CHARACTERIZATION OF EMERALDS FROM A HISTORICAL DEPOSIT: BYRUD (EIDSVOLL), NORWAY

Benjamin Rondeau, Emmanuel Fritsch, Jean-Jacques Peucat, Fred Steinar Nordrum, and Lee Groat

An emerald deposit at Byrud, in southern Norway, yielded significant quantities of crystals and gem rough in the late 19th and early 20th centuries. Complex multiphase inclusions in the emeralds consist of water, gaseous methane, halite, sylvite, calcite, and a sulfide assemblage (pyrrhotite, galena, and sphalerite). This sulfide assemblage makes it easy to distinguish Byrud emeralds from those from other localities with a binocular microscope. The chemical composition of Byrud emeralds is also characteristic: They are colored mostly by vanadium (up to 1 wt.% V₂O₅), and contain low sodium and magnesium (0.1 wt.% oxide or less). Moreover, the relative amounts of iron, magnesium, chromium, rubidium, and cesium appear to be diagnostic. Infrared absorption spectra show that they contain little water. Emeralds from the Byrud deposit are still occasionally recovered by hobbyist collectors from the mine dumps.

During the late 19th and early 20th centuries, the Byrud emerald deposit in Norway was mined commercially and produced many fine specimens [e.g., figure 1], as well as a limited amount of gem rough. Some of the crystals are housed in museum collections around Europe, and it is not uncommon to encounter a Byrud emerald in an antique jewelry piece. The mine is located on the shore of Lake Mjøsa, near Minnesund and a short distance from Eidsvoll, about 60 km north-northeast of Oslo, Norway [figure 2]. The history, geology, and mineralogy of the deposit were comprehensively described by Nordrum and Raade (2006), and are summarized here. The aim of the present article is to characterize the emeralds from Byrud to make their unambiguous identification possible.

From a geologic and spectroscopic standpoint, emeralds from Byrud are interesting because they are vanadium-rich, as is also the case for emeralds from Colombia and some [or all] emeralds from a number of other deposits: Lened in the Northwest Territories, Canada; Salininha in Bahia, Brazil; Malipo in Yunnan, China; Panjshir, Afghanistan, and Gandao, Pakistan. In this article, we refer to “vanadium emeralds” as those that contain more vanadium than chromium, even when the vanadium content is somewhat low. The mechanisms of beryl coloration by Cr and V are very similar (Burns, 1993; Schwarz and Schmetzer, 2002); in 1988, after many years of controversy, vanadium joined chromium as an accepted coloring agent for emerald [CIBJO, 1988].

HISTORICAL BACKGROUND

The Byrud emerald deposit was probably discovered in the 1860s. During these early years, emerald specimens were obtained by several European natural history museums, including Stockholm in

See end of article for About the Authors and Acknowledgments.
1868, Oslo in 1869, and London in 1870. Websky (1876) first described the morphology of the Byrud emeralds, and stated that the deposit was of commercial importance.

Preliminary prospecting and blasting took place around 1880 (Bull, 1952). Evelyn Aston inspected the almost forgotten occurrence in November 1898; at that time, the old workings consisted of an opening 2 m high and 1 m wide that accessed a room measuring about 3.5 m in diameter. On her next visit, she brought a miner who blasted further, and they found gem-quality emerald crystals in small clay-filled pockets (Cameron, 1963).

On April 5, 1899, the prospect (called Narum) was purchased by Evelyn’s father, English mining prospector Edward Y. Aston. A London-based company—the Norwegian Exploration Co. Ltd.—was registered May 9, with Aston as a major shareholder in exchange for the property and rights to the emerald occurrence, as well as some additional prospects. Mining began in spring 1899 (figure 3). In June 1900, the company was renamed the Norwegian & General Exploration Co. Ltd. Edward Aston died on September 21, 1900, and on December 28, 1907, the company was liquidated and the assets sold to the Cornish Development Co. Ltd. The mine was subsequently abandoned in 1909 (Nordrum and Raade, 2006).

In the first years of mining, up to 30 miners were employed at Byrud. A crusher, washing plant, blacksmith’s forge, administration building, and a small workshop were reportedly built at the mine, but few vestiges are visible today. However, production could not sustain this level of mining for very long, and by the time the mine closed 10 years later, it had only nine employees. It does not appear that the mine was ever profitable, even during the first few years (Nordrum and Raade, 2006).

