both long- and short-wave UV radiation, and they showed no reaction to the color filter.

Examination with a gemological microscope revealed that most of the Tibetan samples contained prominent twin lamellae and parallel lath-like hollow channels (figure 8, left), irregular dislocations (figure 8, right), and irregular color patches caused by milky turbidity from fine granular inclusions (figure 9). One of the polished samples displayed aventurescence due to the presence of native-copper platelets. The samples from Inner Mongolia contained parallel flat growth tubes (figure 10, left), as well as abundant linear fissures (figure 10, right) and fine twin planes arranged parallel to a (010) direction. In some cases, the linear fissures caused a weak opalescence, and such stones cut en cabochon would be expected to show a weak cat’s-eye effect. Cleavage planes were also well developed along one (010) direction.

Absorption spectra were measured with a UV-Vis spectrometer in the range 220–860 nm. The Tibetan andesine exhibited absorption from 320 nm toward shorter wavelengths, as well as a prominent broad band near 565 nm due to colloidal copper. In addition, a weak feature near 380 nm was due to Fe\textsuperscript{3+}. Similar absorptions have been documented in red andesine that was reportedly from the Democratic Republic of the Congo and in red labradorite from Oregon (A. M. Hofmeister and G. R. Rossman, “Exsolution of metallic copper from Lake County labradorite,” Geology, Vol. 13, 1985, pp. 644–647; M. S. Krzemnicki, “Red and green labradorite feldspar from Congo,” Journal of Gemmology, Vol. 29, No. 1, 2003, pp. 15–23). Spectroscopy in the near-infrared region (800–2500 nm) revealed an absorption peak near 1260 nm that is caused by Fe\textsuperscript{2+}.

The andesine from Inner Mongolia showed absorptions at 380, 420, and 450 nm. The 380 nm feature was strongest, while the broad band at 420 nm (presumably due to charge transfer between Fe\textsuperscript{2+} and Fe\textsuperscript{3+}) was characteristic. A strong and broad absorption also was observed near 1260 nm.

Energy-dispersive X-ray fluorescence (EDXRF) chemical analysis of andesine from both Tibet and Inner Mongolia revealed very similar compositions, with 55–56 wt.% SiO\textsubscript{2}, 26–27 wt.% Al\textsubscript{2}O\textsubscript{3}, 10 wt.% CaO, and 5.8–6.2 wt.% Na\textsubscript{2}O. Trace elements such as K, Mg, Ti, Fe, and Sr were detected. The Tibetan stones also contained 0.06–0.10 wt.% CuO, but no Cu was detected in the Inner Mongolian samples. The chemical composition showed that all samples were andesine, with some plotting at the border with labradorite (An\textsubscript{47–50}). Previous electron microprobe analyses of samples from Inner Mongolia showed they were labradorite, with a composition of An\textsubscript{50–51} (or An\textsubscript{52–53} if K is excluded; Y. Cao, “Study on the feldspar from Guyang County, Inner Mongolia and their color enhancement,” Master’s thesis, Geological University of China, 2006).

Laser ablation–inductively coupled plasma–mass spectrometry (LA-ICP-MS) analysis showed that samples from...
both localities contained several trace elements: K (2200–3600 ppm), Fe (1000–3200 ppm), Mg (330–620 ppm), Ti (400–510 ppm), Ba (120–160 ppm), Mn (20–40 ppm), Ga (20–30 ppm), Li (10–60 ppm), and Sc (5–15 ppm); B, V, Co, Zn, Rb, Sn, Ce, and Eu were <3 ppm each. No significant elemental difference was observed between Tibetan and Mongolian andesine, other than Cu content: 300–600 ppm in orangy red andesine, and <3 ppm in pale yellow andesine. In addition, Li was slightly dominant in Tibetan andesine.

Additional images from this study can be found in the G&G Data Depository at www.gia.edu/gemsandgemology.

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New find of vivid kunzite from Pala, California. In July 2008, several etched crystals of gem-quality kunzite were found at the historic Elizabeth R mine, located on Chief Mountain in the Pala District of San Diego County (see, e.g., Fall 2001 GNI, pp. 228–231). In mid-2008, Jeff Swanger (Escondido, California) purchased the mine from Roland Reed (El Cajon, California). Mr. Reed has continued to work the Elizabeth R while Mr. Swanger mines the neighboring Ocean View property, where he found a large gem pocket in 2007 (see Spring 2008 GNI, pp. 82–83).

Although this was not the first kunzite discovery at the Elizabeth R, these pieces had a particularly vibrant pinkish purple to pink-purple color that is seldom seen in natural-color kunzite (e.g., figures 11 and 12). Mr. Reed believes they compare favorably to kunzite from the nearby Vandenberg mine, which produced colors that are considered among the finest found anywhere. Approximately 0.5 kg of top-grade material has been recovered, and the best pieces were sent for fashioning to Minas Gem Cutters of Los Angeles. So far 11 stones have been cut, and the two largest ones weighed 57 and 28 ct. The cut material is being sold through Pala International (Fallbrook, California).

The samples shown in figure 12 were examined microscopically by this contributor. The few internal features were typical for kunzite: elongated, tapered etch tubes; and two-phase (liquid and gas) inclusions—either alone, in parallel, or in a “fingerprint” pattern with irregular to rounded shapes. No mineral inclusions were seen. One faceted stone contained very slight cleavage feathers on the pavilion, while the rough piece showed typical shield-shaped etch marks on its surface.

Kunzite is challenging to cut because of its cleavage, twin planes, and sensitivity to vibrations and thermal shock. It also has a reputation as an “evening” gem, since its color fades with prolonged exposure to light or heat. Kunzite can naturally show an attractive color, as in this new find from the Elizabeth R mine, or the purple-pink hue can be produced by irradiating (and annealing) pale or colorless spodumene. Most natural-color kunzite is light pink. This new production from Pala serves as a reminder that gem mining is still active in San Diego County, where kunzite was initially discovered more than 100 years ago.

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Figure 9. This Tibetan andesine exhibits concentrated areas of red and orange-red milky turbidity (left). When viewed with diffused light, some of these areas appear green or near colorless (right). Photomicrographs by A. Abduriyim; magnified 25×.

Figure 10. Dense concentrations of parallel growth tubes were common in the pale yellow andesine from Inner Mongolia (left). In some samples, linear fissures (right) caused weak opalescence. Photomicrographs by A. Abduriyim; magnified 20×.
Natural pearls of the Veneridae family. The best-known natural pearls from bivalves of the Veneridae (classified by Rafinesque, 1815) family are those from Mercenaria mercenaria (Linnaeus, 1758), also known as “quahog” pearls from the mollusk’s common name, northern quahog. These non-nacreous pearls range from “cream” white to brown, and from faint pinkish purple to dark purple, though some are pure white (e.g., figure 13). Like other natural pearls, quahog pearls are seldom perfectly round; in rare cases, circled quahog pearls occur (figure 14). M. mercenaria bivalves are found along the Atlantic coast of North America to the Yucatan Peninsula. The species also has been introduced along California’s Pacific coast.

