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The Journal of Gemmology

VOLUME 25
NUMBER 2  APRIL 1996

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Cover Picture
Faceted sapphire from Cyangugu district, SW Rwanda. See 'Investigations on sapphires from an alkali basalt, South West Rwanda' on pp 90-106.
In this issue...

We introduce this issue with an update of a collection started many years ago and first described in this Journal in 1987. Its original motive was to enlighten the student of gemmology with the number of stones that could be mistaken for diamond. But with increasing knowledge it led to appreciation on other levels with the fascinating range of gem materials that could appear in colourless form or with the subtleties of recognizing small variations in this 'colourlessness'. There are now 46 stones in the collection and they are listed with some basic gem parameters.

Rwanda has been in the news for all the wrong reasons in the past year so it is good to report a development of some potential for the local people. Sapphires have been discovered in the Nyando district in south-west Rwanda (bordering Zaire) and details of their source and gem characteristics are described. Most crystals are a deep blue but some show the milkiness characteristic of the Sri Lankan 'geuda' and these respond well to heat treatment. They contain a wide range of inclusions and a new texture found on some rough crystals, not previously reported, is described and discussed.

Ever since they were first discovered in 1917 the deep blue beryls from the Maxixe mine in Brazil have been a source of fascination both for those in the gem trade and for investigative gemmologists. Maxixe beryls, as they came to be known, are quite different from the commoner blue beryl, aquamarine, in that their colour has a different source within the crystal structure and it fades when exposed to sunlight. The third paper in this issue sets out some current thinking about Maxixe beryls and proposes suitable terminology and its limits of application in the beryl family.

The final paper presents the results of a thorough investigation (involving more than 1000 samples) of a range of emeralds and green beryls from Central Nigeria. Detailed work on inclusions, compositions of hosts and inclusions and on spectra are illustrated, and provide a reference to enable confident identification of emeralds from this locality. Some Nigerian emeralds do contain spiky three-phase inclusions and resemble Colombian emeralds, but again the means for distinguishing between stones from the two localities are indicated.

R.R.H.
Common and rare colourless gemstones

David Kent FGA
London

Over the years I have compiled a collection of colourless gemstones which have been a source of fascination especially for students embarking on their gemmological studies. To a greater or lesser extent the stones may all be considered as diamond simulants and in 1987 a list of 39 stones was published (J. Gemm., 20(6), 344-5). Following some recent acquisitions (and one or two delightful presents!) I feel it is appropriate to bring the record of the collection up to date.

A new style Ever Ready P6 9-volt battery was fitted to the infrared reflectometer and all the specimens were measured again. This led to lower readings (compared with those in the 1987 table) for lithium niobate and smithsonite and a higher reading for the diamond/synthetic sapphire doublet. The higher meter reading for paste results from replacing the original stone with a much more lively lead-glass paste. Is the higher reading for the diamond/synthetic sapphire doublet due to deterioration of the adhesive?

Benitoite and beryllonite in the original list were on loan and have been returned to their owners, and the new additions comprise natrolite, oligoclase, sillimanite, kornerupine, spodumene, chrysoberyl, anglesite and scheelite. There are now 46 stones in the collection and the relevant data are given in Table I.

Table I: Revised list of the David Kent colourless stone collection with additions since 1987. Infrared reflectometer readings calibrated at 70 for diamond. In order of meter readings.

<table>
<thead>
<tr>
<th>Meter reading</th>
<th>Mean RI</th>
<th>Specimen</th>
<th>Carat weight</th>
<th>Hydrostatic SG</th>
<th>Meter reading</th>
<th>Mean RI</th>
<th>Specimen</th>
<th>Carat weight</th>
<th>Hydrostatic SG</th>
</tr>
</thead>
<tbody>
<tr>
<td>17</td>
<td>1.434</td>
<td>Fluorite</td>
<td>2.91</td>
<td>3.18</td>
<td>24</td>
<td>1.613</td>
<td>Paste</td>
<td>1.63</td>
<td>3.54</td>
</tr>
<tr>
<td>17</td>
<td>1.440</td>
<td>Opal</td>
<td>1.96</td>
<td>2.00</td>
<td>25</td>
<td>1.615</td>
<td>Topaz</td>
<td>3.95</td>
<td>3.56</td>
</tr>
<tr>
<td>17</td>
<td>1.487</td>
<td>Natrolite</td>
<td>0.21</td>
<td></td>
<td>25</td>
<td>1.622</td>
<td>Amblygonite</td>
<td>2.58</td>
<td>3.03</td>
</tr>
<tr>
<td>20</td>
<td>1.506</td>
<td>Petalite</td>
<td>2.06</td>
<td>2.39</td>
<td>25</td>
<td>1.629</td>
<td>Tourmaline</td>
<td>0.60</td>
<td>3.00</td>
</tr>
<tr>
<td>20</td>
<td>1.510</td>
<td>Leucite</td>
<td>0.85</td>
<td>2.50</td>
<td>25</td>
<td>1.640</td>
<td>Baryte</td>
<td>8.00</td>
<td>4.49</td>
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<tr>
<td>21</td>
<td>1.539</td>
<td>Oligoclase</td>
<td>0.79</td>
<td>2.63</td>
<td>25</td>
<td>1.647</td>
<td>Datolite</td>
<td>0.96</td>
<td>2.90</td>
</tr>
<tr>
<td>22</td>
<td>1.549</td>
<td>Quartz</td>
<td>2.69</td>
<td>2.65</td>
<td>26</td>
<td>1.633</td>
<td>Danburite</td>
<td>1.59</td>
<td>2.98</td>
</tr>
<tr>
<td>22</td>
<td>1.549</td>
<td>Synthetic</td>
<td>quartz</td>
<td>4.67</td>
<td>26</td>
<td>1.675</td>
<td>Smithsonite</td>
<td>0.44</td>
<td>4.08</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(approx.)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>23</td>
<td>1.540</td>
<td>Iolite</td>
<td>1.19</td>
<td>2.58</td>
<td>27</td>
<td>1.657</td>
<td>Jadeite</td>
<td>1.77</td>
<td>3.34</td>
</tr>
<tr>
<td>23</td>
<td>1.560</td>
<td>Scapolite</td>
<td>0.39</td>
<td>2.60</td>
<td>27</td>
<td>1.657</td>
<td>Enstatite</td>
<td>1.03</td>
<td>3.22</td>
</tr>
<tr>
<td>23</td>
<td>1.561</td>
<td>Labradorite</td>
<td>1.25</td>
<td>2.68</td>
<td>27</td>
<td>1.661</td>
<td>Euclase</td>
<td>0.43</td>
<td>3.08</td>
</tr>
<tr>
<td>23</td>
<td>1.583</td>
<td>Beryl</td>
<td>1.48</td>
<td>2.74</td>
<td>28</td>
<td>1.661</td>
<td>Phenakite</td>
<td>0.85</td>
<td>2.98</td>
</tr>
</tbody>
</table>

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Fig. 1. The author's boxed collection of colourless stones.
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Investigations on sapphires from an alkali basalt, South West Rwanda

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3. SEM-Laboratory, University Basel, Switzerland

Abstract

A new deposit of sapphires in the Cyangugu district of SW Rwanda has been investigated. The sapphires are believed to have originated from a specific alkali basalt lava flow, extruded during the Tertiary extensional regime along the East African Rift. They exhibit mainly a deep blue colour, often showing so-called silk (inclusions of hematite or rutile) or a slight milkiness possibly due to submicroscopic exsolution of these minerals. Greyish 'geuda'-type crystals are known to convert to a blue colour by heat treatment. Values for the refractive indices and specific gravity are given, as well as VIS-IR data. Studies of the inclusions reveal the presence of Ti and Fe oxides, silicates, spinel, zircon and a complex Th-REE phosphate as solid inclusions; the presence of CO2 in fluid inclusions was determined by microthermometric methods. Studies of crystal surfaces by SEM show corrosion or abrasion features, both primary and secondary. The primary corrosion occurred during the transport of the deep-crustal sapphires to the surface; it can be attributed to the high-temperature magma. Secondary abrasion took place during the erosion and weathering of the basalt and the subsequent formation of alluvial sapphire deposits.

