Fancy-color diamonds are among the most highly valued of gemstones due to their beauty and rarity. Interestingly, the rarest of diamond colors correlate with the three most popular choices for favorite color, in general—green, blue, and pink to red. The unique set of conditions in nature that produce the structural imperfections (defects in the lattice of carbon atoms; see Shigley and Breeding, 2013) responsible for the most vibrant hues of green, blue, and pink/red diamonds are so uncommon that many people are not even aware these stones exist. Over the last ten years, diamonds with these natural color components comprised less than 0.4% of all diamonds submitted to GIA’s laboratories worldwide (including both fancy-color and those on the D–Z scale). Pure hues of green, blue, or red are even rarer, accounting for less than 0.07% of all diamonds examined.

Many articles published over the last 20 years in the scientific and gemological literature have looked at specific properties of colored diamonds, quality grading characteristics, or particular treatments. Few researchers, however, have had the opportunity to examine large quantities of similarly colored natural diamonds and report on their distinctive characteristics. Colored diamonds are extremely rare and, consequently, highly valued. This value factor means that laboratory reports are requested for most colored diamonds prior to being sold. As the creator of the diamond color grading system and the largest provider of these grading reports, GIA is uniquely positioned to examine more colored diamonds than anyone else in the world.
Over the last ten years, scientists at GIA have examined more than half a million natural, fancy-color diamonds and systematically documented their gemological and spectroscopic properties. Through a series of articles, we will discuss the major hue groups of natural-color diamonds in detail that has never before been revealed from such a large sampling of stones. The series will include gemological observations, spectroscopy, and statistical compilations from colored diamonds examined over the last decade at GIA. For more information on data collection methods, please see supplemental table 1 at https://www.gia.edu/gems-gemology/spring-2018-natural-color-green-diamonds-beautiful-conundrum.

This first article looks at green diamonds, a beautiful yet enigmatic hue (figure 1). During the last decade, GIA has examined well over 50,000 natural-color diamonds with a green hue component, including more than 9,000 with a pure green hue. While there are a few different causes of green color, the most common cause—accounting for nearly half of the natural green diamonds submitted to GIA—is exposure to radioactive minerals and fluids in the earth’s crust. Over thousands to millions of years, radiation produced by the decay of isotopes of elements such as uranium and thorium [present in minerals or dissolved in geological fluids] physically damages the diamond structure by removing carbon atoms to cre-
Radiation damage
Green luminescence
H-related absorption
Ni-related absorption

TABLE 1. Categories of green diamond.

<table>
<thead>
<tr>
<th>Cause of green color</th>
<th>Defect responsible</th>
<th>Most common colors</th>
<th>Geological formation</th>
<th>Key gemological observations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radiation damage</td>
<td>GR1</td>
<td>Green to blue-green</td>
<td>Exposure to radiation (minerals or fluids)</td>
<td>Green and/or brown spots or stains in fractures; color zones</td>
</tr>
<tr>
<td>Green luminescence</td>
<td>H3</td>
<td>Green to yellow-green</td>
<td>Natural annealing</td>
<td>Localized or dispersed green fluorescence with focused lighting</td>
</tr>
<tr>
<td>H-related absorption</td>
<td>Unknown H-related</td>
<td>Gray-green to brownish green</td>
<td>Crystallization in mantle</td>
<td>Fine particulate clouds, often patterned</td>
</tr>
<tr>
<td>Ni-related absorption</td>
<td>Unknown Ni-related</td>
<td>Yellow-green to green-yellow</td>
<td>Crystallization in mantle</td>
<td>No distinctive features</td>
</tr>
</tbody>
</table>

ate vacant atomic positions (Meyer et al., 1965; Vance and Milledge, 1972; Mendelssohn et al., 1979; Raal and Robinson, 1981; Titkov et al., 1995; Nasdala et al., 2013). These vacancies [sometimes in combination with other defects] cause the diamond to absorb the blue and red parts of incident visible light, allowing primarily green light to be seen when one observes the stone. This natural irradiation process, however, is easily replicated in a laboratory with electrons, neutrons, or gamma rays, and many green diamonds in the trade have been the product of artificial irradiation treatments [where the radiation exposure times are rapid and controllable] since the late 1940s (Ehrmann, 1950).

The similarity of the natural and laboratory irradiation processes has made it very difficult in many cases for gemologists and laboratories to separate natural- and treated-color green diamonds. Indeed, there have been fewer publications about green diamonds than any other color. The purpose of this article is to provide a detailed account of the gemological and spectroscopic characteristics of natural green diamonds colored by several different mechanisms in order to help the trade better understand these gems.

CAUSE OF COLOR

Green color in gem diamonds can span a wide range of color descriptions. For the purposes of this article, we include all natural fancy-color diamonds that display a primary green color component under standard grading conditions. In the GIA fancy-color grading system, this includes stones with a pure green color as well as a dominant green color mixed with blue, brown, gray, or yellow. Of the green diamonds seen at GIA over the last decade, most have been yellow-green or pure green, followed by those with gray or brown modifiers and blue components. These different color groupings correlate, to some extent, to four significant atomic-level defect arrangements in natural diamond: GR1, H3, hydrogen, and nickel defects (figures 2 and 3). Table 1 summarizes the characteristics of these four groups.

GR1 (General Radiation 1). GR1 defects are empty lattice positions in the diamond structure and serve as the most common mechanism for green color in gem diamonds (Shigley and Fritsch, 1990; Collins, 2001; Shigley and Breeding, 2013). Irradiation, both natural and laboratory-induced, provides sufficient energy to displace carbon atoms from their normal positions in the diamond lattice, leaving vacancies (figure 4; see Koptil and Zinchuk, 2000). The displaced carbon atoms then become what scientists call interstitials: atoms that are out of their normal place and “stuck” between other carbon atoms. GR1 vacancies (figure 2A) cause the diamond to absorb light in the red part of the visible spectrum [zero phonon line1, or ZPL, at 741 nm], with associated broad absorption extending from approximately 550 to 750 nm.