Nevertheless, many gem-quality emeralds were recovered, some of which were displayed at the Paris World’s Fair in 1900. Besides crystals, wrote Kunz (1902, p. 742), “Many cut stones, most of them pale in color, but generally free from flaws, were shown.” Sinkankas (1981, p. 487) stated: “Only rarely were good stones found and then never
more than about 6 mm in diameter, but the color quality was considered to be of the highest grade, and if anything a little too bluish. Selset (1963) cut some of the crystals himself and claimed that the gems matched the finest from any other source.”

Most of the transparent crystals from cavities were pale colored, but some were a fine, deep green. Cemented in feldspar, translucent crystals up to more than 1.2 cm in diameter and 5 cm in length with a deep green color were recovered. Translucent crystals were also commonly found in quartz.

A cut emerald from Byrud is said to have been used in a jewel belonging to the royal family of Great Britain, possibly in connection with the coronation of King Edward VII in 1902. The English companies sold the rough crystals and mineral specimens outside Norway, and no records of the sales have been found. The production data and names of the buyers are unknown (Nordrum and Raade, 2006).

Most of the emerald crystals pictured in this article were found by mineral collectors over the past 30 years (see, e.g., figure 4). For a small daily fee to the landowners, collectors have been allowed to search in the mine dumps. Several fine specimens and single crystals have been recovered over the years, but
the chances of making good finds are diminishing and gem-quality specimens are discovered only rarely today. The underground mine has not been worked since 1909.

GEOLOGY AND MINERALOGY

The emerald deposit is located in the northeastern part of the Oslo region, which is a rift structure of Permian age (Vogt, 1884; Goldschmidt, 1911; Nordrum and Raade, 2006). Flat-lying maenaite (syenitic) sills, usually ranging from 0.5 m to several meters thick, transect Cambrian alum shale (a black, clay-rich, iron sulfide-containing, carbonaceous sedimentary rock) over a distance of about 200 m. The sills generally dip 15–20° west but are nearly horizontal in places. They are present in at least three levels and have been intruded by small pegmatites, which also cross-cut the alum shales. Close to the pegmatites, the shales are often bleached. The pegmatites usually range from a few centimeters to 30 cm in thickness, but may locally reach up to 1 m. They form lenses or dikes that are commonly discontinuous and locally contain small cavities. The pegmatites have an alkali syenitic composition, consisting mainly of K-feldspar (microcline), and were intruded during the Permian Period in conjunction with alkaline magmatic activity in the area (Ihlen, 1978). A large alkaline granite intrusion occurs not far to the west of Byrud, and the pegmatites are most likely associated with this intrusive.

Emerald mining has taken place along the maenaite sills at various levels. Beryl occurs chiefly in the pegmatites (figure 5), and occasionally in the maenaites and in the shales adjacent to the pegmatites. The best-quality emeralds have been found in the northern part of the mining area (Lindaas, 1982), in small clay-filled pockets (Cameron, 1963). Vanadium and chromium, chromophores in the emeralds, were probably leached from the alum shales by the mineralizing fluids.

In addition to microcline, the pegmatites typically contain quartz and muscovite. Sodic plagioclase, pyrite, pyrrhotite, fluorite (purple and pale green), topaz, and beryl are also common in the pegmatites. Micro-crystals of laumontite and rutile ("illmenorutile-strüverite") are frequently present as well. Goethite, jarosite, and gypsum are common secondary minerals. In all, 45 minerals have been reported from this deposit (Nordrum and Raade, 2006; Kvamsdal and Eldjarn, 2007). A complete list of minerals associated with the emerald mineralization is available on the G&G Data Depository (www.gia.edu/gemsandgemology).

MATERIALS AND METHODS

The authors studied three emerald-in-matrix specimens, collected more than a century ago, from the mineralogy collection of the Muséum National d'Histoire Naturelle (National Museum of Natural History) in Paris (collection nos. 106.718, 195.174, 217.171).

Figure 4. Galleries excavated for emerald extraction during the late 19th and early 20th centuries are still visible. Mineral collectors, such as Arnfinn Juliussen and Bjørn Skår in this photo, can visit the mine with the permission of the owners, Anne Grethe Røise and Ole Jørgen Bjørnstad. Photo by F. S. Nordrum.