Keywords: sapphire, alkali basalt, inclusions, corrosion, scanning electron microscopy

Introduction

East-Central Africa is well known for rich gemstone deposits. Extensive sapphire mining has been described from the Umbo river (Tanzania), Garba Tula and Turkana (Kenya) (Hughes, 1990; Themelis, 1989; Hänni, 1986; Kanis and Harding, 1990; Barot and Harding, 1994).

Rwanda, at present in the political focus because of the tragic civil war, was until now barely known for gemstone deposits. One of the authors (MSK) recently had the opportunity to investigate a new sapphire deposit in a basalt province in the Cyangugu district, SW Rwanda (Figure 1). This investigation was made possible by kind support of J.F. Damon (Twin Gems, Washington DC).

After a geological field trip of two months which included trenching and sampling in a region of approximately 200km², some areas of major interest for sapphire mining were established (Figure 2). Due to the rapid decomposition of rocks in a tropical humid climate, the main sapphire deposits are of alluvial type. They are exploited mostly by the local population using traditional methods (Figure 3). Twenty representative sapphire samples from the Cyangugu district were selected for the laboratory and investigated by various methods, including spectrophotometry...
(VIS-IR), SEM-EDX, ED-XFA, EMP-WDX, microthermometry and optical microscopy (OM) (Table I). During these analytical procedures information was gained about the trace elements responsible for colour, solid as well as fluid inclusions and surface features of sapphire crystals. In particular, a specific surface feature, which resulted from a shielding effect of plagioclase on sapphire and which influenced the primary corrosion in the basalt in a very spectacular way was observed.
Geological setting

The western border of Rwanda follows the western branch of the East African rift (Figure 4), a continental rift zone, which has been tectonically active for the last 45 million years, i.e. since the early Tertiary. It forms a part of the Afro-Arabian rift system, which extends some 6500 km from Turkey to Mozambique. As a result of the extensional tectonics along this rift, the Red Sea opened in the northern part, whereas in the south, rifting occurred to a much lesser extent (Figure 1).

The western branch of the East African Rift has developed in a rather episodic way, accompanied by intense extensional faulting of the Precambrian basement. The basement is characterized by continental sedimentary rocks, i.e. sandstones and arkoses lying on a metamorphic substratum of gneisses and amphibolites. Upwelling of asthenosphere (upper mantle) caused a regional domal uplift.
Table I: Gemmological properties of sapphires from Cyangugu district, SW Rwanda

<table>
<thead>
<tr>
<th>Property</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colour and visual appearance with unaided eye</td>
<td>dark blue, transparent to semitransparent with microscopic and submicroscopic inclusions ('silk')</td>
</tr>
<tr>
<td>Refractive index:</td>
<td>$n_o 1.770-1.771$</td>
</tr>
<tr>
<td></td>
<td>$n_e 1.761-1.762$</td>
</tr>
<tr>
<td>Birefringence:</td>
<td>$\Delta n = 0.009$</td>
</tr>
<tr>
<td>Specific gravity:</td>
<td>3.998–4.005</td>
</tr>
<tr>
<td>Dichroism:</td>
<td>greenish-blue and dark blue</td>
</tr>
<tr>
<td>Absorption spectrum:</td>
<td>absorption features at 375nm, 390nm, 450nm and a broad absorption band at 890nm</td>
</tr>
<tr>
<td>Reaction to long wave and short wave ultraviolet radiation:</td>
<td>no fluorescence</td>
</tr>
</tbody>
</table>

Fig. 4. The Ruizizi river in the western branch of the East African rift valley, SW Rwanda.
combined with intracontinental basaltic volcanism, which is typically dominated by alkali basalts (Cahen et al., 1984; Wilson, 1991) (Figure 6).

The sapphire deposits in SW Rwanda are connected to this Tertiary-Quaternary volcanic activity. The sapphires have been brought to the surface by an alkali-basaltic magma, in which they were transported as xenoliths (alien crystals, separated in the upper mantle or deep crust from their original melt or rock). Alkali basaltic rocks as parent rocks of corundum are encountered in many places worldwide, and economically important deposits are exploited, e.g. in Australia, Thailand, Nigeria and China (Coenraads, 1992a; Guo et al., 1992).

Due to the intense decomposition of volcanic rocks in the humid climate, sapphires have not been found in their host rock (embedded in alkali basalt), but by carefully examining the eluvial and alluvial gravel accumulations, the sapphire-bearing primary deposit could be assigned to a specific Tertiary alkali basalt. This coarse-grained, olivine-rich basalt flow always occurs near the contact between Tertiary basalt and the Precambrian basement. Therefore it is suggested that this coarse-grained alkali basalt had been erupted during an early stage of the Tertiary-Quaternary volcanic activity in south-western Rwanda.

The sapphires, concentrated in these eluvial and alluvial deposits are found together with zircon (red-brown and colourless transparent), magnetite, hematite, rutile, pyrite and quartz (originating from the Precambrian rocks).

The Cyangugu sapphires

Mineralogy

Sapphire, basically the blue variety of corundum $\text{Al}_2\text{O}_3$, occurs generally in high alkaline and silica-undersaturated rocks. Two types of geological settings are possible (Levinson and Cook, 1994): the sapphires originate from metamorphic rocks (e.g. dolomite marbles, gneisses) or from magmatic rocks (alkali basalts and pegmatites). Trace elements such as the transition metals Fe, Ti, Cr, V are known to cause a broad variety of colours in sapphires. Either alone or in combination, these ionic impurities may act as chromophores when substituting for $\text{Al}^{3+}$ in the
Fig. 8. Slender, prismatic, uncorroded sapphire from the Cyangugu district.

Fig. 9. Faceted and cabochon sapphires from the Cyangugu district. Largest stone is approx. 8 ct.
crystal structure. It has also been observed that these trace elements can actually quit their places in the corundum lattice and concentrate in individual inclusion minerals (Schmetzer, 1986) such as rutile (TiO₂), hematite (Fe₂O₃) or ilmenite (FeTiO₃). As many natural sapphires exhibit growth and colour inhomogeneities, experimental research concentrates on the improvement of the colour in lower quality sapphires by means of heat treatment (Themelis, 1992). During such treatment, former hematite or rutile inclusions may be re-dissolved and their Fe and Ti ions diffuse into the crystal lattice structure where they then may play the role of chromophore elements.

Morphology of rough sapphires

The Cyangugu sapphires are generally 1–8ct in weight and their habit is mostly prismatic-pyramidal. Rhombohedral crystals are much rarer. These observations are in agreement with those made by Kiefert (1987) on sapphire from other basaltic origins (e.g. Nigeria and Australia). She described as significant morphological faces, a, r, n and z, whose combination led to tabular, barrel-shaped prismatic and pyramidal crystals (Figure 7).