Depending on the amount of vacancies and the other impurities and defects present, green diamonds colored by GR1 typically have pure green or blue-green hues (figure 3); irradiation of a light yellow diamond produces a greener color, while irradiation of a colorless diamond causes a bluer color. Most naturally irradiated diamonds have shallow green “skins” or spots on the crystal surface that produce most of the green appearance [Vance and Milledge, 1972; Vance et al., 1973; Kane, 1979b; Fritsch and Shigley, 1991; Roskin, 2000; Eaton-Magaña and Moe, 2016]. Upon faceting, these shallow areas of radiation damage are typically removed and the apparently green rough diamond shows little to

1See Luo and Breeding (2011) for diagram and description of the zero phonon line.
no color as a finished cut stone [figure 5, bottom row]. In some cases, irradiation brings about a deeper green bodycolor that is retained by the diamond after fashioning [figure 5, top row]. Vacancies become mobile at temperatures above 600°C and start to move around and create new defects in the diamond lattice that can change the stone’s color [Collins et al., 2000; Collins, 2001; Eaton-Magaña
and Moe, 2016). During jewelry repair processes, when heating green diamonds colored by this mechanism, jewelers must take great care to avoid destroying the GR1 defects and losing the green color. Heat generated from the friction of polishing can have a similar effect.

The remaining three causes produce less-saturated pure green colors, or green hues with yellow or gray modifiers (figure 3).

**Nitrogen-Related Defects: H3.** Two nitrogen atoms adjacent to a vacancy in the diamond lattice comprise the H3 defect, the second most common mechanism for green color in diamond (figure 2B; see Collins, 2001; Breeding and Shigley, 2009; Shigley and Breeding, 2013). This uncharged defect is usually produced when diamonds containing pairs of nitrogen impurity atoms [A-centers] are exposed to radiation and subsequent heating above 600°C, either at depth in the earth or in a laboratory (Collins, 2001; Zaitsev, 2003). The vacancies introduced by radiation damage are mobilized during the heating, and they combine with the paired nitrogen impurities to create H3. Unlike GR1, H3 produces green color through emission of light [luminescence], rather than absorption (Collins, 2001; Shigley and Breeding, 2013). H3 absorbs light in the blue part of the visible spectrum [ZPL at 503.2 nm, with associated broad absorption from 420 to 500 nm], generating a yellow bodycolor in diamond. When H3 absorbs visible light, it simultaneously emits green fluorescence. High concentrations of nitrogen in the A form se-

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**Figure 3. Different causes of color in green diamonds produce slightly different hue ranges, depending on the other impurities and defects present.**
verely quench the H3 fluorescence (Collins, 2001; Luo and Breeding, 2013), but occasionally a diamond with abundant H3 defects and a low concentration of nitrogen will emit sufficient fluorescence to bring about a combined yellow-green appearance (figures 2B and 3) (Collins, 2001; Luo and Breeding, 2013). H3 defects are very stable to temperatures exceeding 2000°C.

**Hydrogen-Related Defects.** The presence of hydrogen defects in the diamond structure has been known and studied for decades, but very little is known about how H impurities affect color (figure 2C). The best-known hydrogen-related defect in natural diamond is the 3107 cm$^{-1}$ infrared (IR) absorption (Wang and Mayerson, 2002; Goss et al., 2014). This feature is attributed to three nitrogen atoms adjacent to a vacancy.
producing an overall yellow-green bodycolor (figure 3). The origin of the color-causing Ni band is not known, but it is very similar in structure to the absorption sidebands associated with other defects such as N3 and H3. The thermal stability of the broad Ni-related absorption at 685 nm is not known, but studies of synthetic Ni-bearing diamonds suggest it could be stable to temperatures of 2000°C or higher (Zaitsev, 2003).

**OCCURRENCE AND FORMATION**

Green diamonds have been reported from both kimberlite-hosted and secondary alluvial sediments in nearly every country that produces diamonds, but some localities consistently produce more of these valuable stones. Green diamonds colored by H3, hydrogen, or nickel defects occur in many localities. For decades the largest producers of green diamonds colored by GR1 defects have been alluvial deposits in Brazil, Guyana, Venezuela, Zimbabwe, the Central African Republic, and India (Draper, 1951; Themelis, 1987; Coenraads et al., 1994; Chaves et al., 1996; Chaves et al., 2001).

The largest and most famous green diamond is the 41 ct Dresden Green (figure 6), sourced from the Golconda mines in India sometime before its earliest historical record in 1722 (Hough, 1913; Rosch, 1957; Bosshart, 1989, 1994; Kane et al., 1990; Astric et al., 1994; Morel, 1994). Several other famous green diamonds have been sold in recent years, including the 5.51 ct blue-green Ocean Dream (mined in the Central African Republic), the 25 ct Gruosi Green (South Africa and never graded by GIA), and the 5.03 ct Aurora Green (Brazil), which sold at auction in 2016 for more than US $3.3 million per carat (DeMarco, 2016).

Recent mining in South America has yielded a number of green diamonds from Brazil and Guyana (figure 7). Most show extensive surficial radiation damage with variable amounts of internal GR1-related green coloration. Another recent source is Zimbabwe. Rough diamonds from the Marange region of Zimbabwe (figure 7) are well known to have very dark green or brown coatings of radiation-damaged areas and often show internal green coloration as well [Breeding, 2011]. Stones from this source are distinctive in that they also contain high concentrations of hydrogen defects that can enhance the green color from GR1 defects (e.g., a combination of figure 2A and figure 2C contributes to green color). Production from Zimbabwe has been very high in the last decade, but controversy engulfs these goods, as the
United States and other countries still have an embargo on the import of Zimbabwe diamonds due to human rights concerns (Miller, 2011).