Figure 5. Byrud emerald crystals are found mainly in small pegmatites intruding alum shales. This sample is 9.5 cm long, and the emerald crystals attain lengths of 9 mm. Courtesy of the Norwegian Mining Museum; photo by Gunnar Jenssen.
and 203.18). These idiomorphic bluish green crystals, each a few millimeters in length, are intergrown with feldspar, quartz, muscovite, and purple fluorite. One emerald crystal was removed from each of the matrix specimens and all three were polished to measure their gemological, chemical, and spectroscopic properties. In addition, we studied four loose transparent crystals, each measuring a few millimeters in length, that were recovered in the 1980s and belong to the collection of the Norwegian Mining Museum in Kongsberg (collection nos. BVM 2378–A to D; figure 6). All seven samples studied were transparent and homogeneous in color, so we believe that they are representative of the material from which gems would be faceted. We did not study any cut stones or stones set in jewelry, as the very few that are known were not accessible.

Gemological data were acquired on various samples (see Results) using a Topcon refractometer, a 4-watt UV lamp with short-wave (254 nm) and long-wave (365 nm) bulbs, and an Olympus binocular microscope equipped with crossed polarizers and up to 1000× magnification. Specific gravity was determined by a combination of mass measurement using a precision scale and volume measurement using a classic pycnometer.

The composition of fluid inclusions in three samples (nos. 106.718, 195.174, and 203.18) was first determined by Raman spectroscopy using a Jobin-Yvon T 64000 dispersive spectrometer equipped with a confocal-type apparatus. The Ar+ laser (514 nm excitation) was operated with a power of 120 mW, and spectra were measured at a resolution of 2 cm⁻¹. Subsequently, one crystal (from sample 203.18) was broken to expose some of its fluid inclusions, and these were examined using a scanning electron microscope (SEM); the micro-crystals in about 30 of the inclusions were analyzed with an attached energy-dispersive spectrometer (EDS). We used a Zeiss Supra 55 VP SEM with an acceleration voltage of 7 or 22 kV, and a current of ~1 nA.

Electron-microprobe analyses of two samples (21 spots) were obtained with a fully automated Cameca SX-50 instrument, using the wavelength-dispersive mode with the following operating conditions: 15 kV excitation voltage, 20 nA beam current, 20 sec. peak count time, 10 sec. background count time, and 10 μm spot diameter. Data reduction was done using the “PAP” φ(ρZ) method (Pouchou and Pichoir, 1991). Trace-element compositions of three emeralds (14 analyses) were measured using laser ablation–inductively coupled plasma–mass spectrometry (LA–ICP–MS) with an HP4500 spectrometer. The CETAC SLX200 ablation system used a Nd:YAG laser emitting at 1064 nm, frequency quadrupled to 266 nm. We limited our sampling to three emeralds, because this technique is somewhat destructive and cannot be applied to museum specimens.

Polarized ultraviolet–visible–near infrared (UV-Vis-NIR) absorption spectra of four samples (BVM 2378–A to D) were acquired with a Varian Cary 5G spectrometer in the range 300–2500 nm, with a sampling interval of 1 nm and a spectral bandwidth of 1 nm, at a scan speed of 600 nm per minute, using 1 × 1 cm Glan-Thomson calcite polarizers. Infrared absorption spectra of three samples (nos. 106.718, 195.174, and 203.18) were acquired using a Nicoler TO5X spectrometer in the range 5800–1800 cm⁻¹. The spectra presented are directional (not polarized), with the beam (and not the electrical vector) parallel, and then perpendicular, to the optic axis.

RESULTS

Gemological Properties. Visual Appearance and Crystal Morphology. The emerald crystals were translucent to transparent and formed hexagonal crystals with flat basal terminations (figures 7 and 8). Pyramidal faces have also been seen on the terminations of Byrud emeralds (figure 8, right-hand crystal). The studied crystals were green to bluish green, and...
showed weak but significant pleochroism, from bluish green to yellowish green. All samples studied were homogeneous in color, though we have observed some crystals (not included in this study) that exhibit very strong color zonation parallel to the basal plane, with some parts nearly colorless (again, see figure 8).

Figure 7. This 1.2-cm-long emerald crystal is partially embedded in a spherulitic muscovite matrix. Courtesy of the Norwegian Mining Museum; photo by Gunnar Jenssen.

Refractive Indices. We measured the refractive indices from the natural prism faces of samples BVM 2378–A to D and obtained identical values of \( n_o = 1.578 \) and \( n_e = 1.560 \). Three crystals from sample 203.18 were polished parallel to the c-axis, and we measured \( n_o = 1.587 \) and \( n_e = 1.579 \). In all cases, the birefringence was 0.008.