The surface morphology of the sapphires from the Cyangugu district is in general strongly affected by primary and secondary corrosion or abrasion effects and fractures. Normally the sapphires are broken into flat fragments along the basal plane (0001). Only a few small individuals have preserved their long prismatic shape (Figure 8).

Spectroscopy and chemical analyses

The sapphires from Cyangugu district (Figures 5 and 9) are mainly deep blue but greyish and yellowish tints are also found. The latter are suitable for colour modification by heat treatment (Themelis, 1992).

In sapphires from Rwanda, the blue colour is mainly caused by the Fe²⁺/Fe³⁺ intervalence charge transfer (IVCT) (cf. Karr, 1975; Schmetzer and Bank, 1981; Schmetzer, 1987). A typical UV-visible-near IR-spectrum reveals three distinct peaks at 375nm, 390nm and 450nm and a broad absorption peak centred at 890nm, which correspond to Fe³⁺, Fe²⁺/Fe³⁺ ion-pairs and Fe²⁺/Fe³⁺-IVCT absorption (Figure 10, compare also Fritsch and Mercer, 1993).

In one specimen an absorption shoulder at 560nm (o-vibration spectrum) was observed, caused by the superposition of Fe²⁺/Ti⁴⁺-IVCT on the Fe²⁺/Fe³⁺-IVCT absorption band (Figure 11). But as most spectra reveal no distinct peak at this wavelength (560nm), normally the Fe²⁺/Ti⁴⁺-IVCT can be neglected as the cause of colour in these sapphires. The spectral absorption feature at 780nm as observed with o-vibration spectra in Figure 11 remains unexplained (see Schmetzer, 1987).

The strongly dichroic colour of sapphires from Cyangugu district is well illustrated by the different absorption curves in Figure 11.

Qualitative (ED-XFA) and quantitative chemical analyses by the electron microprobe (EMP-WDX) on the same sapphire samples confirmed the spectroscopic information about their trace element content (Table II). Fe is present at constant trace element levels (~0.5 wt% Fe₂O₃), whereas the distribution of Ga varies (~0.03 wt% Ga₂O₃). Ti (and V, Cr, Ca) is nearly at or below detection limit (~0.015 wt% TiO₂) and therefore the blue colour cannot solely be attributed to the Ti. Detection limits below 0.015 wt%, as seen in the literature (Schmetzer and Banks, 1981; Zwaan, 1974), seem quite astonishing in the present context, unless more sensitive analytical methods were used.

Microscopy

Rwanda sapphire samples often exhibit a so-called silk (crystallographically oriented inclusions of submicroscopic dimensions). According to various authors (compare Schmetzer, 1986) rutile needles are oriented parallel to the first order prism of the host corundum. The orientation of
Fig. 10. Wavelength absorption spectra of a sapphire from the Cyangugu district (e-vibration spectrum).

Fig. 11. Wavelength absorption spectra of a sapphire from the Cyangugu district (o- and e-vibration spectra).
Table II: Quantitative chemical analyses (EMP-WDX) of two sapphires from SW Rwanda

<table>
<thead>
<tr>
<th></th>
<th>Mean of 3 analyses</th>
<th>Mean of 5 analyses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al₂O₃</td>
<td>99.31</td>
<td>99.53</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>0.55</td>
<td>0.46</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>n.d.</td>
<td>0.01</td>
</tr>
<tr>
<td>V₂O₃</td>
<td>n.d.</td>
<td>0.01</td>
</tr>
<tr>
<td>Ga₂O₃</td>
<td>n.d.</td>
<td>0.04</td>
</tr>
<tr>
<td>CaO</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>Total</td>
<td>99.87</td>
<td>100.06</td>
</tr>
</tbody>
</table>

n.d.: not detectable (below detection limit)

Hematite tablets may be parallel to the first order prism, or to the basal plane. By heat treatment, this silk occasionally can be removed, and stones of better quality are obtained. The silk normally occurs in growth zones, thus reflecting a varying supply of Ti and Fe during sapphire formation. Growth inhomogeneities can usually be recognized using dark-field optical microscopic techniques.

**Inclusion studies**

Identification of inclusions in gemstones is of considerable importance, as the inclusions yield information about the genesis of the gemstones and therefore may be— with some restrictions— essential for determining authenticity or even the origin of a gemstone (Hänni, 1994).

A number of solid inclusions, randomly encountered in the sapphires, have been investigated by the SEM-EDX technique and also using back-scattered electron images (BSE); these have been compared with the results from inclusion studies in sapphires from other alkaline basaltic deposits (Gübelin and Koivula, 1986; Coenraads, 1992b; Guo et al., 1992; Wang, 1988). Microthermometric studies revealed the presence of just one type of fluid inclusion which is CO₂-dominant.

**Ti and Fe oxides: rutile, ilmenite and hematite**

These inclusions are very common, typically with irregular shapes and dimensions of 0.1 to 2 mm. They are not noticeably oriented along the trigonal crystal structure of the sapphire and generally consist of aggregates of at least two of the three oxides. Partial replacement of rutile (TiO₂) by ilmenite (FeTiO₃) at the grain rims can be observed (Figure 12).
Submicroscopic Ti and Fe oxides
As mentioned above, many sapphires from the Cyangugu district show distinct growth zones which contain varying amounts of submicroscopic inclusions — the cause of the translucent (silky) appearance of sapphires. Tiny hematite and rutile crystals may be responsible for this effect, which has been described by many authors from sapphire deposits throughout the world (Hughes, 1990; Gübelin and Koivula, 1986). In the Cyangugu sapphires hematite occurs as tiny brownish inclusions with a 'flaky' (ragged) shape, and rutile occurs as (sub)microscopic, white needle-shaped inclusions (Figure 13).

Genetically, the finely dispersed submicroscopic inclusions may be related to the microscopic Fe- and Ti-oxides. Both reflect the varying Ti and Fe supply during the growth of the sapphires (Figure 14). The relatively large Ti-Fe-oxide inclusions represent protogenetic crystals, enclosed by the growing sapphire. During the following cooling stage, they probably underwent an isochemical replacement reaction where ilmenite was formed from rutile + hematite. In contrast, the submicroscopic Fe-Ti-inclusions are probably a product of the cooling of the sapphires, during which the Fe and Ti ions, formerly dissolved in the corundum lattice, began to precipitate, forming tiny hematite and rutile inclusions, with orientation controlled by the crystal structure of the corundum.

Silicates: pyroxene, amphibole and chlorite
The shape of these inclusions normally is subidiomorphic. They are partially surrounded by a rim of chlorite, and their size normally does not exceed 0.1mm.

The Ca-Fe-pyroxenes and -amphiboles revealed similar EDX-spectra. They were distinguished, using their typical crystal shape in a thin section cut perpendicular to the c-axis of an inclusion of amphibole or pyroxene. The amphibole inclusions may represent retrograde products of primary pyroxene inclusions. During cooling, the pyroxene could have transformed to amphibole with the supply of H\textsubscript{2}O from small scale fissures or diffusion processes within the sapphire. Such (healed) small scale fissures have been observed by OM and SEM-techniques. Mica inclusions are also present but are so fragile that they tend to disintegrate during the polishing of the sapphire samples.