However, M. mercenaria is not the only mollusk of the Mercenaria genus to produce pearls. White, “cream,” and sometimes brown non-nacreous pearls can be found in another species belonging to the same genus, M. campechiensis (Gmelin, 1791), or southern quahog. This species is found in the southern part of the M. mercenaria distribution area. M. campechiensis is slightly larger than M. mercenaria, and its interior surface lacks purple coloration; thus, it cannot produce purple pearls.

Nor are Mercenaria bivalves the only mollusks of the Veneridae family that can produce beautiful pearls. In the Fall 2001 GNI section (p. 233), one of these contributors (EF) described an almost perfectly round purple pearl found along the coast of France in a mollusk from the Venerupis genus (Lamarck, 1818), V. affinis decussata (Linnaeus, 1758). This mollusk (known as “palourde” in French) is commonly harvested for its meat, which is considered a delicacy.

The interiors of M. mercenaria and V. aff. decussata shells, as well as the pearls associated with them, are similar in appearance (figure 15). The purple color is present at the shell margins, mainly around the muscle scars. Both pearls and shells display the same medium chalky whitish yellow fluorescence to long- and short-wave UV radiation, though it is weaker for the V. aff. decussata.

Raman spectroscopy of the samples in figure 13 (left
image) and figure 14, using 488, 514, and 561 nm laser excitations, showed that the purple color was due to a mixture of unsubstituted polyenic (polyacetylenic) compounds (figure 16). To our knowledge, the origin of the purple color of Veneridae pearls and/or inner shells has not been previously reported. Similar pigments have been observed in freshwater cultured pearls (S. Karampelas et al., “Identification of pigments in freshwater cultured pearls with Raman scattering,” Fall 2006 Gems & Gemology, pp. 99–100).

M. mercenaria can reach 12 cm in diameter, while V. aff. decussata mollusks from the west coast of France do not exceed 7.5 cm. Thus, the latter mollusks produce smaller pearls (rarely up to 6 mm) compared to those from M. mercenaria (rarely up to 12 mm). It should be noted that “gem-quality” natural pearls from V. aff. decussata have been documented only once, whereas there have been numerous reports of gem-quality quahog pearls.

Figure 16. These Raman spectra for a purple quahog (M. mercenaria) pearl were taken at laser excitations of 488, 514, and 561 nm. In the region most “sensitive” to C=C stretching bonds (about 1500 cm⁻¹, see inset), variations in the position, shape, and relative intensities of the peaks are quite apparent. This suggests that the purple color is due to a mixture of unsubstituted polyenic (polyacetylenic) compounds and not to a single pigment. Raman spectra on colored samples from V. aff. decussata showed the same peaks. All the peaks are normalized to the main aragonite peak at 1086 cm⁻¹. The spectra are offset vertically for clarity.

Figure 15. Though V. aff. decussata (inner shell, 4.5 cm) is found on the western coast of France and M. mercenaria (9.5 cm, courtesy of Antoinette Matlins, South Woodstock, Vermont) is found on the North American Atlantic coast, the two exhibit similar coloration. Photo by S. Karampelas.

SYNTHETICS AND SIMULANTS

Synthetic citrine with abundant nail-head spicules. A necklace of transparent yellow faceted beads (figure 17) was sent to Gemlab for identification. Specular reflectance Fourier-transform infrared (FTIR) spectroscopy identified the material as quartz. FTIR spectra recorded in transmission mode were not definitive, but nevertheless were characteristic of synthetic citrine. The spectra contained an unusually intense water absorption centered at ~3200 cm⁻¹ (too strong to be resolved) and only a single sharp peak at 3580 cm⁻¹. In general, natural citrine has a much lower water content and shows more complex FTIR spectra.
Microscopic examination revealed a most unusual inclusion scene: All of the beads were full of hollow growth channels (figure 18, left) and nail-head spicules (wedge-shaped, liquid-filled growth channels terminated by an inclusion on one end; figure 18, right). Nail-head spicules are characteristic inclusions in both synthetic beryl and synthetic quartz, though similar-looking inclusions have been described in some natural stones (G. Choudhary and C. Golecha, “A study of nail-head spicule inclusions in natural gemstones,” Fall 2007 Gems & Gemology, pp. 228–235). Thus, isolated inclusions of this type do not necessarily offer proof of synthetic origin.

Nevertheless, the appearance of the nail-head spicules in these beads was typical for synthetic material, especially since the “heads” of the spicules contained “breadcrumb” inclusions (e.g., figure 19), the most characteristic and common inclusion in synthetic quartz. However, this contributor has never seen such a large number of these inclusions in any type of synthetic material. All of them exhibited the typical wedge shape, and in most of them the liquid contained a gas bubble.

The inclusions, hollow cavities, and nail-head spicules were oriented parallel to the c-axis. In determining the optic axis direction, it was evident that none of the material was twinned—unlike most natural citrine, which is created by heat-treating natural amethyst that commonly contains Brazil-law twinning. The large, hollow cavities likely represent oversized nail-head spicules that were either exposed by the polishing process or reached the surface during the growth process. In some of these very large cavities, the breadcrumb inclusion was found at the narrow end of the channel (figure 20, left); in the others, it was absent (figure 20, right). These features, like the smaller nail-head spicules, can probably be attributed to rapid growth conditions.

The necklace, which had been sold to the client as natural quartz, was therefore identified as synthetic citrine. Despite this deception, the piece was a fantastic source of photomicrographs of nail-head spicules, which normally do not occur in such heavy concentrations.

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Radiocarbon dating of “Neptunian” beads from Asia proves modern origin. During the inaugural Macau Jewellery & Watch Fair in January 2008, this contributor purchased four baroque-shaped drilled beads of an unknown material that were sold as Neptunian beads. The brownish orange samples had white striae, and the brownish orange portions showed an appealing sheen (figure 21). When asked about their origin, the seller reported that the material was from fossilized conch shell found at an altitude of 5,000 m in the Himalaya Mountains. A brochure provided with the samples added that the beads had several medicinal uses.

In the laboratory, we found that the specific gravity of the beads was 2.78. Close examination showed that they had two folded layers, which is common for conch shell (figure 22). Raman spectroscopy identified the material (both the orange and white portions) as aragonite. EDXRF analysis...
showed that Ca was the only major element, with traces of Sr present, as expected for aragonite from conch shell.

However, the material did not have the appearance of a fossil, so the author decided to apply a method rarely used in gemology: age determination by radioisotope. The Swiss Federal Institute of Technology (SFIT) in Zürich performed $^{14}$C isotope measurement to calculate the age (radiocarbon dating can determine ages up to 30,000 years). To make the determination, the laboratory took 200 mg of powder from the drill hole of one of the beads.

A spectrum relating time with atmospheric radiocarbon content and the sample's data is shown in figure 23. Because above-ground nuclear weapons testing in the 1950s substantially raised the concentration of $^{14}$C in the atmosphere (and consequently in all living organisms), the testing produced two possible results (see P. J. Reimer et al., “Discussion: Reporting and calibration of post-bomb $^{14}$C data,” Radiocarbon, Vol. 46, No. 3, 2004, pp. 1299–1304). The $^{12}$C/$^{14}$C ratio of the sample intersected with the atmospheric ratio at 1957 and 1997. Clearly, the beads are far younger than the 35 million years claimed in the brochure.