Specific Gravity. We measured a specific gravity of 2.75 on five fragments of sample 203.18. Additional SG measurements could not be done because many samples were crystals on matrix (in particular, samples BVM 2378–A to D contained too much matrix material on their extremities).

UV Fluorescence. All samples were inert to both short- and long-wave UV radiation, as with most natural emeralds worldwide (Bosshart, 1991; Zylbermann, 1998).

Magnification. In the seven emerald samples studied with a gemological microscope, we observed a few solid inclusions; these showed a bright yellow metallic luster and were most probably pyrite. All the samples contained primary fluid inclusions, virtually all of which were multiphase (figure 9). We observed no secondary inclusions. Some inclusions had an irregular outline (figure 10), and others showed a somewhat hexagonal outline when viewed parallel to the c-axis of the host (again, see figure 9), indicating that crystal growth occurred mostly along the basal planes.

Figure 9. Multiphase inclusions are commonly observed in emeralds from Byrud. They typically show regular geometric outlines and contain a liquid and a gaseous phase, a cubic transparent solid phase, elongated transparent crystals, and opaque phase(s). Photomicrograph by B. Rondeau; magnified 200×.
The most characteristic multiphase inclusions (figure 11) contained predominantly a liquid phase, a gas bubble, an optically isotropic cube, some transparent, birefringent elongated material (as revealed between crossed polarizers, figure 11 right), and very small opaque phases. When these opaque phases are sufficiently large, one can see that they have a brownish yellow metallic luster (best observed using fiber-optic lighting from the side).

**Composition of the Multiphase Inclusions.** Raman Spectroscopy. Raman analyses revealed that the transparent cubes were halite and the aggregates of elongated opaque material were calcite \((\text{peaks at } 1083, 279, 182, \text{ and } 151 \text{ cm}^{-1})\). The liquid was water \((\text{broad band around } 3600 \text{ cm}^{-1})\), and the gaseous phase was a mixture of water and methane \((\text{CH}_4; \text{broad band around } 3600 \text{ cm}^{-1} \text{ and sharp peak at } 2915 \text{ cm}^{-1})\).

SEM-EDS Analysis. Fresh breaks on a fragment of sample 203.18 exposed numerous fluid inclusions. The liquid and gaseous phases were flushed out of the inclusions during sample preparation, and the solid phases were lost from some of the cavities as well. In those that still contained solids (figures 12 and 13), the most common phase identified was halite \((\text{NaCl})\); sylvite \((\text{KCl})\) was also found frequently. Several sulfide phases were observed: pyrrhotite \((\text{Fe}_{1-x}\text{S}, \text{hexagonal flakes})\), galena \((\text{PbS})\), and sphalerite \((\text{ZnS})\); these were often grouped together (again, see figure 12). Copper was also detected during the analysis, but we could not ascribe it to a specific mineral. Some rarer solid phases were also identified. In particular, we found one cassiterite crystal \((\text{SnO}_2)\) and one crystal of a phase that contained phosphorus, oxygen, and scandium that we attributed to the very rare mineral pretulite \((\text{ScPO}_4)\) (again, see figure 13).

**Chemical Composition.** Electron-microprobe analyses are summarized in table 1, and LA-ICP-MS analyses are provided in table 2; additional data from both techniques are available on the G&G Data Depository (www.gia.edu/gemsandgemology). We measured relatively high concentrations of V \((\text{up to } 2.44 \text{ wt.\% } \text{V}_2\text{O}_3, \text{ with a mean of more than } 1 \text{ wt.\%})\) and comparatively moderate concentrations of Cr \((\text{up to } 0.33 \text{ wt.\% } \text{Cr}_2\text{O}_3)\). The V/Cr ratio was very high, ranging from 3.7 to 24.3. The emeralds also contained remarkably low concentrations of Mg and Na \((\text{about } 0.1 \text{ wt.\% oxide or less})\). Concentrations of Rb \((21–61 \text{ ppm})\) and Cs \((35–127 \text{ ppm})\) were quite high compared to other vanadian emeralds. Iron concentrations were low \((467–1024 \text{ ppm})\) compared to emeralds from most other deposits.