Similar kinds of inclusions have been described from other alkali basalt derived sapphire deposits, e.g. in the Australian New England deposit (Coenraads, 1992b), from the Sri Lanka sapphires (Gübelin and Koivula, 1986) and in sapphires from the Shandong Province in China (Guo et al., 1992).
Spinel

In one sapphire sample a subidiomorphic solid inclusion, surrounded by a chlorite reaction rim was analysed by EDX-spectra and revealed a Mn-bearing intermediate member of the spinel solid solution series between hercynite ($\text{Fe}^{2+}\text{Al}_2\text{O}_4$) and gahnite ($\text{ZnAl}_2\text{O}_4$) (Figure 15).

Zircon and cheralite-(Ce)

Zircon and cheralite inclusions in the sapphires may be detected by back-scattered electron image investigations (SEM-BSE). Because they contain elements with high atomic weight (e.g. Zr, Th, REE...), they show a strong brightness contrast relative to the surrounding sapphire (Figure 16).
The zircon inclusions (0.02 to 0.1mm) are characterized by a more or less idiomorphic shape, suggesting proto to syngenetic growth relative to the sapphire. They commonly occur in small groups and are surrounded by stress fissures in the host sapphire, typical for metamict inclusions (Gübelin and Koivula, 1986) (Figure 17).

One inclusion, adjacent to some zircon inclusions, was identified by SEM-EDX as cheralite-(Ce), a complex thorium-phosphate \((\text{Ca,Ce,Th})(\text{P, Si})\text{O}_4\) belonging to the monazite group (Figures 18 and 19).

Stress fissures around this inclusion are partially filled with a mineral (probably goethite \(\alpha\)-\(\text{FeO(OH)}\)), causing a pale brown halo (Figure 20).

**Fluid inclusions**

Microthermometric studies (by B. Bruder) of the sapphires revealed the presence of \(\text{CO}_2\) dominated fluid inclusions as fillings of voids, negative crystals and 'fingerprints'. At room temperature, most of the fluid inclusions are monophase (pure \(\text{CO}_2\) with a maximum density of 0.95g/cm\(^3\)). This is consistent with results from fluid inclusion studies on other volcanic sapphires. No distinctive fluid composition
and density was discovered for the Rwanda sapphires (Figure 21).

Inclusions similar to those observed in sapphires from the Cyangugu district have been described by many authors from analogous sapphire deposits in volcanic provinces. As Webster and Anderson (1983) and Hanni (1994) suggest, using the inclusions alone is insufficient to indicate the geographic origin of a gemstone. They merely give information about the geological and geochemical conditions during their genesis (for the Rwanda sapphires: during the continental volcanic activity along the western part of the East African rift suture).

Surface features
There are two main processes that have affected the surfaces of the Rwanda sapphires. Primary corrosion, a mainly chemical process during volcanic activity, and secondary abrasion and fracturing, mechanical impact processes which occurred during the erosion of basalts and subsequent transport of the sapphires to the alluvial deposit.

The primary corrosion features were studied with the SEM on eight randomly selected samples.

The genesis of sapphires from alkali basalts is still under debate. The model generally accepted is that they were formed at the conditions prevailing in the lower crust or upper mantle (high temperature and pressure) in a high aluminium environment. Most authors (Coenraads, 1992b; Guo et al., 1992; Levinson and Cook, 1994) suggest, however, that the sapphires were not formed in the alkali basalt itself. The corrosion features on the surfaces of sapphires are interpreted as reactions of sapphire xenocrysts in disequilibrium with the basic magma (alkali basalt). If correct, this model implies that the basaltic magma acted merely as a transport medium and not as the parent magma of the sapphires. During the rise of the carrier magma, the sapphires were corroded by reactions with the melt.

Most Cyangugu sapphires show distinct primary corrosion features. The surface is typically rough and marked by many indentations (Figure 22). On the basal plane, there are triangular indentations, which are typical corrosion features of trigonal minerals.

One specimen is covered with a thin brownish layer (Figure 23), probably consisting of a Fe-hydroxide ($\alpha$-FeO(OH)), which may have been derived originally from native iron.

Another specimen shows a very specific corrosion, not previously recorded. The whole surface is covered by a randomly oriented needle-like pattern (Figures 24 and 25), interpreted as due to long prismatic plagioclases, which covered the sapphire during its transport by and in the plagioclase-rich alkali basalt. The plagioclase partly protected the sapphire from primary corrosion in the magma (Figure 26) but due to the intense weathering of the basalt, it has since decomposed, and only the sapphire remains (cf. Coenraads, 1992a).

This specific corrosion feature sheds some light onto the relation between sapphire genesis and alkali basaltic magmatism. Plagioclases that are probably responsible for the needle-like surface pattern are virtually absent as solid inclusions within the sapphires from Cyangugu dis-
Fig. 22. SEM-micrograph of a strongly corroded sapphire from the Cyangugu district. Therefore it is suggested that these sapphires originate from a source other than the alkali basalt magma. Whether the incorporation of sapphires into the ascending alkali basalt occurred by magma mixing processes (magmatic origin of sapphires) or by supply of deep crustal rock fragments (metamorphic origin of sapphires) is not yet established (cf. Levinson and Cook, 1994). Radiometric age dating of both the alkali basalt and the sapphires (U-Pb-dating of zircon inclusions) could reveal further information on sapphire genesis (cf. Coenraads et al., 1990).

Fig. 23. SEM-micrograph of a sapphire, partially protected by a Fe-hydroxide layer.

Conclusions
Sapphires from the Cyangugu district (SW Rwanda) originate from a specific alkali basalt, which has been erupted during Tertiary volcanic activity along the East African rift. Prospecting has established several alluvial-type sapphire deposits, which may have potential for future mining.

The sapphires normally are deep blue and commonly show a so-called ‘silky’ appearance. Three different types of solid inclusions which are typical for corundum
Fig. 24. SEM-micrograph of a specific needle-like corrosion feature on a sapphire of the Cyangugu district.

Fig. 25. SEM-micrograph of the marked area of the same sample under higher magnification.
Fig. 26. Scenario explaining the genesis of the corrosion feature shown in Figures 24 and 25:
(a) the sapphire was covered by plagioclases in the ascending alkali basalt, protecting the gem stone partly from corrosion;
(b) subsequently, the plagioclases have been weathered during erosion of the erupted basalts.

from volcanic provinces have been distinguished. The sapphires often show signs of pronounced corrosion, and a needle-like pattern is interpreted as the relic of a former coating of plagioclase crystals. There is close similarity of these sapphires with those derived from other basaltic deposits, and an unambiguous identification of sapphires specifically from the Rwanda deposit is not possible.

Until now, sapphires from the Cyangugu district in SW Rwanda have not been well known on the gem market, but if the political situation stabilizes they may well become much more familiar to stone dealers and gemmologists.

Acknowledgements
We would like to thank Dr R Gieré (Min. Petrogr. Institute, University Basel) for his critical review of this paper, and Dr L. Kiefert (SSEF, Basel) for analytical assistance in taking both optical and ED-XRF-spectra and for useful discussions. Thanks also go to J.F. Damon (Twin Gems Co., Washington DC) for financial support and to J. Stähli (Bern) for the loan of faceted sapphires from SW Rwanda and for his encouragement.

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Appendix

Analytical conditions

References for some of the analytical techniques applied are given in Hänni (1994). Two sapphire samples were cut and two faces polished parallel to the optic axis in order to record absorption spectra. With a Hitachi 4001 spectrophotometer the absorption features between 250 and 900nm (VIS-IR) were measured. Polarization filters were used in the visible part.