Henry A. Hänni

**Purplish blue synthetic quartz.** Synthetic quartz has long been available in a wide range of colors, such as yellow, purple-violet, green, pink, colorless, parti-colored, and even blue. Recently, the Gem Testing Laboratory of Jaipur, India, had an opportunity to study a new and unusual “cobalt” blue synthetic quartz.

The $^{12}$C/$^{14}$C ratio of the sample intersected with the atmospheric ratio at 1957 and 1997. Clearly, the beads are far younger than the 35 million years claimed in the brochure.

**Figure 19.** The nail-head spicules in the synthetic quartz beads, such as those shown here, consist of liquid-filled cavities that may contain a gas bubble and be terminated by a “breadcrumb” inclusion. Photomicrographs by T. Hainschwang; field of view is ~1 mm high for both.

**Figure 20.** Some of the surface-reaching hollow cavities in the synthetic citrine beads (e.g., as shown in figure 18) showed a breadcrumb inclusion at their narrow end (left) while others did not (right). Photomicrograph by T. Hainschwang; field of view is 5.4 mm high (left) and 4.1 mm high (right).

**Figure 21.** These baroque-shaped beads, marketed as fossilized Neptunian beads, proved to be recent shell material. The largest bead is ~17 mm long. Photo by H. A. Hänni, © SSEF.

**Figure 22.** The beads consist of alternating layers of nacreous orange and non-nacreous white aragonite. Photo by H. A. Hänni, © SSEF; image width 16 mm.
A purplish blue specimen weighing 73.08 g was submitted for identification. At first glance, the specimen appeared to be a cobalt glass because of its color (figure 24) and apparently frosted surface. However, we observed tiny circular growth features that were very similar to the “cobbled” surface seen in rough slabs of synthetic quartz. When the specimen was viewed from the side, a colorless zone with the appearance of a seed plate was evident, which led us to believe that the material was actually synthetic quartz.

The specimen displayed a clear anisotropic reaction when rotated in the polariscope, confirming that it was not glass. A bull’s-eye optic figure—as expected for quartz—could not be resolved, due to the rough surfaces. With the desk-model spectroscope, however, we observed three strong bands in the green, yellow, and orange regions; this absorption pattern is typical of cobalt. The sample had a strong red reaction to the Chelsea filter, and was inert to long- and short-wave UV radiation.

Examination with magnification confirmed the presence of a seed plate. Also seen was color zoning parallel to the seed plate, as well as nail-head spicules along its length (figure 25). An interesting aspect was the location/orientation of the spicules. In general, nail-head spicules are oriented in one direction pointing away from the seed.
plate, on both sides of the synthetic overgrowth. In this case, however, most of the spicules appeared to be in the seed plate rather than in the overgrowth material. At some viewing angles, the spicules appeared to be oriented in different directions (figure 26).

While the influx of interesting colors of synthetic quartz has given jewelers more options, proper disclosure remains essential.

**TREATMENTS**

**Dyed chalcedony resembling chrysocolla.** Historically, chalcedony has been dyed in a wide variety of colors, often to simulate various other gem materials. Recently, the Dubai Gemstone Laboratory received for identification a 45.78 ct greenish blue cabochon with areas of orangy brown (figure 27). At first glance, the semitranslucent-to-opaque stone resembled chrysocolla. An SG of 2.29 (determined hydrostatically) and a vague RI reading around 1.5 supported this initial impression (R. Webster, Gems, 5th ed., revised by P. G. Read, Butterworth-Heinemann, Oxford, UK, 1994, pp. 326–327). The stone did not exhibit strong reactions to long- or short-wave UV radiation, only a weak, patchy bluish green fluorescence to long-wave UV. The stone's Chelsea filter reaction was pinkish red.

Examination with magnification (up to 68×) did not reveal the minute chrysocolla inclusions present in chrysocolla chalcedony, although such inclusions are typically too small to see with a gemological microscope. However, microscopic examination did reveal that the surface-reaching fractures and cavities had concentrations of a greenish blue dye (figure 28), which could be removed with acetone.

Colorless-to-milky chalcedony can easily be dyed with inorganic cobalt or copper salts to simulate chrysocolla (see A. Shen et al., “Identification of dyed chrysocolla chalcedony,” Fall 2006 Gems & Gemology, p. 140). However, because of the stone's opacity, we did not obtain the characteristic absorption lines of chalcedony dyed with cobalt, and we were unable to perform UV-Vis-NIR spectroscopy to compare our data with those of Shen et al.

Nevertheless, all of these properties pointed to dyed chrysocolla or dyed chalcedony. Raman analysis is unable to differentiate between chrysocolla and chalcedony. However, EDXRF spectroscopy showed a major amount of Si and only traces of Fe, identifying the material as chalcedony. By contrast, chrysocolla is a hydrous copper silicate \([Cu,Al]_2H_2Si_2O_5(OH)_4\cdot nH_2O\), and no Cu was detected in the sample. Therefore, the client was informed that the cabochon consisted of dyed chalcedony.

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**ANNOUNCEMENTS**

Starting with this issue, the conference and exhibit calendars will only appear online at www.gia.edu/gemsandgemology. Please refer to this online resource for regular updates to these calendars.
Beginning with this issue, the Book Reviews and Gemological Abstracts sections will be available only in electronic (PDF) format. These sections will be available free of charge both on the G&G web site (www.gia.edu/gemsandgemology) and as part of G&G Online (gia.metapress.com), and will be paginated separately from the rest of the issue. The year-end Index has also been moved online and is available on the G&G web site in the Indexes section.

These sections are also included in this full-issue PDF. Accordingly, the Table of Contents included in this file lists these additional sections, and thus differs from the Table of Contents in the print version. For these reasons, this PDF is not the official version of this issue—the “journal of record” for this issue is the combined print/online version that was released to subscribers. This full-issue PDF file is created for archival purposes in order to maintain continuity with previous issues.
Photoatlas of Inclusions in Gemstones, Volume 3
By E. J. Gübelin and J. I. Koivula, 672 pp., illus., publ. by Opinio Publishers, Basel, Switzerland [http://publications.microworldofgems.com], 2008. US$299.00

Gemology now has its gemstone inclusion bible: the three-volume Photoatlas of Inclusions in Gemstones. Culminating 35 years of groundbreaking research, Volume 3 is the final “gem” in Dr. Edward J. Gübelin and John I. Koivula’s monumental achievement. The three books together are intended to be the most comprehensive visual reference library of gemstone inclusions available. Like the previous two volumes, this one is abundantly illustrated with high-quality photomicrographs that brim with information, clarity, and beauty.

The book is divided into three major parts. The first includes a prefatory note by gemologist Edward Boehm, Dr. Gübelin’s grandson, and an introduction to using the Photoatlas library. The heart of the book is its second section, “Inclusions in Major Commercial Gems,” which deals with diamond, ruby, sapphire, and emerald. Each of these four chapters summarizes the different geologic processes that form or transport the gem, and the inclusions that characteristically result. As in Volume 2, the host material is clearly indicated in large type at the top of each page (e.g., “Inclusions in Sapphires”).