**Spectroscopy.** UV-Vis-NIR. Typical polarized absorption spectra of a Byrud emerald (no. BVM2378-A) are presented in figure 14. The spectrum acquired with E||c shows V- \((\text{and partially Cr-})\) related absorption bands at 430 and 608 nm. With E\perp c\perp, the spectrum shows absorption bands at 423 and 630–642 nm (compare to Fritsch et al., 2002). A weak absorption band at 683 nm \((\text{spectrum E\perp c\perp})\) is ascribed to chromium. The weak shoulders at 373 and 385 nm are due to Fe\sup 3+.

In the near-infrared region, the three main absorption bands at approximately 1150, 1400, and 1896 nm are due to type I and type II water.
Infrared. Figure 15 shows two directional IR spectra in the 5800–1800 cm\(^{-1}\) range. Both H\(_2\)O and CO\(_2\) absorptions are clearly visible. Details of the water-related absorptions in the range 4000–3000 cm\(^{-1}\) are given in figure 16. These spectra show typical absorptions due to both type I H\(_2\)O (3700–3694 and 3607 cm\(^{-1}\)) and type II H\(_2\)O (3597–3595 cm\(^{-1}\)) molecules. The peak at 5273 cm\(^{-1}\) (\(\nu_2+\nu_3\)), attributed to type II water, is quite sharp and weak (figure 15), which means that there are relatively few type II water molecules in emeralds from Byrud.

Details of the absorptions in the 3000–2100 cm\(^{-1}\) range are given in figure 17. The CO\(_2\) absorption at 2359 cm\(^{-1}\) is sharp; weak absorptions at 2372, 2345, and 2326 cm\(^{-1}\) are also attributed to CO\(_2\). These are possibly due to isotopic effects (combination of the presence of \(^{13}\)C and \(^{18}\)O). The weak peak at 2818 cm\(^{-1}\) is attributed to chlorine (Schmetzer et al., 1997, and references therein; Fritsch et al., 1998). Several additional peaks shown in figure 17 (i.e., at 2928, 2854, 2739, 2687, and 2640 cm\(^{-1}\)) have never before been documented in the literature for natural emeralds. They may correspond to companion peaks of the 2818 cm\(^{-1}\) chlorine peak, even if their positions differ slightly from those observed in hydrothermal synthetic emeralds.

### TABLE 1. Chemical composition of two Byrud emeralds, obtained by electron microprobe.\(^a\)

<table>
<thead>
<tr>
<th>Oxide (wt.% )</th>
<th>Sample 203.18</th>
<th></th>
<th>Sample 195.174</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Range</td>
<td>Average</td>
<td>Std. dev.</td>
<td></td>
</tr>
<tr>
<td>SiO(_2)</td>
<td>65.49–66.73</td>
<td>65.91</td>
<td>0.33</td>
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<tr>
<td>Al(_2)O(_3)</td>
<td>17.02–18.78</td>
<td>17.93</td>
<td>0.43</td>
<td></td>
</tr>
<tr>
<td>BeO(^b)</td>
<td>13.69–13.92</td>
<td>13.77</td>
<td>0.06</td>
<td></td>
</tr>
<tr>
<td>V(_2)O(_3)</td>
<td>0.16–2.44</td>
<td>1.17</td>
<td>0.54</td>
<td></td>
</tr>
<tr>
<td>FeO</td>
<td>0.05–0.18</td>
<td>0.11</td>
<td>0.04</td>
<td></td>
</tr>
<tr>
<td>MgO</td>
<td>0.05–0.10</td>
<td>0.07</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>Na(_2)O</td>
<td>0.02–0.14</td>
<td>0.08</td>
<td>0.03</td>
<td></td>
</tr>
<tr>
<td>Sc(_2)O(_3)</td>
<td>nd–0.08</td>
<td>0.05</td>
<td>0.02</td>
<td></td>
</tr>
<tr>
<td>Cr(_2)O(_3)</td>
<td>nd–0.30</td>
<td>0.13</td>
<td>0.09</td>
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<tr>
<td>H(_2)O(^b)</td>
<td>0.85–0.96</td>
<td>0.91</td>
<td>0.02</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>99.52–100.87</td>
<td>100.13</td>
<td>0.38</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) Total iron is shown as FeO. The detection limit for Sc\(_2\)O\(_3\) is ~0.05 wt.%, and for Cr\(_2\)O\(_3\) is ~0.06 wt.%. Abbreviations: Std. dev. = standard deviation, nd = not detected.