The formation mechanism for green diamonds in the earth varies significantly depending on the defects responsible for the color. Stones colored by hydrogen- or nickel-related defects incorporate these impurities, along with nitrogen, when they crystallize in the earth’s mantle. Long residence times at high temperatures and pressures contribute to the final arrangement of the complex defects that produce green color in these diamonds, but their inherent color centers are imparted at “birth.” Diamonds colored by H3 defects have a somewhat more complex history. Nitrogen impurities, incorporated during crystal growth, aggregate into A-centers over millions to billions of years at elevated temperatures deep in the earth. Vacancies associated with H3 defects are most commonly created by plastic deformation as the diamonds are violently transported via erupting kimberlite magmas to shallower depths in the earth. The H3 defects are then created by mobilization of the vacancies to combine with A-centers as the diamonds reside at elevated temperatures. In most scenarios, the color centers in H3-based green diamonds form during magma transport (Fisher,
2009]. The final, and largest, category of green diamonds—those colored by GR1 defects—are generally thought to obtain their color due to late-stage interactions with radioactive minerals or fluids in the earth’s crust over thousands to millions of years. Areas of radiation damage in these diamonds range from green to brownish spots caused by direct contact with radioactive minerals (most likely uraninite) to zones of green- to brown-colored “skin” on the surface or within surface-reaching fractures caused by radioactive fluids (figure 8; see Vance and Milledge, 1972; Vance et al., 1973; Mendelsohn et al., 1979). The shallow nature [approximately 20 µm thick] of the green spots and zones indicates that the radiation damage is primarily caused by alpha particles [see box A for radiation types and sources], most likely from uranium decay [Vance et al., 1973; Nasdala et al., 2013]. These spots have been experimentally shown to rapidly turn from green to brown at temperatures of 550–600°C [Nasdala et al., 2013; Eaton-Magaña and Moe, 2016], indicating that diamonds with green radiation damage must have resided at depths less than about 24 km [assuming average geothermal gradient of 25°C/km].

Some green diamonds, however, show color zones that penetrate much deeper than 20 µm, sometimes even throughout the stones. In these cases, it is likely that some form of beta particle decay [or possibly gamma radiation] is contributing to the irradiation [box A]. Some beta decay occurs as part of the same U-Th-Pb decay chain that is likely responsible for the stain-producing alpha damage described above. The overall doses of beta particles possible from these later parts of the decay chain are unknown and the subject of ongoing research. Another viable source for beta decay in the earth’s crust with a half-life on
the order of millions of years appears to be potassium (\textsuperscript{40}K), but its natural abundance is very low (box A), leaving the complete story of radiation-related green color zoning unclear—in part because so few diamonds with green bodycolor have been available for scientific study.

**GEMOLOGICAL OBSERVATIONS**

Green diamonds have many interesting natural features that can be observed using a gemological microscope. These properties are quite variable, however, depending on the nature of the green color. In this section, we will review the major observations on green diamonds colored by each group of defects. Keep in mind that it is not uncommon for color to be affected by more than one of these color causes simultaneously. For example, many hydrogen-rich diamonds from Zimbabwe also show signs of radiation damage.

**Primarily Colored by GR1.** The most distinctive and important gemological features of green diamonds colored by radiation damage are surficial radiation spots or coatings (commonly referred to as “stains” if localized and “skin” if generally distributed) and internal color zonation (figure 9, A–E). Radiation stains occur as either green or brown patches of color on the surface of a rough diamond or within fractures. The radiation damage initially has a forest green color (often very dark green, depending on the dosage of the exposure) and then turns olive green at temperatures between 500 and 600°C and a reddish brown to dark brown at higher temperatures (Vance et al., 1973; Nasdala et al., 2013; Eaton-Magaña and Moe, 2016). Some experiments have shown that the brown stains disappear at temperatures above 1400°C (Eaton-Magaña and Moe, 2016). Radiation stains are thought to be a product of alpha particle bombardment, and structural damage increases to-

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**Figure 8.** In the earth, diamonds may be exposed to radiation emitted from tiny radioactive mineral grains adjacent to the stones or through interaction with radioactive fluids. The radiation produces either isolated circular radiation “stains” (often from adjacent minerals in the kimberlite host rock, left) or radioactive fluids (usually in alluvial environments) that surround a stone and penetrate into fractures. Photo on the top left courtesy of William F. Larson. Photo on the top right by Robison McMurtry.
**Box A: Summary of Radioactive Decay**

Radioactivity primarily occurs by three mechanisms: alpha decay, beta decay, and gamma emission (figure A-1, top left). Alpha decay involves the expulsion of an alpha particle [two protons and two neutrons—equivalent to a helium atom nucleus] as the parent isotope decays to the daughter. Alpha particles are relatively large and slow, causing them to have low penetration depths (stopped by a sheet of paper, skin, or even air; figure A-1, top right). In natural-color green diamonds, alpha particles produce very thin green spots or skins on the stone’s surface due to their shallow penetration depth. Beta decay involves ejection of a beta particle [an electron] as parent converts to daughter. Beta particles are small, high-energy, and high-speed, with moderate penetration depths (stopped by a sheet of aluminum). This type of radiation produces internal color zones in diamond at varying depths and is the most common type of radiation used for diamond treatment. In gamma emission, a parent isotope in an excited nuclear state decays to a daughter isotope by release of gamma rays. These extremely small forms of electromagnetic radiation are the fastest, highest-energy, and most penetrative radioactive decay product. They can fully penetrate a large gem diamond to impart uniform coloration and require a sheet of lead to stop their progress.