The scanning electron microscope, attached to an energy-dispersive X-ray spectrometer (SEM-EDX) is a powerful tool to obtain morphological and chemical information (Putnis, 1992). All the SEM investigations were performed on a Philips 515 instrument (20kV). Thin sections with randomly oriented, polished planes were prepared from ten samples to determine the nature of solid inclusions. The samples were coated with carbon for SEM-EDX analyses and back scattered electron imaging (BSE). Eight rough samples were sputtered with gold (20nm thickness) for SEM investigations of surface features.

Qualitative X-ray fluorescence analyses (ED-XRF) were carried out by a Philips combined Spectrace X-ray fluorescence spectrometer (25 kV, 0.3 mA, Al primary filter) in order to determine the trace elements present in the sapphires from SW Rwanda. For quantitative chemical information, two sapphires were analysed by a wavelength-dispersive electron micro-probe (1:200, JXA-8600) operated at 20 kV and with a specimen current of 10 nA measured on a Faraday cage (EMP-WDX). Seven reference standards (Al, Fe, Ti, Cr, V, Ga, Ca) of well characterized natural and synthetic compounds were used and all the data were fully corrected for matrix effects by PROZA-computer program.

Two thin slices (approximately 0.3mm thick) of sapphires were investigated under a Leitz Ortholux microscope equipped with a Reynolds heating-freezing stage for microthermometry. The measurements were performed on 75 fluid inclusions by Bernard Bruder (Univ. Freiburg, Germany) who investigated fluid inclusions in corundum from various sources for his university diploma in mineralogy.
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On the identification and fade testing of Maxixe beryl, golden beryl and green aquamarine

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Abstract
Forty-three large (up to 126.66 ct) high quality faceted beryls of recent Brazilian origin were examined. Of these, 29 were intense blue Maxixe-type beryls, a material widely seen about 1970. The remaining five yellow and nine greenish beryls were found to be simply golden beryls and green aquamarines, respectively.

Heat and light exposure tests showed that an accelerated fading test must not use an excessively elevated temperature, say 50 °C (122 °F) maximum, since otherwise both aquamarines and golden beryls (which do not fade from light alone) would lose their yellow colour component from the heat and could then be misidentified as fading Maxixe beryls. This has indeed happened in one report in the recent gemmological literature, where a temperature of 100 °C was used as part of a fade test.

It is proposed that the gemmological distinction between the original 'Maxixe' and the later 'Maxixe-type' beryls be dropped, the term 'Maxixe' then covering both variants. While the distinction remains significant in mineralogy and other fields, it serves no useful purpose in gemmology.

Introduction
The early history and the gemmological characteristics of the original Maxixe beryls of 1917 and the later Maxixe-type beryls are well covered in the gemmological literature. This includes their identification and distinction from aquamarine by the anomalous dichroism, unusual absorption spectra, and fading from exposure to light. These two variants differ somewhat in their absorption spectra and by the fact that the intense blue colour centre is associated with a nitrate impurity in the original Maxixe beryl, or with a carbonate impurity in the later Maxixe-type beryl.

It is often assumed that any intense blue original Maxixe beryl owes its colour to nature and that the equivalent colour of Maxixe-type beryl originates from man-induced irradiation as, for example, stated by Brown. Yet it is clear from the literature reports that the colour of the original Maxixe beryl can also be produced or restored by irradiation by man and that the Maxixe-type blue beryl also occurs naturally.

The distinction between these two variants is thus independent of the origin of the irradiation and depends only on the chemistry. Whether the colour derives from irradiation by nature or by man generally cannot be distinguished by gemmological or by any other form of testing. An exception occurs if an inappropriate irradiation technique has induced radioactivity, the residue of which may then be detectable, as happened at least once.

Although I myself originally introduced these two variant designations, I now believe that the distinction between the original 'Maxixe' beryl and the later
'Maxixe-type' beryl serves no useful purpose in gemmology. It has never been used in GIA identification reports, for example. I believe it should be dropped because:

1. both variants generally occur in intense blue form in nature;
2. both can be produced by irradiation of pale colour beryl containing the necessary precursor impurities (nitrate or carbonate);
3. both fade in light in about the same time frame, although there is some variability;
4. both lose their colour on being heated, again with some variability;
5. lost colour can be restored by irradiation for both; and
6. routine gemmological testing using a hand spectroscope is not sufficient to distinguish these two variants, but this distinction is possible by using a research spectroscope.

In what follows, the designation 'Maxixe' beryl includes both variants except when specifically designated as 'original Maxixe' beryl or 'Maxixe-type' beryl.

On loss of colour by light and by heat

Experts prefer to reserve the term 'fading' for a loss of colour on exposure to light and use other terms such as 'destruction' or 'loss' of colour for the equivalent action of heat. A typical example is Herbert Smith, who in his Gemstones describes the actions of light and heat on gemstones in separate paragraphs and uses the term 'fade' for the effects of light but not for those of heat. Again, Hunter and Harold define: 'Fading: a colour change in a material that involves a weakening or lightening with time, usually as a result of exposure to light or weather.'

Rapid testing for fading uses illumination conditions much more intense than those to which a gemstone would be exposed in normal jewellery wear. In studies begun in 1973, I found that Maxixe beryls faded rapidly under such conditions, losing essentially all colour in a few weeks, and stated that such material was therefore unsuitable for jewellery use. Some variability in behaviour was noted.

Accelerated fade testing often involves intense illumination combined with an elevated temperature to speed up the fading even more, typically less than 60 °C.

However, it is obvious that if an elevated temperature by itself changes the colour, then only such a temperature should be used for acceleration as does not itself have any effect. Nevertheless, the use of a temperature that might be present under normal illumination conditions, perhaps 50 °C (122 °F) maximum, would be reasonable.

Materials and experimental procedures

Examination of 40 beryls, the property of Dr al Gobaisi, was conducted under restricted access with limited time and equipment in January 1994 in Singapore, in part with the assistance of E. Wong and M. Stern. Conventional gemmological techniques were used, including particularly the nature and orientation of the dichroic colours and the absorption spectrum, the latter of these by use of the hand spectroscope on all 40 beryls (referred to hereafter as aG beryls).

Six of these beryls, two selected at random from each colour, were studied in more detail: intense blue, a 33.29 ct oval and a 40.59 ct cushion; yellow, a 33.77 ct oval and a 68.86 ct cushion; and greenish, a 33.85 ct cushion and a 63.23 ct pear. These tests confirmed that all were natural beryls of exceptional quality, being flawless under 10x magnification. The absorption spectra of these six beryls were also run on a BYK Gardner Colorview Spectrophotometer at Herberts South East Asia Pte Ltd in Singapore, using in-house software with
limited wavelength resolution, with the assistance of H. Gian and Wong. Each specimen was run three times in the geometry shown in Figure 1 and the results are shown in Figures 2 to 4.

The unusual geometry of Figure 1 had been worked out in preliminary experiments with a 25.85 ct intense blue cushion beryl, property of P. Hung, obtained by him from the same Brazilian source as the 40 aG beryls. The spectrum of this beryl was obtained by the author on a Colorview machine with much better wavelength resolution software at Colortech Associates, Inc. of Oldwick, New Jersey, USA, and is shown in Figure 5. The capability to run polarized spectra was not available with either of these instruments. While not as good as results from a research type instrument, these data were more than adequate to distinguish Maxixe beryl from aquamarine; both could also enable distinction between original Maxixe and Maxixe-type beryls.