\(^b\) BeO and H\(_2\)O were calculated, not measured.
DISCUSSION

Variations in Properties. The RI, SG, pleochroism, and fluorescence of our Byrud emeralds samples fall within the range of properties for emerald (Zylbermann, 1998). Higher values of RI were previously reported for Byrud emeralds ($n_o = 1.591$, $n_e = 1.584$; Webster, 1955).

Variations in color zonation, RI values, inclusion composition, and chemical composition may indicate that the mineralizing fluid was less homogeneous than is usual for emerald deposits. The relatively high concentration of chromophores (mostly V) may explain the high RI values sometimes observed, because—for example—V, which is heav-

### TABLE 2. Chemical composition of three Byrud emeralds, obtained by LA-ICP-MS.

<table>
<thead>
<tr>
<th>Element (ppm)</th>
<th>Sample 203.18</th>
<th>Sample 195.174</th>
<th>Sample 106.718</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Range</td>
<td>Average</td>
<td>Range</td>
</tr>
<tr>
<td>Li</td>
<td>20–28</td>
<td>24</td>
<td>15–21</td>
</tr>
<tr>
<td>Mg</td>
<td>277–474</td>
<td>389</td>
<td>306–486</td>
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<tr>
<td>K</td>
<td>23–92</td>
<td>55</td>
<td>26–684</td>
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<tr>
<td>Sc</td>
<td>126–332</td>
<td>266</td>
<td>160–278</td>
</tr>
<tr>
<td>Ti</td>
<td>23–270</td>
<td>146</td>
<td>12–16</td>
</tr>
<tr>
<td>V</td>
<td>4965–10021</td>
<td>7356</td>
<td>4496–6895</td>
</tr>
<tr>
<td>Cr</td>
<td>293–1208</td>
<td>579</td>
<td>1006–1654</td>
</tr>
<tr>
<td>Mn</td>
<td>5–25</td>
<td>5737</td>
<td>18–25</td>
</tr>
<tr>
<td>Fe</td>
<td>573–1024</td>
<td>802</td>
<td>467–698</td>
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<tr>
<td>Ni</td>
<td>nd</td>
<td>nd</td>
<td>nd</td>
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<tr>
<td>Zn</td>
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<td>5291</td>
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</tr>
<tr>
<td>Ga</td>
<td>9–27</td>
<td>5841</td>
<td>19–26</td>
</tr>
<tr>
<td>Rb</td>
<td>29–61</td>
<td>41</td>
<td>21–54</td>
</tr>
<tr>
<td>Sr</td>
<td>nd</td>
<td>nd</td>
<td>nd–7</td>
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<tr>
<td>Zr</td>
<td>nd–1.1</td>
<td>nd</td>
<td>nd</td>
</tr>
<tr>
<td>Cs</td>
<td>50–127</td>
<td>85</td>
<td>35–44</td>
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<td>Pb</td>
<td>1.8–2.8</td>
<td>2.2</td>
<td>3–5</td>
</tr>
<tr>
<td>V/Cr</td>
<td>6–24.3</td>
<td>16.2</td>
<td>3.68–4.47</td>
</tr>
<tr>
<td>V/(V+Cr)</td>
<td>0.9–1.1</td>
<td>0.9</td>
<td>0.79–0.82</td>
</tr>
<tr>
<td>Fe/(Fe+Cr)</td>
<td>0.4–0.8</td>
<td>0.6</td>
<td>0.24–0.41</td>
</tr>
</tbody>
</table>

*Co, Cu, La, Ce, Gd, Th, and U were analyzed for but not detected. Na, Cl, F, P, and Ca could not be analyzed correctly by this technique. Si is taken as the reference at 650,000 ppm. Abbreviations: DL = detection limit, nd = not detected.

Figure 14. Polarized UV-Vis-NIR spectra of sample BVM2378-A show absorption bands at 430 and 608 nm (bottom) and 423 and 630–642 nm (top) that are due mostly to V$^{3+}$, and partly to Cr$^{3+}$. The weak shoulders at 385 (top) and 373 nm (bottom) are due to small amounts of Fe$^{3+}$. The sharp near-infrared peaks at approximately 1150, 1400, and 1896 nm are due to type II molecular water. Spectra have been shifted vertically for clarity. Sample thickness is 3 mm.
ier than the Al it substitutes for, may contribute to slowing down light as it passes through the emerald. However, the RI values are sometimes very low, as seen in samples BVM2378-A to D, which were not chemically analyzed. Their low RI values may be related to the water and CO$_2$ content in the channels of these emeralds, which are very low compared to those of other natural emeralds (B. Sabot, pers. comm., 2006; a value of 1.1 wt.% H$_2$O is given by Alexandrov et al., 2001). As these samples were vivid green, the channel content apparently may be more important than the chromophores for controlling RI. Nevertheless, our limited data set does not allow us to demonstrate a clear correlation between RI and water, CO$_2$, and V contents.