As shown in figure A-1 (bottom), the most likely sources of alpha decay in the earth are the radioactive isotopes of uranium ($^{238}\text{U}$ and $^{235}\text{U}$) and thorium ($^{232}\text{Th}$). Radiogenic thorium is three times more abundant than uranium in the earth’s crust, but its decay is very slow, with a half-life (the time it takes for half of the parent isotope to decay to the daughter) of 14 billion years. $^{238}\text{U}$ is the most abundant radiogenic uranium isotope and has a half-life of 4.5 billion years, making it a more feasible source of radiation damage for diamond. Most uranium is found in the minerals uraninite and zircon. The best sources of beta decay to create green bodycolor in diamond include late-stage radiogenic decay within the uranium-lead decay chain and potassium. $^{40}\text{K}$, the only significant radiogenic isotope of potassium, has a similar abundance in the earth’s crust and a much shorter half-life of 1.3 billion years (figure A-1, bottom). $^{40}\text{K}$ is hosted in common crustal minerals such as micas and feldspars. Because these radioactive isotopes are extremely scarce in the earth’s mantle, most green diamonds are thought to obtain their color from radiation exposure in alluvial deposits, where crustal minerals are more abundant, rather than in their kimberlite host rocks. In addition, the elevated temperatures of the mantle cause vacancies in the diamond lattice to be extremely mobile, and thus unstable.

<table>
<thead>
<tr>
<th>Decay type</th>
<th>Radiation emitted</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alpha decay</td>
<td>$^4_2\alpha$</td>
<td>![Alpha] Parent, Daughter Alpha particle</td>
</tr>
<tr>
<td>Beta decay</td>
<td>$^0_{-1}\beta$</td>
<td>![Beta] Parent, Daughter Beta particle</td>
</tr>
<tr>
<td>Gamma emission</td>
<td>$^0_0\gamma$</td>
<td>![Gamma] Parent (excited nuclear state), Daughter Gamma ray</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Radioactive isotope</th>
<th>Main decay type</th>
<th>Half-life (billions of years)</th>
<th>Avg. abundance Earth’s mantle (ppm)</th>
<th>Avg. abundance – Earth’s crust (ppm)</th>
<th>Common mineral host</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{238}\text{U}$</td>
<td>Alpha</td>
<td>4.5</td>
<td>0.022</td>
<td>1.787</td>
<td>Zircon, uraninite</td>
</tr>
<tr>
<td>$^{235}\text{U}$</td>
<td>Alpha</td>
<td>0.7</td>
<td>0.0002</td>
<td>0.011</td>
<td>Zircon, uraninite</td>
</tr>
<tr>
<td>$^{232}\text{Th}$</td>
<td>Alpha</td>
<td>14.0</td>
<td>0.083</td>
<td>5.999</td>
<td>Zircon, titanite, monazite, thorianite</td>
</tr>
<tr>
<td>$^{40}\text{K}$</td>
<td>Beta</td>
<td>1.3</td>
<td>0.031</td>
<td>1.800</td>
<td>Micas, clay minerals, K-feldspar</td>
</tr>
</tbody>
</table>

Figure A-1. Radioactive decay usually involves emission of alpha or beta particles or gamma rays. Each type of radiation has a different penetration depth in diamond, with alpha particles only affecting the surface and gamma rays penetrating throughout. Isotopes of uranium, thorium, and potassium are the most likely to affect diamonds in natural settings. These isotopes vary in abundance in the earth.
Figure 9. Radiation stains in diamond (A–E) occur initially as green spots that may turn brown with exposure to heat above 500°C (Vance et al., 1973; Nasdala et al., 2013; Eaton-Magaña and Moe, 2016). The stains are caused by alpha decay and occur as spots or thin coatings on the surface (A, C, and E). Green color usually penetrates only a few micrometers deep (A). Natural indentations or fractures allow radioactive fluids to penetrate and leave stains within the features (B). Occasionally, radioactive fluids between grain boundaries of adjacent minerals preserve their positions as green spotted patterns (D). H3-colored green diamonds (F) usually show yellow bodycolor with green luminescence. H-rich diamonds often have distinctively patterned clouds of microinclusions (G). Photomicrographs by various GIA staff.

ward the center of the area. While the stains can occur in any shape or size, depending on the diamond’s surface exposure to radioactive material, most take the form of circular spots that likely represent halos extending from multiple point sources (figures 8 and 9A, 9C, and 9E). Damage to the atomic lattice by alpha particles causes a slight volume expansion in the diamond and an elevated surface at the locus of the stain (Nasdala et al., 2013). The green or brown color penetrates to depths of usually not much more than 20 µm (0.02 mm), creating a shallow zone of green color adjacent to the stain(s) (figure 9, A–B; Nasdala et al., 2013; Eaton-Magaña and Moe, 2016).

When many stains are concentrated together, a rough diamond may be covered with large patches of
green zones. Similarly, very thin skins of relatively uniform green color may coat the surfaces and fractures (figure 9, C and E). These more pervasive areas of radiation damage probably represent radioactive fluids [rather than point sources of particles of radioactive minerals] surrounding a diamond and penetrating into the fractures and crevices (figures 8 and 9B). Patterns of green staining often mimic grain boundaries of minerals that would have surrounded the diamond, suggesting that radioactive fluids were also present between mineral grains in some ancient alluvial environments (figure 9D; see also Mennell, 1915; Guo et al., 1986). Green color zones that extend deeper must be related to beta or gamma radiation (or possibly neutron bombardment). While these zones do occur in some natural diamonds, they are much less common and therefore must represent some unusual radiation exposure conditions. For example, the color of the 41 ct Dresden Green was described by Kane et al. [1990] as “evenly distributed throughout the gem,” despite its large size (29.75 × 19.88 × 10.29 mm). This may represent a diffused green color that penetrated uniformly because the diamond was surrounded by radioactive fluids.