The section begins with diamond. Mineral inclusions in natural diamonds—peridotitic, eclogitic, and deep-mantle—are discussed and well illustrated. The detailed examination of internal strain and “graining” patterns provides clues to distinguishing natural, treated, and synthetic diamonds. Inclusions in synthetic diamonds grown under high-pressure, high-temperature conditions are discussed and well illustrated [though not those grown by chemical vapor deposition], as are inclusions resulting from the full range of diamond treatments. The chapter concludes with identification of the main diamond substitutes: cubic zirconia and synthetic moissanite.

The ruby, sapphire, and emerald chapters cover inclusions in natural stones from all major localities. The authors often specify the particular mining region from which the inclusions originated and discuss the typical inclusions associated with the various modes of occurrence. Characteristic inclusions are organized according to genetic type. This is very valuable information, particularly for those who are interested in geographic origin. Also discussed are known treatments for each gem material and the inclusions that can identify them, as well as known synthesis methods and the characteristic inclusions they generate. Simulants such as assembled stones, synthetic overgrowths on natural stones, and glass are also discussed.

In the third section, “Inclusions in Rare and Unusual Gems,” 21 different gems—including axinite, benitoite, cordierite, danburite, ekanite, enstatite, fluorite, gypsum, pezzottaite, sapphireine, and taaffeite—are described and strikingly illustrated. This section ends with a useful glossary and index.

Pioneered by the late Dr. Gübelin nearly 70 years ago, the study of internal features in gemstones, along with our current understanding of how and where certain minerals formed in the earth, enables gemologists to infer a great deal of information by simply viewing inclusions with magnification. By referring to the book’s exceptional photomicrographs, the experienced and diligent user of Volume 3 will frequently be able to ascertain the identity of individual inclusions and, in many cases, establish whether the stone is natural or synthetic. If the ruby, emerald, or sapphire is natural, the user may be able to determine the probable geographic origin and detect indications of treatment. The extensive further-reading list in each chapter provides easy access to additional information and photomicrographs. The scope of localities, treatments, and synthetics represented makes this book extraordinarily valuable to the gemologist, jeweler, and gem collector.

Admirers of beautiful art books will certainly appreciate all three of these volumes. The astute salesperson will also find occasions to use these photomicrographs to illustrate the unique beauty of inclusions in gems they are offering to a client.

With a total of 2,033 pages and more than 5,300 exquisite photomicrographs, the three-volume Photoatlas is the most remarkable achievement in the history of gemological literature. This reviewer cannot imagine that any serious gemologist would want to be without it, and
this volume in particular is highly recommended—it is an essential reference for anyone involved in the identification, research, or appraisal of diamonds, rubies, sapphires, and emeralds.

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Bulgari

By Amanda Triossi and Daniela Mascetti, 2nd ed. (revised and updated by Amanda Triossi), 320 pp., illus., publ. by Abbeville Press, New York [www.abbeville.com], 2007. US$75.00

From their humble beginnings as traveling silversmiths in 19th-century Greece, the Bulgari family became a dynasty of internationally acclaimed fine jewelers. Dedication to excellence in design, materials, and fabrication vaulted the company to the iconic status it enjoys today. Patronized by royalty and the power elite, Bulgari is an Old World company that has remained on the cutting edge of fashion.

In 1996, Daniela Mascetti and Amanda Triossi coauthored the first edition of *Bulgari* [reviewed in the Winter 1998 *Gems & Gemology*, p. 303], in which they described the growth of this contemporary jewelry giant. Recently, Ms. Triossi gave the book an extensive revision. In this second edition, the visual design has been refined, the text has been updated, and more than 120 images have been added, including many press photos of the rich and famous wearing Bulgari jewelry.

In some chapters, there is little or no change from the first edition. Others—most notably “History of Bulgari,” “Evolution of the Bulgari Style,” and “Colour and Fabulous Gemstones”—have been updated to reflect changes in the family and their business structure, in new designs, and in the use of gemstones. The chapters on watches and perfumes have also been expanded significantly, and a new chapter on accessories was added to show Bulgari’s further diversification during the 1990s into ties, scarves, leather goods, and eyeglasses.

Also useful would have been a timeline showing when the various jewelry styles and luxury items were introduced to the Bulgari product lines. An alphabetized glossary of the unique Bulgari styles—*tubogas, parentesi, pippoli, celtaura, gourmette*, and *B.zero1*, to name a few—would also have been very welcome. These are minor criticisms, however, when the book as a whole is considered. Overall, Ms. Triossi has done a superb job of revising the first edition to reflect the significant changes in the style and range of luxury products this remarkable company has undergone over the last decade, making this an important addition to the literature of jewelry history.

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Guidebook to the Pegmatites of Western Australia

By Mark Ivan Jacobson, Mark Andrew Calderwood, and Benjamin Alexander Grguric, 356 pp., illus., publ. by Hesperian Press [www.hesperianpress.com], Victoria Park, Western Australia, 2007. US$85.00

While the pegmatite districts of Western Australia and their mineral assemblages are known to many, information about them—particularly their locations—has been sketchy at best. Most were worked in the early 20th century, and some locales hadn’t been visited for decades. Because of the importance of these pegmatites to scientists and collectors, and a renewed interest in the industrial minerals they contain, the authors produced this field guidebook.

The bulk of this work is a listing of the many pegmatite districts and pegmatite-containing geologic areas of Western Australia. A major achievement of the book is the location data, including detailed directions to the pegmatites and mines, as most of the previous literature offered only approximate locations. (The authors make it clear that their provision of directions to a given locality does not guarantee permission to visit. In some cases, they were unable to get permission themselves.)

*Guidebook* is a handsome hardbound volume printed on high-quality paper. It contains 103 black-and-white photos and 76 maps. The introduction covers the history, mineralogy, classification, and mineral assemblages of these pegmatites. An immensely useful listing of the 120 minerals found in Western Australian pegmatites is provided. It includes not only the minerals but also their location[s], with some indicated as type localities. The following 10 chapters cover individual deposits within the major districts or geologic areas. Some of these locations are well known and have yielded significant commercial production (the Wodgina pegmatite field and the Greenbushes pegmatites, for instance). Others have provided mineral specimens for the collectors’ market (including emerald from the Poona pegmatite field and ferrocolumbite from the Giles columbite-beryl prospect in Spargoville). For each locality, be it a small prospect pit or a major operation, the authors provide an introduction and location data [some with GPS coordinates], as well as the history, geology, and mineralogy of the deposit. There are useful indexes for names, localities, and mineral species, plus an extensive reference list.

While the average gemologist might not have an urgent need for this book, pegmatologists and mineral collectors will find it invaluable. The fact that Western Australia’s pegmatites are not miarolitic (i.e., they do not have significant open pockets for freestanding crystals to form) has prevented them from achieving the kind of fame that the Pala District pegmatites of California enjoy. Nevertheless, collectible mineral specimens and some gems have been produced, and renewed interest will
likely lead to the discovery of more in the future.