Stones used for fading and heating tests included three beryls from the collections of P. Hung (hereafter designated 'PH'), obtained by him from the same source as the 40 aG beryls: the above-mentioned 25.85 ct intense blue cushion, a 13.90 ct yellow oval, and a 6.15 ct greenish oval. There were also five older beryls from the collections of the author (hereafter designated 'KN'): an 8.75 ct intense blue emerald-cut Maxixe-type beryl, a 1.55 ct greenish yellow round, a 1.75 ct greenish yellow emerald-cut, and yellow rounds of 2.20 and 2.28 ct.

The rapid fade test used in the restricted time available for this study employed incandescent light from a 150 watt internal reflector bulb for 43 hours with the face-up stone at 25 cm from the bulb as in Figure 6. Photometric measurements showed that this was approximately equivalent to a continuous (i.e. 24 hours per day) exposure to maximum (noon) sunlight for one week. A fan kept the temperature of the stone well below 50 °C, actually at a measured 44±2 °C. These conditions were different from those I had used previously because of the limited time available and are not...
Fig. 3. Spectral curves of two yellow aG beryls (33.77 and 68.86 ct) run in triplicate as in Figure 1.

Fig. 4. Spectral curves of two greenish aG beryls (33.85 and 63.23 ct) run in triplicate as in Figure 1.

Fig. 5. Spectral curve of the 25.65 ct intense blue PH beryl run as in Figure 1 but with higher resolution software than that used for Figures 2 to 4; there is agreement with the absorptions of Maxixe-type beryl (circles) but not with those of the original Maxixe beryl (squares), both taken from the 300K curves of Figure 2 of Reference 6.
necessarily preferred. This subject, as well as the question of the rapidity of fading will be discussed in detail elsewhere.

Some stones were also heated in the dark in an oven at 100±10 °C for one week, so that the effects of light and heat could be identified separately.

To avoid reliance on colour memory, which is notoriously fallible, or on equally difficult-to-reproduce photography, before and after colours were measured by comparison with the ISCC-NBS Centroid Color Charts15; such colours are given within quotation marks below.

The dichroism of Maxixe beryls shows the stronger blue colour in the ordinary ray, while the reverse is true for blue aquamarine. In using this test to identify Maxixe beryl, it is only the blue-producing absorption that is relevant. In the presence of much yellow, as in greenish aquamarine, a similar orientation of the dichroic yellow-causing absorption, which may also have this same dichroism orientation as described below, is not relevant for identifying Maxixe beryl.

**Intense blue beryl**

All the 28 large (20.10 to 41.70 ct), intense blue aG beryls were examined by the author, by Wong and by Stern with both the dichroscope and the hand spectroscope; these tests showed Maxixe beryl spectra with the stronger dichroic colour in the ordinary ray. Since the two variants of Maxixe beryl are very similar in their spectroscopic absorptions, they cannot be distinguished with a hand-spectroscope.

With the assistance of Gian and Wong, the author was able to examine two of the intense blue aG beryls on the limited-capability BYK Gardner Colorview Spectrophotometer in Singapore. The results of Figure 2 confirmed that these were Maxixe beryls; the position of the absorption at about 680 nm indicated that this spectrum corresponded to Maxixe-type beryl. The higher resolution results of Figure 5, run in New Jersey on the intense blue PH beryl obtained from the same Brazilian source, supported this identification. On this basis, the intense blue aG and PH beryls match the later carbonate-containing Maxixe-type beryls; they do not match the earlier nitrate-containing original Maxixe beryls. Accelerated fading tests demonstrated the expected fading, albeit at a relatively slow rate; and the details of these tests will be reported elsewhere.

At this point it is pertinent to refer to Brown's report concerning an intense blue rectangular cushion cut aG Maxixe beryl of 32.60 ct and his assumption that it had been irradiated by man; this was not supported by any observational or analytical data. Except in a fortuitous situation where inappropriate irradiation resulted in some residual radioactivity on a stone, the author knows of no way to distinguish such blue beryls that have been irradiated naturally from those exposed to irradiation by man.

It can be concluded that all the intense blue aG and PH beryls are indeed Maxixe-type beryl and can therefore be expected to fade on exposure to light. It cannot be established whether the colour derives from irradiation by nature or by man.
Golden beryls
Golden beryl has a weak dichroism, basically a darker-yellow to lighter-yellow, often with brownish or greenish overtones, as described in gemmology texts. The orientation of the darker colour is not usually given. Some published golden beryl spectra show a stronger absorption and therefore deeper colour in the ordinary ray. Some texts, such as that of Hurlbut and Kammerling, give the darker colour in the extraordinary ray. A recent check of a randomly chosen yellow beryl in the GIA collection by Kammerling (unpublished observation) gave the darker colour in the ordinary ray. It thus appears probable that the orientation of the stronger hue in dichroic yellow beryl can be either along the ordinary ray or extraordinary ray directions.

It is widely recognized that the colours of both golden beryls and green aquamarines are stable to light and that both lose their yellow component on being heated, turning colourless and blue, respectively. Only a few minutes are required to complete this change at 500 °C; it takes hours at 250 °C. This is the well-known process by which most aquamarine beryl is heated to convert it from the as-mined greenish colour into the more commercially desirable blue aquamarine.

To reconfirm this universally accepted behaviour and extend it to even lower temperatures, four yellow to greenish beryls (the PH13.90 ct ‘pale orange yellow’, the PH6.15 and KN1.55 ct ‘light greenish yellows’, and the KN2.28 ct ‘light to brilliant yellow’) were exposed to the 43 hour rapid fade test of Figure 6, where the fan kept the temperature of the stones at 44±2 °C. There was no change in colour, i.e. no fading, in any of these beryls. However, when held at 100±10 °C for one week in the dark, the 1.75 ct ‘light greenish yellow to light yellowish green’ beryls changed to ‘light yellow’. Some yellow coloration was being slowly destroyed in both instances even at this low temperature.

The author and Wong and Stern examined four large (33.77, 37.71, 68.86 and 126.66 ct), high quality yellow aG beryls, one of which (37.71 ct) had also been examined in detail by Brown, and was in Brown’s ‘after fading’ (and heating) state when examined. The dichroism results agreed with that of Brown: it is the ordinary ray that carries the stronger yellow colour, but in contrast to Brown’s results, no absorption lines for Maxixe beryl were detected using the hand spectroscope.

Instrumental spectra were again run, with the assistance of Gian and Wong, on two of the four yellow aG beryls in the arrangement of Figure 1, with the results shown in Figure 3. The spectra show an absorption minimum in the 600 to 650 nm region, this being the region where previously measured blue Maxixe beryls show a broad absorption band maximum, as seen in Figure 2 where similar experimental conditions obtained. Therefore these beryls totally lack Maxixe beryl absorptions, indicating ordinary golden beryl, which does not fade in light.

Thus, in the author’s opinion, there is no reason to doubt that the four yellow aG beryls, as well as the related yellow pH beryl, are ordinary golden beryls. Here again, it cannot be established whether these golden beryls were coloured by irradiation by nature or by man.