Sometimes a complex history of residency in the earth can be gleaned from radiation stain patterns. A number of examples have been observed where green and brown radiation spots with identical shapes are offset from each other in one direction, sometimes with three or more instances of the same radiation spot along the same trajectory (figure 10). Due to the dynamics of how the spots change color with temperature, we know that the farthest brown spot was oldest and that the stone subsequently moved one or more times due to some tectonic or compaction process while the radiation source remained present. Heating episodes exceeding 500°C must have accompanied the shifting process to cause the spot color changes, followed by a cooling and subsequent re-irradiation of a new spot [Vance et al., 1973]. The change in diamond position relative to its environment, combined with the unchanged spot shape, suggests that these spots were adjacent to radioactive mineral grains rather than fluid pockets.

**Colored by H3.** As mentioned earlier, H3 defects primarily cause yellow bodycolor in diamond, with the green component caused by a strong undertone from a luminescence reaction to visible light. Using a fiber-optic or other strongly focused light source, green luminescence (sometimes referred to as “transmission”) with a watery appearance is usually seen in green diamonds colored by H3 defects (figure 9F; see also Kane, 1979a). Although the green color may be uniform in the stone, the structure of H3 as a combination of A-centers with vacancies means that if nitrogen aggregate pairs and vacancies are irregularly distributed, the luminescence will show a similar pattern. For example, diamonds with grain lines caused by plastic deformation sometimes have higher concentrations of A-centers along the graining [Massi et al., 2005] and higher amounts of vacancies available [Fisher, 2009]. Consequently, fluorescence from H3 defects in these diamonds can often be seen as an alternating banded pattern between green and nonfluorescent regions.

**H-Related Green Color.** Unlike the two previous groups, direct evidence of the defects related to green color in H-rich diamonds is typically not visible with a microscope. Some features, however, do occur primarily in diamonds with a high concentration of hydrogen defects. The most common features of these stones are milky particulate clouds that have sharp,
angular boundaries with relatively non-included regions. Sometimes these clouds outline a six-rayed star pattern that is well known in hydrogen-rich stones (figure 9G; see also Wang and Mayerson, 2002; Rondeau et al., 2004). When viewed under UV light, H-rich diamonds often fluoresce yellow as well.

**Ni-Related Green Color.** Gemological observations from Ni-rich green diamonds are generally not distinctive. Some have been reported to show a uniform weak green luminescence to visible light, similar to that from H3, but it has not been directly linked to the Ni impurities.

**LABORATORY GRADING**

Stones weighing up to 2.0 ct comprised 82% of GIA’s intake of natural green diamonds, with those between 1.0 and 2.0 ct accounting for nearly 40% of that amount (figure 11). Rounds were most prevalent (36%), followed by cushions (26%), cut-cornered rectangles (12%), and pears (9%) (figure 11). Color grade distribution, as mentioned earlier, included 54% yellow-green, 28% pure green, and 12% blue-green, with the remainder having gray or brown undertones (figure 12). Many green diamonds are colored by a combination of defects, making it difficult to assign specific percentages to each cause of color. As an estimate, we used a random sample of 250 natural green diamonds colored by the four main defects to determine that 34% were colored by GR1, 21% by H3, 16% by hydrogen defects, and 1% by nickel defects, while the remainder included some combination of these color centers (figure 12). No discernible trends were observed from clarity grades.

**ABSORPTION SPECTROSCOPY**

Absorption spectroscopy measurements are nondestructive and provide a detailed look at a diamond’s atomic structure by passing light of different wavelengths through the stone and measuring what wavelengths [energies] are absorbed by the impurities and defects present. IR absorption gives information about the impurities in the diamond lattice, while ultraviolet-visible (UV-Vis) absorption reveals the color-producing defects. Each group of green color defects has unique spectroscopic features.

**IR Absorption.** Details of the major impurity present (nitrogen or boron), as well as the configuration of the atoms of those impurities relative to the rest of the carbon lattice in diamond (diamond type), are quickly and directly measured from IR spectroscopy. In addition, information about other impurities such as hydrogen can be inferred through evaluation of IR features created by complex defects involving those atoms. Infrared spectra are typically collected with the stone kept at room temperature conditions (Anderson, 1943a, 1943b, 1943c; Collins, 2001; Zaitsev 2003).

Figure 11. Most of the natural green diamonds submitted to GIA during the last decade have weighed 1–2 ct or less. Fewer than 5% are larger than 4 carats. Rounds were the predominant shape, followed by cushion, cut-cornered rectangle, and pear shapes.
Green diamonds colored by H3, H, and Ni defects are all type Ia [containing N impurities]. Those with significant H3 defects usually contain low to moderate concentrations of nitrogen, typically both as A-centers (pairs) and B-centers [4N+V] along with associated platelet defects (figure 13; see also Woods, 1986 for platelet defect description). Higher-nitrogen diamonds tend to show very little to no green color from H3 luminescence, even when H3 concentrations are high, because the high levels of A-centers tend to quench the green fluorescence, leaving only the yellow bodycolor due to absorption of H3 (Collins, 2001; Luo and Breeding, 2013). Low concentrations of H-related defects seen at 4496, 3107, and 1405 cm⁻¹ are not uncommon. Green diamonds that owe their color to H-related defects have very high concentrations of both H-related defects [4496, 4167, 3236, 3107, 2812, 2785, and 1405 cm⁻¹] and N impurities in the form of A- and B-centers, as well as platelets [1360–1370 cm⁻¹] and related peaks at approximately 1435, 1498, 1525, and 1549 cm⁻¹ that shift depending on the A/B center ratio. The concentrations of aggregated N impurities in these stones are so high as to saturate most IR spectrometer detectors [i.e., the peaks extend beyond the upper edge of the infrared spectrum]. Green stones colored primarily by Ni-related defects contain very low amounts of nitrogen impurities, predominantly as A-centers [with only occasional traces of B-centers]. In many cases, IR spectra also reveal traces of C-centers [i.e., single, substitutional nitrogen] as well as a series of peaks including 3137, 3143, 3180, 3202, 3242, 3257, and 3272 cm⁻¹ that are ascribed to H + C-center defects. Relatively low but variable amounts of the H-related defects seen in H3 diamonds are also present.