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OTHER BOOKS RECEIVED

Crazy About Jewelry! The Expert Guide to Buying, Selling and Caring for Your Jewelry. By Susan Eisen, 245 pp., illus., publ. by Full Circle International Publishing [www.crazyaboutjewelry.net], El Paso, TX, 2007. US$16.95. This is the book you want your customers to read. Susan Eisen’s enthusiasm for jewelry shines through her fun and functional work. There are a lot of good practical suggestions here, and though some in the industry might think her advice is simply common sense—such as not packing jewelry in your suitcase when you travel—I’ve heard of jewelry being lost in this manner countless times. Other chapters include “Redesigning Your Jewelry,” “Medical Identification,” “Cleaning It the Right Way,” and “Knowing Your Jeweler.” Ms. Eisen succeeds in relaying her advice in a casual, user-friendly manner, and the true-life stories from her career are both interesting and entertaining. Many of us will relate to her experiences, and the illustrations are colorful and attractive.

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ERRATA

Two book reviews in the Fall 2008 issue, of Amazonite: Mineralogy, Crystal Chemistry, Typomorphic Features and Infrared Reflection Spectrometry in Advanced Mineralogy, Gemology and Archaeometry, both by Mikhail N. Ostrooumov, inadvertently misspelled the author’s last name. In addition, the amazonite book was published by Polytechnics, St. Petersburg, not Nedra, Moscow.

Because of an oversight, Jared Nadler of Birmingham, Alabama, was omitted from the list of 2008 Challenge Winners in the Fall issue.

Gems & Gemology regrets the errors.
COLORED STONES AND ORGANIC MATERIALS


Atomic force microscopy (AFM) and transmission electron microscopy (TEM) were used to investigate the fine structure of calcite prisms in *Pinctada margaritifera* shell. AFM showed that the prisms were made of closely packed circular micro-domains (in the 0.1 µm range) surrounded by a dense cortex. TEM images and diffraction patterns revealed the internal structure of the micro-domains, each of which was enriched in calcium carbonate. Hosted in distinct regions of each prism, some of these domains were fully amorphous while others were fully crystallized as subunits of a larger calcite crystal. At the border separating the two regions, the micro-domains displayed a crystallized core and an amorphous rim, probably representing an arrested crystallization front. Compared to recent data concerning the stepping mode of growth of the calcite prisms and the resulting layered organization at the micron scale, these results offer unexpected insight into the modalities of biocrystallization.


This section is designed to provide as complete a record as practical of the recent literature on gems and gemology. Articles are selected for abstracting solely at the discretion of the section editors and their abstractors, and space limitations may require that we include only those articles that we feel will be of greatest interest to our readership.

Requests for reprints of articles abstracted must be addressed to the author or publisher of the original material. The abstractor of each article is identified by his or her initials at the end of each abstract. Guest abstractors are identified by their full names. Opinions expressed in an abstract belong to the abstractor and in no way reflect the position of Gems & Gemology or GIA.

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Emerald is rare because unusual geologic and geochemical conditions are required to bring together sufficient amounts of Be [to make beryl] with Cr and/or V [the coloring agents]. This article reviews the major emerald deposits of the world and presents chemical composition data for samples from each deposit.

In the classic model, Be-bearing pegmatites interact with Cr-bearing ultramafic or mafic rocks to form emeralds. In Colombian and certain other deposits, however, emeralds result from regional or tectonothermal metamorphic processes without magmatic activity. Various schemes have been proposed to classify emerald deposits, but not all have been useful in clarifying the actual conditions of emerald formation. Recent studies have demonstrated that emeralds crystallized under conditions where a combination of geologic mechanisms (magmatic, hydrothermal, and metamorphic) brought Be in contact with Cr and/or V in the right geologic setting.

In the field, emerald can be recognized by its color, hardness, and form, but it will not concentrate in heavy mineral fractions because of its relatively low SG. Exploration for emerald deposits is typically based on structural geology considerations and geochemical studies of soils and stream sediments in a target area.


The composition and microstructure of opalized saurian [Plesiosaurus] bones from Andamooka, South Australia, were compared to saurian bones that were partially replaced by magnesian calcite from the same geologic formation, north of Coober Pedy, South Australia. The opalized bones were essentially pure SiO2 (88.59–92.69 wt.%) with minor Al2O3 (2.02–4.04 wt.%) and H2O (3.36–4.23 wt.%). No traces of biogenic apatite remained after opalization. During the formation of the opal, the coarser details of the bone microstructure were preserved down to the level of the individual osteons (~100 μm), but the central canals and boundary area were enlarged and filled with chalcedony, which postdates opal formation. The chemical and microstructural features are consistent with opalization occurring as a secondary replacement after partial replacement of the bone by magnesian calcite, and also with the opal forming first as a gel in the small cavities left by the osteons, with individual opal spheres growing as they settled within the gel. Changes in the viscosity of the gel provide a ready explanation for the occurrence of color and pitch banding in opals. The indication that opalization is a secondary process after calcification in the Australian opal fields is consistent with a Tertiary age of formation.


The authors first give a brief history of the beryllium diffusion treatment of corundum. The presence of Be in untreated sapphire was first reported by P. Wathanakul and colleagues at the 2004 International Gemmological Conference, in a trapiche sapphire from Houay Xai in Laos. In mid-2006, F. Claverie and coauthors detected Be in untreated blue sapphires from Ilakaka, Madagascar. Twenty-eight blue sapphires obtained at the Ilakaka mines by one of the present authors were analyzed at the Gübelin Gem Lab, and 12 contained Be in local concentrations of 1–20 ppm. Complementary analyses at other labs [in Bangkok, Berne, and Lucerne] confirmed the findings.

Beryllium in untreated blue sapphires seems to be concentrated in comet-tail inclusions. These cloud-like inclusions also contained some Nb, W, Sn, and Ta. A correlation between these elements and Be could be useful for separating Be-diffused blue sapphire from untreated blue sapphire containing beryllium. The data might also serve as a chemical fingerprint for origin determination.


This article describes modern pearl farming with Pinctada maxima oysters in Southeast Asia, using as examples two pearl farms in Bali and Irian Jaya. The oysters are cultured from fertilized eggs derived from carefully selected donor oysters. Hygienic standards are closely controlled during the entire growth process. These and other measures resulted in a good yield of high-quality pearls. After harvest, the oysters are not re-seeded; the meat is used for seafood and the shells for their nacre. The cultured pearls are processed, quality graded, and marketed in Australia.


A purple waterworn pebble of unknown origin was acquired in Bangkok in 2004, and it was cut into a strongly pleochroic, biaxial gemstone weighing 10.01 ct. It was acquired in Bangkok in 2004, and it was cut into a strongly pleochroic, biaxial gemstone weighing 10.01 ct. It was identified as thortveitite [confirmed by Raman analysis], a scandium yttrium silicate that was previously unknown in gem quality. Electron microprobe analysis revealed significantly higher concentrations of Sc and lower concentrations of Y than were reported in the literature for nongem thortveitite, which is normally opaque to translucent and found only as very small crystals. The unusual chemical composition suggested a possible synthetic origin, though the presence of three-phase inclusions in a planar array...
indicated otherwise. Unlike quenched flux or melt inclusions, these features were composed of a gas bubble, brine, and cubic daughter crystal suspected to be halite; this suggested formation in a hydrothermal environment. The inclusions showed signs of exposure to heat, either in nature or in the laboratory.