Green aquamarine or Maxixe beryls?
A greenish beryl colour can originate in two different ways. First, it could result from a combination of the yellow of golden beryl with the light blue of aquamarine, together giving green aquamarine. Here light does not produce fading but heat destroys the yellow component as discussed above and widely practised commercially.
Alternatively, a similar green could result from a combination of the yellow of golden beryls and the blue Maxixe beryl coloration as described by Nassau et al.\textsuperscript{6} Such a combination was also reported by Crowningshield who noted the many narrow absorption lines in a research-type prism spectroscope;\textsuperscript{20} these lines are clearly those of a Maxixe-type beryl spectrum.\textsuperscript{4-6} Rink\textsuperscript{21} produced such material by electron irradiation and was careful to distinguish the two colour components, also stating that the yellow component is stable to light. Here a fading of the blue component would be expected from light while heat would destroy both the blue (Maxixe) and the yellow (golden beryl) components.

Both types of greenish beryls may derive their yellow component (as well as their blue Maxixe beryl component, if present) from irradiation by nature or by man; again these cannot normally be distinguished by gemmological or by any other testing.

The eight large (33.85 to 74.20 ct) aG beryls with a colour range from light green to greenish-yellow were examined by the author, by Wong and by Stern. The stones closely resemble typical green aquamarines in their original as-mined state before such material is subjected to heat treatment to convert it to blue aquamarine. The possibility that irradiation had been used to intensify the yellow colour component, or restore it after it had been removed by a heating, cannot be ruled out.

Examination of the dichroism indicates that it is the ordinary ray that carries the stronger colour in all these aG greenish beryls. Such dichroism is reasonable for the yellow component and it would be expected for a Maxixe beryl blue component if present, but not for an aquamarine blue component. Dichroism by itself cannot normally be a reliable characteristic to distinguish the latter two possibilities. Indeed, we could not clearly establish the dichroic orientation of the blue-producing entity. In retrospect, it is possible that colour filters could perhaps have been used to separate the blue and yellow dichroisms but such an approach has not yet been investigated. Therefore, currently, only a detailed spectroscopic examination or a light-only fade test can help establish the presence or absence of Maxixe beryl. Hand spectroscope examination showed no absorptions in the Maxixe region in any of these aG beryls, and this was confirmed by results obtained from the spectrophotometer for two of the greenish aG beryls (Figure 4). The spectra are exactly those expected for the combination of golden beryl plus blue aquamarine. Again, there is a minimum in absorption in the region where the broad band Maxixe beryl absorption is seen in Figure 2. This conflicts with the results obtained by Brown\textsuperscript{12} from one of the stones (50.35 ct rectangular cushion) in the aG collection.

On the basis of these results there is thus no reason to doubt that the eight greenish aG beryls, as well as the related greenish PH beryl, are anything other than ordinary green aquamarine beryls. Here again, it cannot be established whether the yellow-producing component in these greenish beryls derives from irradiation by nature or by man.

Conclusions
It is concluded that the 29 large, high quality intense blue aG and PH beryls of recent Brazilian origin are of the Maxixe-type carbonate variety, a material that was seen widely about 1970 (but never in such large sizes) and has been seen only occasionally since then. This conclusion agrees with that of Brown\textsuperscript{12} on the 32.60 ct intense blue beryl.

The author also concludes that the five yellow and nine greenish large, high quality aG and PH beryls of recent Brazilian origin are non-fading golden beryls and non-fading greenish aquamarines, respectively. These conclusions conflict with the results reported by
Brown for the 37.71 ct golden beryl and the 50.35 ct greenish beryl, which he examined by permission of Dr D.M.K. al Gobaisi.

It is worth emphasizing three points:
(i) yellow (golden) beryls do not fade on exposure to light but will lose colour slowly on exposure to a temperature of 100 °C and more rapidly at higher temperatures;
(ii) aquamarines do not fade on exposure to light, but any yellow component present in greenish aquamarines will again be lost slowly at 100 °C and more rapidly at higher temperatures; and
(iii) while Maxixe beryl has the stronger dichroic blue colour in the ordinary ray, it is only the blue-producing absorption that has been and is relevant for reaching such an attribution; the yellow-causing absorption in golden beryl, which is also present in greenish aquamarine and which may also have this same dichroism orientation as described above, is not relevant in deciding whether or not a beryl can be described as Maxixe.

For gemmological purposes, it is also recommended that the term 'Maxixe' beryl be used for both the original Maxixe and the later Maxixe-type variants, since this distinction is difficult to establish without spectrophotometers.

Acknowledgements
I wish to thank Michael Stern of Gems, Toronto, Canada, Eric Wong of Ideal Gemological Laboratory Pte Ltd, Singapore, and Henry Gian of Omega Medical and Scientific Pte Ltd, Singapore, for assistance with various parts of the experimental work in Singapore; Dr D.M.K. al Gobaisi of Abu Dhabi, United Arab Emirates, and the Singapore Court for access to the PH beryls; and Peter Hung of House of Hung Pte Ltd, Singapore, for the use of the PH beryls.

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Emerald and green beryl from Central Nigeria

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Abstract

Mineralogical and gemmological characteristics of emeralds and green beryls from the central Nigerian Kaduna and Plateau States are presented. A genetic model for the occurrence of these gemstones in greisen associations of Mesozoic alkali granite ring complexes has been established. Gemmologists should be able to distinguish between Nigerian emeralds and those from other localities. Distinction is based on the unique inclusions and the chemical features of the Nigerian emeralds and green beryls. For those Nigerian specimens that contain solid inclusions, the mineral association of albite + fluorides (fluorite, boldyrevite, ralstonite) + F-silicate + Fe-rich mica + ilmenite + monazite, is locality-specific. Growth structures and fluid inclusions are extremely common and the later ones show a large range of features including Colombian-type three-phase inclusions. For Nigerian emeralds that show an inclusion pattern similar to that of Colombian emeralds, the easiest distinction is by absorption spectroscopy. The Nigerian emeralds show 'mixed spectra' with peaks attributable to Cr³⁺, Fe²⁺, Fe³⁺ and Fe²⁺/Fe³⁺, whereas the spectra of Colombian emeralds, as a rule, are largely free of Fe-components.

Introduction

Since the beginning of the 1980s, Nigeria has become a country of special interest for the gem industry, with significant occurrences in the Kaduna and Plateau States of different coloured beryls, tourmalines and sapphires, some of a very good quality (Bank, 1984; Kanis and Harding, 1990). At the beginning of the 1980s and again at the beginning of 1991, probably a few thousand carats of Nigerian emeralds or green beryls entered the gem market for periods of a few months.

The beryl, tourmaline and topaz deposits of central Nigeria show a great variety in their modes of occurrence. Some may be related to pegmatites, quartz veins, stockworks and greisens within or near granite contacts. Others occur in fissures, joints and shear zones at the edges of granite bodies or in syngenetic miarolitic and schlieren formations.

Emeralds and green beryls in the Plateau and Kaduna States of central Nigeria are much less common than their blue counterpart, aquamarine. They occur in rocks of two distinct ages. The older occurrences are usually pegmatites, whereas the younger are in the 'root' zones of granites. Although some beryls from both kinds of sources have an 'emerald colour', most are rather pale green and many dealers do not accept them as emeralds, classifying them as green beryl. Arps and Zwaan (1995) gave the source of the central Nigerian emeralds as east of Gwantu (south east Kaduna State) and north west of Nassarawa Eggon (Plateau State).
This paper presents the first comprehensive study of the mineralogical and gemmological characteristics of Nigerian emeralds and green beryls and establishes a genetic model for the occurrences of these gemstones in central Nigeria.