Diamonds colored green by GR1 defects from natural radiation span the gamut from high-nitrogen type Ia stones to type Iia diamonds with no measurable nitrogen impurities (figure 13). Likewise, the H content varies widely. The reason for this lack of correlation is the secondary nature of the radiation damage. Any diamond can be exposed to radioactive minerals or fluids after its formation to develop GR1 defects and possibly a green color. The preexisting impurities and other defects will determine whether the green stone resulting from radiation damage has a yellow, gray, or brown undertone, but the green color is entirely introduced secondarily from the original condition of the diamond. One radiation-related defect that occurs as a weak peak in the IR spectrum of some intensely green type Ia natural diamonds is
H1a, an interstitial nitrogen defect at 1450 cm$^{-1}$ (Zaitsev, 2003; Dischler, 2012).

**UV-Vis-NIR Absorption.** Ultraviolet/visible/near-infrared spectroscopy measures the color center absorptions that produce the color we see in a diamond. Although most primary impurity absorptions occur in the IR region, many complex defects (and even some very simple ones) absorb light in the visible part of the spectrum to produce color. These spectra are typically collected at liquid nitrogen temperature (approximately 77 K) to observe narrow peaks from temperature-sensitive defects (Collins, 2001; Shigley and Breeding, 2013).

In the “Causes of Color” section, we discussed the main visible absorption features responsible for color in green diamonds containing GR1, H3, H-related, and Ni-related defects. Here we will detail the less prominent features that tell us more about the nature of the coloring mechanism and how it relates to the color we see. For diamonds colored by H3 or Ni defects alone, UV-Vis-NIR spectra tend to be relatively consistent from stone to stone because both of these defect centers intrinsically coexist with aggregated nitrogen, which always absorbs light toward the blue end of the spectrum and commonly occurs with N3 defects $\{3N+V\}$ at 415 nm (figure 14). The same is generally true of H-rich diamonds, except that these stones often have accompanying absorptions related to H at 615 nm and more distinctive “cape” absorptions related to nitrogen at 478 nm ($N2$). Additionally, H-rich green diamonds often have a broad band centered at around 530 nm (figure 14). The combination of this 530 nm band with the N-
and H-related visible absorptions causes the lower absorbance part of the spectrum between 500 and 600 nm to be elevated, resulting in the gray undertone that is common in H-rich green diamonds.

Radiation damage in diamond produces a number of defect centers in Vis-NIR spectra in addition to the primary GR1 color center (figure 14). The four most common of these defects occur at 496 (H4), 503.5 (3H), 594, and 667 nm. Natural diamonds remain buried in the near-surface region of the earth for variable amounts of time even after they are exposed to radioactive minerals or fluids. Depending on the depth of residency, green diamonds are commonly exposed to elevated temperatures of a few hundred degrees Celsius that cause a natural annealing effect (Vance et al., 1973). The 667 nm, 594 nm, and 3H defects can all occur from radiation damage alone, without annealing, and are often seen in stones with green radiation stains. The 667 nm center consists of a carbon interstitial that anneals out at 400–500°C (Zaitsev, 2003). The 594 nm center (known to gemologists as the 595 line) is a defect of uncertain structure, possibly consisting of nitrogen atoms and vacancies or interstitials (Collins, 1999; Zaitsev, 2003). It anneals out at approximately 1000°C. The 3H defect is a split carbon interstitial that increases in intensity with annealing and remains stable to at least 600°C (Zaitsev, 2003). The H4 defect at 496 nm is more complex, consisting of four N atoms and two vacancies, and generally forms with radiation damage and subsequent annealing to temperatures above 600°C. H4 is very stable and withstands temperatures of at least 1400°C (Collins, 1997; Zaitsev, 2003).

It is important to remember that radiation damage can also occur simultaneously with any of the other defects that cause green color. It is not uncommon to see H-rich or H3 diamonds with GR1 defects (figure 12, right). We have even observed the occasional Ni-rich diamond with GR1. The combination of multiple causes of green color can dramatically intensify the overall green appearance but often makes identification more difficult, as we will discuss later.

LUMINESCENCE SPECTROSCOPY AND IMAGING

Luminescence spectroscopy measures the light emitted by a diamond in response to stimulation by another source of light, such as a laser or UV lamp. Photoluminescence (PL) uses lasers of different wavelengths to produce emission spectra that reveal the defects present in a stone. DiamondView imaging exposes diamonds to very high-energy UV light (<230 nm) to stimulate fluorescence that can be imaged with great detail (Welbourn et al., 1996). Both techniques are nondestructive and provide important information about the structure of natural green diamonds.

PL Spectroscopy. PL analysis is one of the most useful and sensitive techniques for defect characterization in diamond. Defects occurring at concentrations
as low as the ppb (parts per billion) level can easily be detected (Breeding et al., 2010; Eaton-Magaña and Breeding, 2016). Like UV-Vis-NIR absorption, PL measurements of green diamonds are usually focused in the visible range of the electromagnetic spectrum and need to be collected at low temperatures (around 77 K) to resolve many spectral peaks. There are hundreds of PL peaks that might occur in diamond, and some are activated only by certain laser excitation wavelengths since each laser excites luminescence features from diamond in a particular limited range of wavelengths. It would be impossible to characterize every feature—and little is known about many of them—so we have chosen to report representative PL features that are associated with each group of green diamonds.