Some solids in the multiphase inclusions were identified only with Raman spectroscopy (calcite) and others only with EDS (sylvite and pretulite), simply because we did not analyze the same specimens with both methods. This illustrates the diversity of the solid phases in the multiphase inclusions. However, opaque, metallic-appearing sulfides were observed in the fluid inclusions of all seven samples studied.

**Origin of Color.** The main UV-Vis spectroscopic features are the absorption bands at 423 or 430 nm and at 608 or 642 nm. These positions prove that the color is due mainly to the presence of V. However, the weak peak at 683 nm indicates that Cr also contributes to the coloration. This was confirmed by chemical analyses: V was much more abundant than Cr.

**Source of Vanadium and Chromium.** Chromium in emerald typically originates from mafic and ultramafic rocks (e.g., Morteani and Grundmann, 1977; Laurs et al., 1996; Marshall et al., 2003), with the exception of Colombian-type deposits, where chromium is leached from sedimentary rocks (Giuliani, 1997). However, the geologic source of vanadium in emerald is not well documented. This element is usually concentrated in rocks that contain iron-rich
minerals or organic compounds (e.g., Moskalyk and Alfantazi, 2003). Byrud’s alum shales, rich in organic matter, are the most likely source of vanadium for these emeralds. By comparison, the source of vanadium in Colombian deposits is also related to organic matter in black shales (Campos Alvarez and Roser, 2007).

Trace-Element Geochemistry. Trace-element incorporation into beryl varies with geologic environment (Staatz et al., 1965; Calligaro et al., 2000). We plotted some trace-element pairs to detect possible correlations in their relative abundances within the three samples analyzed. Chromium and V showed a positive correlation (figure 18) in two samples but not in sample 203.18. Gallium and V showed a clear negative correlation in all three samples (figure 19). A less pronounced negative correlation was found for Zn, Pb, and Mn relative to V (figure 20). From these data (based on the analysis of only three samples, as mentioned above), the group of divalent ions Zn, Mn, Pb, and Ga seems to correlate negatively with V, but we did not observe any correlation with Cr. All this confirms that Cr and V do not have similar geochemical distribution properties. Also, this could indicate that Ga and V integrate into the beryl structure via the same crystallographic site.

We calculated a very low water content in our samples (0.85–0.96 wt.%). This is not inconsistent with the value of 1.1 wt.% reported by Alexandrov et al. (2001). It approaches the lowest water content recorded for emerald from any source (Schwarz, 1987, p. 43). The low water content in such emeralds is related to the very low amount of Na, as type II water incorporation directly correlates to Na concentration (Wood and Nassau, 1968; Charoy, 1998).
IDENTIFICATION

We found several distinguishing criteria for Byrud emeralds. Most importantly, the presence of sulfides in the multiphase fluid inclusions is unique to emeralds from this locality and makes them straightforward to identify. With the optical microscope, we consistently observed a small (sometimes very small) black, opaque point somewhere in each multiphase inclusion. Three-phase and multiphase inclusions are known in emeralds from only a few other deposits: Colombia, Kaduna in Nigeria (Schwarz, 1998; Vapnik and Moroz, 2000; Sabot et al., 2000), Panjshir in Afghanistan (Seal, 1989; Bowersox et al., 1991), Kafubu in Zambia (Zwaan et al., 2005), and Xinjiang in China (Blauwet et al., 2005). Multiphase inclusions in these emeralds commonly contain halite cubes and sometimes carbonate crystals, but none of them systematically show the black, opaque phases seen in emeralds from Byrud. Sulfides have been described as solid inclusions in emeralds from several deposits (Giuliani et al., 1997; Rondeau et al., 2003), but never as part of a multiphase inclusion. Also, we observed that fluid inclusions in Norwegian emeralds are generally more regular in shape than those in Colombian emeralds, and the halite cube they contain is smaller. Hence, the careful use of a binocular microscope can be sufficient to ascribe Norwegian provenance to an emerald.