**Vaterite in Süßwasser-Zuchtperlen aus China und Japan**


Chinese and Japanese freshwater cultured pearls [beaded and non-beaded] were analyzed by Raman spectroscopy and LA-ICP-MS. The results showed that they consisted only of aragonite and vaterite, with no calcite. For the most part, the vaterite was concentrated near the center of the cultured pearls; less commonly, it occurred in small blemishes on the surface. Continuous growth structures transected both the aragonite and the vaterite areas. Low concentrations of Na and Sr were found in the vaterite, as well as relatively enriched Mg values, which allowed its distinction from aragonite by LA-ICP-MS.

The authors concluded that vaterite is a common phase in freshwater cultured pearls from China and Japan, and that it tends to concentrate near their centers. It was found in cultured pearls of high quality, as well as in lackluster samples.

**DIAMONDS**


This article reviews a century of scientific study of Kimberlites in southern Africa and the diamonds and mantle-derived rocks they contain, which has increased our understanding of geologic processes and the conditions of diamond crystallization in the subcontinental lithosphere. The formation of kimberlite-hosted diamond deposits involves a lengthy and complex series of events, beginning with the growth of the diamonds in the mantle, followed by their removal and transport to the surface by kimberlite magmas. Age dating of mineral inclusions indicates diamond growth occurred several times during the earth's geologic history. Older diamonds—of Archean age—are mainly peridotitic, whereas younger diamonds originated from eclogitic, websteritic, or lherzolitic rocks, and their formation periods correspond in age with major tectono-thermal events in southern Africa.

Only about 1% of the kimberlite bodies discovered in southern Africa have been commercially exploited for diamonds, but among them are some of the world's richest mineral deposits. The bulk of this article is a review of 34 diamond mines in the region, including summaries of their geology and characteristics of their diamonds and mantle-derived rocks. The mines vary greatly in size, diamond grade, and value, as well as in their mantle-derived mineral suites. All the deposits are hosted by the Kalahari Craton, indicating that it provided the right environment for diamond growth and subsequent transport to the surface by kimberlite magmas.


Natural diamonds sometimes contain dark inclusions that are often described as being graphite or a sulfide mineral. In this study, the authors examined 11 near-colorless, slightly rounded octahedral diamond crystals (2–4 mm) with dark eye-visible inclusions [believed to be a sulfide] for possible remnant magnetism. All the samples were from the Orapa mine in Botswana. The dark inclusions were found to be single or multiple metallic black and opaque fracture systems, each of which contained a tiny grain (20–50 μm) of pyrrhotite. When released by crushing of the diamond, these tiny grains appeared “dirty-yellow.” Pyrrhotite has a greater differential expansion than diamond, and its presence caused the localized fracturing of the host crystal. The black material within the fractures had the same chemical composition as the associated inclusion. The shape and orientation of the pyrrhotite inclusions indicate that they formed at the same time as the host diamond. They were found to be capable of carrying strong and stable remnant magnetization.

These results suggest that with the availability of suitable samples, it may be possible to obtain information about the earth's geomagnetic field during key intervals of geologic time. Furthermore, specific details of the remnant magnetism would allow individual diamonds with pyrrhotite inclusions to be uniquely identified, even in cases where the inclusions are quite small (i.e., only a few microns in diameter).


Nanometer-size isolated inclusions were studied in four cloudy octahedral diamonds from the Internationalaya pipe and one from the Jubileynaya mine, both in Yakutia. TEM, AEM, EELS, and HREM analyses of the samples were conducted, as well as line-scan and elemental mapping. All the crystals exhibited an octahedral habit with
opaque central cuboid cores that contained numerous nano-inclusions (30–800 nm). They were composed of multiphase assemblages that included silicates, oxides, carbonates, brines [KCl], and fluid bubbles. Distin-
guishable crystalline phases included a high-Mg silicate, dolomite, Ba-Sr carbonate, phlogopite, ilmenite, ferroperi-
clace, apatite, magnetite, K-Fe sulfide [possibly djer-
fisherite], and kyanite. Carbonates identified by TEM from all the diamonds studied showed a general enrich-
ment in incompatible elements such as Sr and Ba. Some ele-
mental variations in the crystalline phases may be explained by fractional crystallization of the fluid/melt or the mixing of fluids with different compositions. RAH


Little quantitative research exists on the phosphorescence properties of natural blue diamonds. This study used broadband UV radiation and a novel spectrometer system to examine the luminescence of 67 natural blue diamonds, including stones from the Aurora Butterfly and Aurora Heart collections, as well as the 45.52 ct Hope and the 30.62 ct Blue Heart.

The red phosphorescence of the Hope Diamond was once believed to be quite rare. This study showed that virtually all natural blue diamonds have red phosphorescence; however, the color is often masked by a concomitant luminescence in the green-blue region of the spectrum. Sixty-two of the 67 samples exhibited two phosphorescence peaks—at orange-red (~660 nm) and green-blue (~500 nm wavelengths). Significantly, the study demonstrated that because these two bands are nearly always present, the relative intensity of emissions and their decay kinetics [i.e., the ratio of peak intensities plotted against the half-life of the 660 nm peak] yields a unique “fingerprint” for each specimen. Phosphorescence analy-
sis therefore provides a robust method to discriminate among individual blue diamonds using a relatively inex-
pensive, portable desktop spectrometer.

The authors also examined three blue synthetic diamonds and an HPHT-annealed gray-turned-blue natural diamond. All four exhibited the phosphorescence band at 500 nm but not the one at 660 nm, which suggests that phosphorescence spectroscopy might be an effective tool for discerning synthetic and HPHT-treated diamonds from natural blues.

Although the authors acknowledge there is insufficient evidence to completely describe the defect states, impurities, or energy-transfer mechanisms of phosphorescence, their findings suggest that the same donor-acceptor pair recombination mechanism is active in both natural and synthetic blue diamonds.

**GEM LOCALITIES**

Advances in our understanding of the gem corundum deposits of the West Pacific continental margins intraplate basaltic fields. I. Graham [i.graham@unsw.edu.au], L. Sutherland, K. Zaw, V. Nechaev, and A. Khanchuk, Ore Geology Reviews, Vol. 34, No. 1/2, 2008, pp. 200–215.

The continental margins of the western Pacific contain the world’s largest and richest deposits of gem ruby and sapphire. These deposits are genetically related to Late Mesozoic to Late Cenozoic basaltic volcanism, and today they are spread over a distance of more than 12,000 km, extending from Russia (Siberia) to Australia (Tasmania). The gem corundum consists of xenocrysts of mafmatic and/or metamorphic origin trapped in the host basalt. The corundum is mined from placer deposits formed by weath-
ering of the basalt.

Corundum suites from each origin type have distinctive trace-element geochemistry, mineral inclusions, crys-
tallization ages, and formation conditions. Magmatic corundum appears to have crystallized under upper-man-
tle to mid-crustal pressure-temperature conditions (~700–
1200°C), from melts of syenitic to nepheline syenitic com-
position. In contrast, the metamorphic corundum appears to have formed at slightly higher temperatures (800–1300°C) and at depths ranging from the mantle to the lower crust. The conditions of corundum formation at the major deposits within the western Pacific continental margins are discussed.