Green diamonds colored by H3 defects show relatively few prominent PL features. The main peaks are H3 (503.2), 535.8, 575 (NV\(^0\)), 637 (NV\(^-\)), 741 (GR1), and 986 nm (H2) (figure 15). H2 has the same structure as H3 (N-V-N) but carries a negative electrical charge. Negatively charged defects indicate the presence of trace amounts of single substitutional nitrogen impurities serving as electron donors (Collins et al., 2000; Collins, 2001). The nature of the 535.8 nm feature is unknown, and PL-detectable trace amounts of GR1 are common in many natural diamonds. H-rich green diamonds also contain very few notable PL features. Peaks at 694, 700, 716, 787, and 793 nm are usually encountered with occasional H3 defects (503.2 nm; figure 16). Interestingly, most of these defects (694, 700, 787, and 793 nm) have previously been assigned to complex Ni-N defects (Bokii et al., 1986; Zaitsev, 2003). From our observations, at least two of them, 700 and 793 nm, correlate with A-aggregate concentrations in high-H diamonds. It is possible that these H-rich stones are also high in Ni content but, due to natural annealing and trapping of the Ni impurities by more-abundant aggregated nitrogen, no longer show the 883/885 nm Ni defect. If so, they represent a more N-rich, mature form of Ni-related green diamond. Ni-colored green diamonds show many of the same PL features, including the 700, 787, and 793 nm peaks discussed above that are attributed to Ni-N complexes. Yet these stones also show many other narrow PL features, among which the most consistent and prominent are 603 and 668 nm along with the very strong and well-known Ni doublet at 883/885 nm described earlier (figure 17; see also Zaitsev, 2003; Wang et al., 2007). Little is known about the 603 nm PL feature, but the one at 668 nm has been attributed to a Ni defect (Zaitsev, 2003).

PL spectra collected from diamonds colored by GR1 can vary considerably due to irregular distributions of radiation damage across a stone’s surface and interior. PL features related to radiation damage are also dramatically affected by heating, even to relatively low temperatures of a few hundred degrees. In order to represent the PL defects of natural radiation damage, the features below are from Brazilian green diamonds with abundant green radiation stains (meaning they have not been heated to 550–600°C). Similar stones from other localities show the same...
features. The strongest PL peaks in these green diamonds are stable defects at 741 and 744 nm (GR1), 503.2 nm (H3), 637 nm (NV⁻), and 986 nm (H2). Weaker, much less stable features at 526, 587.9, 647.2, and 722.5 nm are all related to interstitial defects seen in the H-rich stones (though they are much stronger here), as well as the 883, 885 nm Ni-related doublet. Intrinsic diamond Raman lines have been removed for clarity.

Figure 17. PL spectra from Ni-rich green diamonds contain abundant peaks, including the same Ni-related defects seen in the H-rich stones (though they are much stronger here), as well as the 883, 885 nm Ni-related doublet. Intrinsic diamond Raman lines have been removed for clarity.

one of the authors (CMB) has shown that these two are related to each other and likely represent interstitial N atoms. The 647.2 and 722.5 nm PL features are completely destroyed by heating to 500°C, and thus do not occur in natural green diamonds with radiation stains that have turned brown.

**DiamondView.** While photoluminescence records a semi-quantitative spectrum of the light emitted from a diamond at a single spot, DiamondView imaging records the qualitative fluorescence pattern produced...
over the entire instrument field of view to reveal spatial variations in luminescence. These patterns can be used to separate natural from laboratory-grown diamonds, but also to interpret some of the stone’s impurity incorporation and growth history. The DiamondView uses ultra-short-wave UV light (<230 nm) to excite fluorescence from diamonds at room temperature (Welbourn et al., 1996). During UV exposure, the diamond can be rotated to observe fluorescence patterns in several directions.

In green diamonds colored by H3 defects, the most obvious feature shown in the images is green fluorescence from H3 (figure 19), which often follows particular growth zones or plastic deformation structures. Blue fluorescence from N3 defects is also usually visible. The ratio of green to blue fluorescence in the images directly correlates with the total concentration of H3 defects. H-rich diamonds typically show weaker overall fluorescence that is almost always blue from N3 defects. Irregular patterns reflect
cuboid growth, which is often apparent with H diamonds. Isolated green H3 luminescence occurs occasionally at the boundaries between growth zones. Ni diamonds usually contain multiple zones of contrasting fluorescence colors. Blue from N3 is almost always present along with inert zones and/or regions that fluoresce yellowish green (figure 19). The green may be from a Ni-related defect, as there are typically few or no H3 defects seen in these stones. The DiamondView images from green diamonds colored by GR1 are extremely variable because they primarily reflect the defects present in the stones prior to natural irradiation. The observations suggest that high levels of radiation damage tend to quench fluorescence, resulting in areas with radiation staining that appear inert in DiamondView images (figure 19).

IDENTIFICATION CONCERNS
Identification of colored diamonds is a two-step process involving determination of genesis (grown in the earth versus a laboratory) and color origin (a product of natural processes versus treatment). In the case of green diamonds, the first step is relatively easy, but the color origin determination is often problematic. While all the diamonds discussed in this article were identified by GIA as having natural color, green diamonds in general are the single largest group of diamonds that receive “Undetermined” color origin reports from gemological laboratories. Laboratory irradiation using an electron beam takes only a few minutes and can be performed commercially at low cost. This treatment has been performed regularly since the 1950s to create green diamonds, so nearly every green stone in the trade must be regarded with suspicion. Other treatment methods such as embedding diamonds in radioactive compounds, cyclotron and linear accelerators, and gamma ray exposure in nuclear reactors have been employed less frequently to achieve a green color.

As might be expected with the similar processes in nature and the laboratory, the absorption and luminescence spectra from natural- and treated-color green diamonds are virtually identical in many cases (figure 20). Furthermore, almost any kind of diamond can be subjected to irradiation (in nature or the laboratory) to produce a green hue. This means that the preexisting defects, and thus the resulting radiation-related defects, can vary as much as natural heterogeneity in diamond itself. In the absence of obvious color zoning related to laboratory treatment (green culet and a cyclotron “umbrella” pattern), all green diamonds should be sent to a gemological laboratory.