Among chemical criteria, the very low Na concentration (about 0.1 wt.% or less) is most significant. This is comparable to emeralds from Emmaville, Australia (Schwarz, 1998), and Delbegetey, Kazakhstan (Gravilenko et al., 2006), and is therefore not distinctive but certainly indicative. The appropriate data can be obtained by using either an electron microprobe or LA-ICP-MS.

In addition, the V content of the Byrud emeralds we tested was commonly high (a mean of more than 1 wt.% V2O3, with values ranging up to 2.4 wt.%), which is consistent with earlier data (Schwarz, 1991; Calligaro et al., 2000). This is among the highest of all emeralds, along with those from Malipo, China (Zhang and Lan, 1999). However, the V content is sometimes lower (down to 0.16 wt.% V2O3), particularly in light green stones, so this criterion cannot be considered a definitive identification tool. Moreover, a high V/Cr ratio (3 to 24 in our measurements) is not distinctive: Emeralds from other deposits (Salininha, Brazil, and Lened, Canada) also can show high V/Cr ratios, sometimes exceeding 100 (Marshall et al., 2004). However, the relative contents of Fe+Mg (low), Cr (low), and Cs or Rb (high) appear specific to the emeralds from Byrud, as shown in a ternary diagram that plots Fe+Mg, Cr, and Cs (figure 21). The trace-element composition of emeralds from Lened (Canada) has not yet been determined, but it may be useful to compare it to Byrud in this ternary diagram.

The oxygen isotopic composition of an emerald from Byrud has been reported in the literature (Giuliani et al., 1998; Groat et al., 2002). The δ18O value of 9.4‰ is rather low, but overlaps that of many other deposits (Giuliani et al., 1998; Sabot, 2002), so this criterion alone is not distinctive.

According to the published literature, the emeralds from Emmaville and Torrington, Australia, share...
many characteristics with emeralds from Byrud (Schwarz, 1991). They formed in pegmatite sills that intruded alum schist (Schwarz, 1991, and references therein; Grundmann and Giuliani, 2002), and their associated minerals include quartz, feldspars, micas, fluorite, topaz, cassiterite, wolframite, and arsenopyrite. These Australian emeralds contain very low concentrations of Na$_2$O and MgO (<0.1 wt.%; Schwarz and Henn, 1992; Brown, 1998). Such low concentrations are otherwise encountered only in emeralds from Byrud and in synthetic emeralds. Emerald crystals from Emmaville and Torrington typically show strong color zonation parallel to the basal face, which is rare in emeralds from Byrud. Also, emeralds from Emmaville contain more chromium and less vanadium (0.1 wt.% V$_2$O$_3$) than those from Byrud (Schwarz, 1991).

CONCLUSION

The gemological and spectroscopic properties of our Byrud samples did not differ on average from those of emeralds in general, either chromium- or vanadium-bearing. However, our study has shown that

Byrud emeralds (e.g., figure 22) are colored mostly by vanadium. We also found a number of criteria that are diagnostic of these emeralds. Most importantly, we noted the presence of sulfide phases in multiphase fluid inclusions, together with halite, sylvite, calcite, liquid water, and gaseous CH$_4$. This is the first time that sulfide phases have been reported in multiphase inclusions in emeralds. They are seen with the optical microscope as minute black, opaque points. The chemical composition is also distinctive: The emeralds analyzed from Byrud had relative contents of Fe, Mg, Cr, Cs, and Rb that are specific to this locality and easily identified through LA-ICP-MS analysis. They also contained low Na and Mg and often high V.

These criteria make emeralds from Byrud unique and quite easy to distinguish using a microscope and/or chemical analysis. An unverified legend says that one of the emeralds in the jewels belonging to the British royal family comes from the Byrud deposit. Using the identification criteria provided in this article, it would now be quite easy to investigate this story.

When we compare our results with those reported in the literature for emeralds from other localities, it appears that those emeralds from Byrud that show a strong color zonation closely resemble emeralds from
Emmaville and Torrington, Australia: similar color zonation, very low Na and Mg contents, similar geo-
logic setting, and similar associated minerals. Hence, a detailed gemological study of emeralds from
Australia would be very helpful in testing the identi-
fication criteria proposed here for the emeralds from
Byrud. In general, much remains to be done on vana-
dium-bearing emeralds (except perhaps for some
Colombian deposits), which are much less studied
than their chromium-bearing counterparts.

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