The author presents an in-depth characterization of three specimens of Afghan beryl from pegmatites at Ghursalak in Konar Province [aquamarine and morganite] and from the Panjshir Valley [emerald]. INAA, XRD, ICP-AES, ICP-
MS, and IR spectroscopic techniques were used to exam-
ine the beryls, and the results are summarized in accompa-
nying tables. IR spectroscopy of the aquamarine and emer-
ald indicated the presence of organic matter, probably bitu-
ninous material within structural channels. The author notes that while Afghanistan has not been a major gem producer for the world market, many important gems have been known from the region since Egyptian, Greek, and Roman times. Production is growing, and examples of fine material have appeared in markets worldwide.

**Age and origin of gem corundum and zircon megacrysts from the Mercaderes–Río Mayo area, south-west Colombia, South America.** F. L. Sutherland [lin.sutherland@austmus.gov.au], J. M. Duroc-Danner, and S. Meffre, Ore Geology Reviews, Vol. 34, No. 1/2, 2008, pp. 155–168.
Alluvial gem corundum has been known for several centuries from the Rio Mayo area of southwestern Colombia. Samples recovered from this area (some near colorless but most multicolored, 99% of them sapphire) exhibit features such as color zoning, polysynthetic twinning, and healed fractures, as well as various mineral inclusions (rutile, apatite, zircon, some plagioclase, and occasional allanite, which appears to be an inclusion unique to this locality). U-Pb dating of the zircon, allanite, and apatite inclusions suggested the corundum crystallized approximately 10 million years ago, placing the formation in the Miocene epoch. Corundum formation appears to be related to geologic events associated with the uplift of the northern portion of the Andes Mountains and accompanying volcanism. The article provides chemical composition data for both the corundum and the important mineral inclusions.


Black opaque spinel, pyroxene, and magnetite occur in gravels associated with corundum in the alluvial deposits of Denchai and Bo Phloi, Thailand. Raman spectra and XRD patterns have been used to unravel some of the minerals surrounding these materials in the gem markets. The black spinel lies in the spinel-hercynite series, the black pyroxene is mostly augite, and the magnetite lies in the magnetite-ulvöspinel series. The details of their chemical composition suggest that these minerals did not originate in the same environment as the corundum or the basaltic host rocks.


Gem corundum is found at a number of localities in Madagascar, primarily in the central and eastern portions of this island nation. Ruby and sapphire formed at different stages and in distinct environments. The authors describe four main geologic settings:

1. primary deposits in magmatic rocks such as syenites, granites, and alkali basalts
2. primary deposits in metamorphic rocks such as granulites
3. primary deposits that resulted from alkaline metasomatism due to fluid circulation occurring along discontinuities in gneisses and granulites
4. secondary deposits derived from the erosion of surrounding rocks

The article provides an excellent review of the major ruby and sapphire deposits of Madagascar, including their geologic setting and age, host rocks, typical mineral assemblages, and inferred conditions of formation.


The Thunder Bay Amethyst Mine Panorama is a major amethyst deposit on the western shore of Lake Superior in southern Ontario. Although most of the quartz is amethystine, loose pieces of yellowish green and green quartz have been found, and greenish gray quartz occurs in situ as part of a color-gradation sequence that includes colorless and smoky quartz along with chalcedony. Analysis of samples of all these colors show corresponding trends in the salinity and temperature of the quartz-forming hydrothermal solutions. The greenish material exhibits greater turbidity and more numerous fluid inclusions than the amethyst. Furthermore, differences in crystal growth rates also appear to have influenced the color of the quartz.

The authors conclude that the greenish gray coloration is not from the secondary heating of preexisting amethyst, but rather is another distinct radiation-induced color variety of quartz. This color resulted from specific chemical constituents in the hydrothermal solutions, the conditions of natural radiation exposure, and the incorporation of molecular water in the quartz, both as nanoscale and micro- to macro-scale fluid inclusions. The greenish gray material appears to have formed during the initial stages of mineralization, and these solutions underwent a decrease in salinity and quartz growth rate during quartz precipitation.

Opal-C, opal-CT, & opal-T from Acari, Peru. F. Cauzia [cauzia@crystal.unipv.it], C. Ghisoli, L. Adamo, and M. Boiocchi, Australian Gemmologist, Vol. 23, No. 2, 2008, pp. 266–271.

Optical features, SG and XRD data, and IR spectroscopic features are described for 25 translucent-to-opaque volcanic opals from the Acari region of Peru. The XRD and IR results correspond with opal-C and opal-CT, with some samples being pure tridymite [i.e., opal-T]. Opals displaying various colors and transparencies were classified according to their luster, and the relationship between luster and the presence of phyllosilicate phases within the opals was assessed. Andean opals with a vitreous but dull porcelain-like luster were opal-C and opal-CT that were free of phyllosilicates.
INSTRUMENTS AND TECHNIQUES

Accelerating refractive rendering of transparent objects.


Ray tracing may be used to create photorealistic computer images of transparent gemstones and other objects. However, this technique requires a great deal of computation and is inherently slow. The authors propose a technique for the interactive rendering of transparent objects using a refractive rendering algorithm. There are two stages in the algorithm: pre-computation and shading. In the pre-computation stage, a ray-tracing process tailored to gemstone rendering is performed. A database is constructed for the storage of information such as the ray directions and the positions of the corresponding image points. In the next stage, these data are retrieved from the database for the shading of the transparent object, taking illumination into consideration. The time required for the pre-computation process is proportional to the number of polygons (p) composing the model, the image size (number of pixels, s), and the number of internal reflections (r). The performance of the shading process is determined by the light obstruction test and the illumination calculation, with a time complexity of \( p^2 \) and \( sr \), respectively. Experimental results show that refractive rendering is significantly faster than ordinary ray-tracing techniques. 

Dennis Zwiggert


The author describes three rules for refractometer observations of biaxial gemstones that, if followed correctly, allow one to determine optic character and sign by simply recognizing observed patterns (without the need to construct graphs of RI measurements). All four possible biaxial patterns are discussed, and diagrams illustrate the behavior of gem materials on the refractometer based on calculated movements of shadow edges for different orientations of the optical elements and the facet being measured. The author notes that although one biaxial pattern is quite common and requires use of a polarizing filter to determine true beta, the other three are rare and may only show up at all because gem cutters sometimes use a large crystal face for the gemstone’s table.

SYNTHETICS AND SIMULANTS


Attractive beads sold as fire agate were proved to be heat-treated chalcedony (agate). The heating causes surface cracking that is somewhat reminiscent of fire agate, but the beads showed none of the surface iridescence or botryoidal structure that are typical of true fire agate.


The author reports on some interesting gem assemblages, eye-catching composites marketed as “Fusion.” The gems have concave pavilion facets, which improves their bril-
liance and overall appearance. Combinations include citrine and amethyst imitating ametrine, and topaz and amethyst resembling tanzanite. The composites vary from three to five pieces, giving an impression of multicolored stones.

Identification of these composites is straightforward. The presence of junction planes, which appear as sharp separations between the layers, is diagnostic. Spherical or flattened gas bubbles, dendritic flow patterns, and iridescence may all be present along the junction planes. In addition, the glues may fluoresce chalky blue to white under short-wave UV. Guy Lalou


Paraíba tourmaline simulants are becoming more common in the marketplace. Materials such as apatite and glass have been joined by cubic zirconia and hydrothermal synthetic beryl; production of the latter two has reached a commercial scale. Short descriptions of these four simulants are presented, and their gemological properties are compared in a table. The authors describe a glass simulant with swirled color zoning, which at first glance displayed pleochroism reminiscent of that seen in tourmaline. All four simulants are easily separated from tourmaline through standard gemological testing.