At GIA, we continue to conduct our own irradiation treatment experiments and to examine known natural-color green diamonds from around the world to refine our identification criteria and reduce the number of “Undetermined” reports issued each year. As with all instances of gem testing, the identification of natural-versus treated-color in diamond is based on a variety of features. These identification
Among green diamonds, chameleon diamonds are a fascinating curiosity. They do not directly owe their green color to any of the four major defects discussed in this article. In fact, they span a wide range of color, from predominantly yellow to green hues. What makes these stones particularly interesting is that when heated or left in the dark for an extended period of time (days to weeks), they temporarily lose their green color and change to yellow or orangy hues (figure B-1, top) through energetic shifts in the broad visible absorption bands that cause their color. Within a few seconds to several minutes of being removed from darkness or allowed to cool after heating, chameleon diamonds regain their greenish color. The nature of the color change remains unclear, but these unique diamonds have a cherished niche among collectors (Fritsch et al., 1995; Hainschwang et al., 2005; Fritsch et al., 2007).

Over the last decade, a few thousand chameleon diamonds have been documented within GIA’s laboratories; the vast majority had a stable color that included a green contribution (figure B-1, bottom left). Nearly half were grayish yellow-green, and only about 1% had pure green hues; 83% of the chameleons seen at GIA had gray or brown color components as well. Chameleon diamonds were generally smaller than other natural green diamonds (e.g., figure 11), with most weighing less than 1.0 ct (figure B-1, bottom right).

UV fluorescence was a distinctive feature among the chameleon diamonds. Of these, 74% showed yellow fluorescence and another 18% exhibited orange or orangy yellow fluorescence in response to long-wave UV exposure. The intensity of the long-wave fluorescence was predominantly strong (92%). Under short-wave UV illumination, 86% showed yellow fluorescence and another 13% exhibited orange or orangy yellow fluorescence. The intensity of the short-wave UV fluorescence was predominantly medium (85%). Additionally, all the chameleon diamonds exhibited short-wave UV phosphorescence that was usually weak in intensity (54%) and yellow in color (96%).

GIA laboratory reports include a special comment indicating the unique color-change property of a chameleon diamond. To receive this comment on a report, a diamond must have a green color component (either primary or secondary), it must display phosphorescence to short-wave UV light, and it must change color from greenish to yellow or orange upon gentle heating or removal from light. Tremendous caution must be taken when heating green diamonds, as those colored by GR1 defects may permanently lose some or all of their green color. While chameleon diamonds cross the boundary between green and yellow stones, they remain an interesting oddity that accentuates the novelty of green diamonds.

Figure B-1. Chameleon diamonds (top) change from a greenish color to a yellow or orange hue when heated or kept in the dark. Of the chameleon diamonds seen at GIA over the last decade (bottom), most have stable colors of grayish yellow-green and brownish greenish yellow and less than 0.5 ct in size. Photos by C.M. Breeding.
Figure 21. While the vast majority of green diamonds fall into the four categories described in this article, a few very unusual stones with green hues are type IIb, type Ib, or chameleon stones. These very rare diamonds have distinctive spectroscopic features that make them relatively easy to spot.

criteria are considered proprietary information to protect the integrity of GIA’s grading report service. To support our conclusion on color origin, dealers sometimes bring an important rough green diamond for us to examine before manufacturing and then during several stages of the process. This allows us to study the progression of the color appearance of the cut stone, and to confirm that it originated from the starting piece of rough.

UNUSUAL EXAMPLES

We have discussed the four major causes of green color in gem diamonds. These are responsible for some 95% of the naturally green diamonds seen at GIA’s laboratories in the last decade. As with anything natural, there are a few unusual diamonds with green color components that do not fall within the aforementioned groups. These include type IIb and Ib stones as well as chameleon diamonds (figure 21 and box B). Type IIb diamonds are typically blue due to boron impurities that absorb light from the infrared all the way through the visible spectrum, with the absorption higher at the red end of the spectrum than the blue end. In a few extremely rare cases, a type IIb diamond also has strong absorption from 400 to 500 nm, resulting in a green hue (Crowningshield, 1965; Scarratt, 1989). The cause of the higher-energy absorption is unclear, but it might be elevated trace
concentrations of nitrogen impurities that are masked in the IR by the excessive boron. Type Ib diamonds contain single substitutional nitrogen impurities that absorb strongly in the UV to blue part of the spectrum, generating a strong yellow color. If a type Ib diamond is plastically deformed and has enough vacancies to increase the NV–[637 nm] defect concentration significantly enough to absorb light in the red region, the stone has a greenish hue (usually also grayish or brown from the plastic deformation vacancy cluster absorption, causing a rise to higher energies). The last and perhaps most abundant and interesting oddity is the chameleon diamond (box B). Chameleons are type IaA diamonds (usually with moderate N content) that exhibit broad 480 nm (unknown structure) and 700 nm (H- or Ni-related) bands in the visible spectrum that often give rise to a greenish hue (usually brownish or grayish and termed “brackish”). These popular color-changing collectors’ stones occupy their own niche in the diamond trade.

CONCLUSIONS
Natural-color green diamonds (figure 22) are rare and beautiful, with a variety of defects that can be responsible for their color. Today, most are discovered in South American or African mines. The main causes of green color in gem diamonds, either as the sole mechanism or in combination, are vacancies produced by radiation damage [GR1], H3 defects, H-related defects, or Ni impurities. Each of these gives rise to a slightly different green hue. GR1-colored green diamonds are most abundant, followed by those colored by H3 green luminescence. H- and Ni-related green stones are less common. Each group of green color centers displays gemological clues as well as absorption and luminescence spectra that aid in their identification. Both GR1 and H3 defects can be easily induced in diamond by laboratory irradiation and annealing, a process that has been applied since the 1950s. GIA has made significant progress in determining the origin of color of green diamonds since the 1950s and ’60s, when the vast majority were described as undetermined. Today, most green diamonds have a definitive origin determination. Due to the similarities between natural radiation damage and laboratory treatment effects, color origin determinations of green diamonds are not always possible and have turned these rare and exquisite gemstones into the beautiful conundrum they are.
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