Reexamination of a synthetic emerald grown in the late 19th century using a flux method showed its composition to be nearly stoichiometric and homogeneous except for significant variations in chromium (1.45–2.59 wt. % Cr2O3). Trace amounts of Ti, Mg, Fe, Zn, Na, K, and F were also noted. Despite the flux growth method, FTIR spectra in the OH-stretching region showed the presence of weak but significant H2O vibrations. The polarized FTIR spectra collected with E∥c consisted of a sharp, intense band at 3463 cm−1, whereas the E∥c spectra consisted of two minor bands at 3643 and 3587 cm−1. These bands were assigned to the ν3 antisymmetric stretching and ν1 symmetric stretching modes of type II water in the structural channels. These water molecules were probably associated with Li impurities in the mineral from the flux used for the synthesis. Using the molar absorption coefficient, the authors derived a water content of ~30 ppm.

RAH TREATMENTS


Chinese andesine has attracted considerable attention due to rumors that an unknown treatment can reproduce the red color seen in some natural material. EDXRF data were collected for samples of pale yellow rough andesine before and after three stages of an unspecified treatment process performed in Thailand. These samples showed increasing CuO levels and deeper diffusion of red color with each stage of treatment. After the third stage, copper levels reached those recorded for andesine from the market.

Two pieces of pale yellow Mongolian rough underwent treatment [also unspecified] in China. Viewed in immersion, they showed less color at their surface and more color internally than the Thai-treated stones. The red color, when magnified, consisted of tiny reddish orange spots. Copper concentrations measured in these samples were also similar to those of andesine available in the market.

A new andesine deposit in a rugged mountainous area of Tibet was also described. Hand mining by locals during the summer has yielded stones with natural red and green hues, most of them <1 ct when cut. More information is needed to distinguish untreated Tibetan andesine from the treated stones.

ERB Pearl treatments: How pearls, natural and cultured, are treated to enhance them, and how to detect the treatments. S. Kennedy, Organic Gems, No. 7, 2008, www.maggiecp.co.uk/subs/OG7%20Jan%202008/Pearl_Treatments.htm.

The article gives an overview of the primary pearl treatments encountered in the market. Most involve color enhancement, such as the use of silver nitrate staining to darken a pearl’s appearance. The author also describes the filling of internal hollows in large pearls to enhance their weight. Both filling and silver nitrate staining are detectable by X-radiography. The author notes that some enhancements are still difficult to detect, such as yellow treated South Sea cultured pearls. Some practical knowledge of certain characteristics indicating treatment is provided. Two examples are the high likelihood of treatment in a matched string of brown Tahitian cultured pearls, and the fact that black is not a natural color for cultured pearls from freshwater mussels.


Following up on his 2006 Journal of Gemmology paper [covering the period 1996–2005], the author updates a list of patents pertaining to the surface treatment of gem materials. Although it is not known if any of the reported technologies have already been implemented, the author warns that that at least some will probably be applied in the future. Two types of surface treatment are identified and discussed: (1) processes not involving heat treatment
[i.e., coatings of wear-resistant material, coatings that cause an optical phenomenon, and coatings that form a diffractive optical element], and [2] processes involving heat treatment [i.e., heat treatment while in contact with a plate containing transition metals, and heat treatment following deposition of a coating].

MISCELLANEOUS


Satellite imagery has become a valuable tool in diamond exploration. This article details the development of a methodology for monitoring diamond mining activity from very high resolution satellite images using a multi-criteria “fuzzy” classification system. Previous investigations were based on medium-resolution sensors [e.g., Landsat TM data]. In this study, two sets of satellite remote-sensing data were acquired four months apart and then processed using a “fuzzy sets” procedure. (“Fuzzy sets” are those in which elements have degrees of membership, rather than either belonging or not belonging to the set.) The authors discuss in detail the data-processing flow–image preprocessing methodology (geometric and radiometric corrections), image processing using multi-resolution segmentation and supervised fuzzy logic classification software; and the classification scheme that would eventually provide the object maps and statistics to assess the status of mining activity in the imaged areas (e.g., increasing, decreasing, or stable mining activity).

Of interest in one African diamond-producing zone was a large mechanized mine and small artisanal “hand-dug” sites that only required light manpower and equipment. The identification parameters for the mines [e.g., average size, aerial geometric shape, distance from rivers, water-clarity status, and excavation materials] were operationalized and became part of multi-temporal data sets.

The multi-criteria method showed an overall accuracy of 91% based on comparison with image datasets from standard manual photo interpretation. The authors conclude that the object-oriented interpretation model developed in the study proved successful in monitoring the level of diamond mining activity.

The advantages of using such image analysis include the wealth of additional information that can be derived from image objects, its robustness, its success in monitoring the level of diamond mining activity in inaccessible areas, and the complementing of maps with qualitative and quantitative information on mining activity for both small artisanal and large mechanized mines.


There has been a proliferation of initiatives aimed at producing truly “clean,” “fair,” or “ethical” gold, diamonds, and other minerals, so that these resources can better contribute to sustainable development in the local communities where they are mined. These initiatives have been instigated and/or sustained by various stakeholders in the diamond business, as well as by governments, donors, NGOs, and other organizations. This paper analyzes nine initiatives designed to improve the lives of artisanal diamond miners and small-scale diamond cutters in various parts of Africa. Some have had better results than others, and the successes and failures are identified to provide lessons for the future.

Two initiatives in the Democratic Republic of the Congo are briefly described, though the report acknowledges that much more is happening there [e.g., diamond-oriented efforts by Congolese civil society organizations working on the social and environmental impacts of mining and forestry]. The report also examines two company-led diamond fair trade initiatives: one by Finesse Diamonds, a U.S.-based De Beers sightholder, and the other by Target Resources, a London-based gold and diamond miner. Many other fair trade initiatives focusing on gems and gold offer lessons to be learned, but are beyond the scope of this paper.

A brief overview is provided of attempts by individual retailers worldwide to sell only diamonds that are ethically sourced. Many choose the safe option of Canadian stones, or even synthetic diamonds, yet these do not contribute to the development of diamond-producing countries in Africa.

Setting goals for standards and establishing guidelines is critical to establishing trust in fair trade principles. Seven organizations that are attempting to do so are analyzed. An overview of charity initiatives is given, many of which are using diamond industry profits to fund African development projects. As such, there is a difference between “diamond charity” and “development diamonds.” The report concludes with eight issues for discussion: the lack of clarity as to what constitutes an “ethical diamond,” the urgent need for monitoring, the difficulty of achieving overall aims, the ongoing lack of information exchange, the inability of many charities to address root problems, the need to collaborate, the fact that too many initiatives are top-down rather than bottom-up, and the failure to recognize existing internal structures that might be most effective.

The author provides a list of socially responsible diamond retailers and designers, as well as references and websites. An overview matrix of the different initiatives and what they aim to achieve is also included.
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