**Background and previous studies**
Emeralds and green beryls (Figure 1) are found together with gem-quality beryls of various other colours in similar geological environments in the Kaduna and Plateau States of central Nigeria.

When the Nigerian emeralds and green beryls reached the gem market at the beginning of the 1990s, they attracted attention for their large size (up to 100 grams) and beauty of their well-formed crystals. They also provoked a revival of the old debate on emerald terminology i.e. on the dividing line between emeralds and 'normal' green beryls (Hänni, 1992). The physical and mineralogical properties of beryls from central Nigeria have been described by several authors (Lind et al., 1984; Henn et al., 1984; Kiefert and Schmetzer, 1990; see Table IV below).

**Geology and genesis**
The geological history of central Nigeria covers a long period, even in geological terms! The main geological events and associated rock types are summarised in Table I.

The Nigerian Basement Complex contains a variety of rock suites of different ages spanning a period from 3000–450 million years (m.y.) ago. The oldest rocks were originally sediments and intrusive bodies which have undergone periods of folding and metamorphism and have been altered to migmatite and granite gneiss. Younger sediments were deposited onto

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Fig. 1. Well-formed crystals of emerald or green beryl (length about 4 cm) from the Kaduna and Plateau States, Central Nigeria. Photo Gebr. Henn, Idar-Oberstein
Table I: Summary of the geology of central Nigeria

<table>
<thead>
<tr>
<th>Time divisions</th>
<th>Age in million years</th>
<th>Geological events with ages in million years and associated rock types</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quaternary</td>
<td>Recent -0.2</td>
<td>Well preserved volcanic cones and alkali olivine basalt flows.</td>
</tr>
<tr>
<td></td>
<td>Pleistocene -2</td>
<td>Sedimentary series with intercalated tholeiitic basalt lava flows, now largely eroded.</td>
</tr>
<tr>
<td></td>
<td>Pliocene -11</td>
<td>Alkaline basaltic and rhyolitic volcanism, alkaline granite intrusives with cassiterite, columbite, wolframite, sphalerite and beryl/emerald. Younger Granites</td>
</tr>
<tr>
<td></td>
<td>Miocene -25</td>
<td>Alkaline basaltic and rhyolitic volcanism, alkaline granite intrusives with cassiterite, columbite, wolframite, sphalerite and beryl/emerald. Younger Granites</td>
</tr>
<tr>
<td></td>
<td>Oligocene -40</td>
<td>Alkaline basaltic and rhyolitic volcanism, alkaline granite intrusives with cassiterite, columbite, wolframite, sphalerite and beryl/emerald. Younger Granites</td>
</tr>
<tr>
<td></td>
<td>Eocene -55</td>
<td>Alkaline basaltic and rhyolitic volcanism, alkaline granite intrusives with cassiterite, columbite, wolframite, sphalerite and beryl/emerald. Younger Granites</td>
</tr>
<tr>
<td></td>
<td>Palaeocene -65</td>
<td>Alkaline basaltic and rhyolitic volcanism, alkaline granite intrusives with cassiterite, columbite, wolframite, sphalerite and beryl/emerald. Younger Granites</td>
</tr>
<tr>
<td>Cenozoic</td>
<td>Paleocene 65</td>
<td>Metasedimentary rocks and granitoid intrusives, pegmatite bodies with cassiterite, tantalite and beryl/emerald. Older Granites.</td>
</tr>
<tr>
<td>Tertiary</td>
<td>Jurassic -135</td>
<td>Metasedimentary rocks and granitoid intrusives, pegmatite bodies with cassiterite, tantalite and beryl/emerald. Older Granites.</td>
</tr>
<tr>
<td></td>
<td>Triassic -180</td>
<td>Metasedimentary rocks and granitoid intrusives, pegmatite bodies with cassiterite, tantalite and beryl/emerald. Older Granites.</td>
</tr>
<tr>
<td></td>
<td>Permian -225</td>
<td>Metasedimentary rocks and granitoid intrusives, pegmatite bodies with cassiterite, tantalite and beryl/emerald. Older Granites.</td>
</tr>
<tr>
<td></td>
<td>Carboniferous -270</td>
<td>Metasedimentary rocks and granitoid intrusives, pegmatite bodies with cassiterite, tantalite and beryl/emerald. Older Granites.</td>
</tr>
<tr>
<td></td>
<td>Devonian -350</td>
<td>Metasedimentary rocks and granitoid intrusives, pegmatite bodies with cassiterite, tantalite and beryl/emerald. Older Granites.</td>
</tr>
<tr>
<td></td>
<td>Silurian -395</td>
<td>Metasedimentary rocks and granitoid intrusives, pegmatite bodies with cassiterite, tantalite and beryl/emerald. Older Granites.</td>
</tr>
<tr>
<td></td>
<td>Ordovician -440</td>
<td>Metasedimentary rocks and granitoid intrusives, pegmatite bodies with cassiterite, tantalite and beryl/emerald. Older Granites.</td>
</tr>
<tr>
<td></td>
<td>Cambrian -500</td>
<td>Metasedimentary rocks and granitoid intrusives, pegmatite bodies with cassiterite, tantalite and beryl/emerald. Older Granites.</td>
</tr>
<tr>
<td></td>
<td>Late -570</td>
<td>Metasedimentary rocks and granitoid intrusives, pegmatite bodies with cassiterite, tantalite and beryl/emerald. Older Granites.</td>
</tr>
<tr>
<td>Palaeozoic</td>
<td>Late -1000</td>
<td>Kibaran 1600 Granite gneisses of central and SW Nigeria</td>
</tr>
<tr>
<td></td>
<td>Middle -1800</td>
<td>Eburnean 2400 Granite gneisses of central and SW Nigeria</td>
</tr>
<tr>
<td></td>
<td>Early -2600</td>
<td>Liberian 3000 Banded gneisses of SW Nigeria</td>
</tr>
<tr>
<td></td>
<td>Late -3300</td>
<td></td>
</tr>
</tbody>
</table>
the Basement and folded together with this granitized basement during the Pan-African orogeny from 750 to 450 m.y. ago, to form long north-south oriented linear schist belts of low grade metasedimentary rocks.

During the late Pan-African orogeny, from around 600 to 450 m.y. ago, there was a phase of granitoid magmatism. These granitoids were intruded into both the Basement Complex and the younger metasedimentary cover. The granitic rocks associated with the Pan-African orogeny generally have contrasting petrological and chemical compositions compared with the later Jurassic (Mesozoic) suite of alkaline ring complexes and are therefore called 'Older' Granites, in contrast to the 'Younger' Mesozoic Granites. The closing stages of the orogeny from 650-450 m.y. ago were marked by cooling, uplift and fracturing, by the eruption of volcanic rocks and the formation of pegmatitic lenses and dykes.

Within the Basement Complex is a zone in which this suite of pegmatites cut Old Eburnean rocks and early Pan-African metamorphic and igneous assemblages. The mineralized pegmatites are widely, but unevenly distributed within a broad zone, extending from the Ife area towards the Younger Granite province. It appears that the mineralized pegmatites occur within the north-south oriented younger metasedimentary belts composed of biotite schists and amphibolites. The pegmatites occur as dykes and sheets of varying dimensions - the dykes range from a few centimetres to tens of metres in width, with strike lengths of up to 2 km. The pegmatite sheets can have a considerable thickness of up to 40 m, and have been of economic interest because they contain cassiterite and columbo-tantalite. However the sheet-like pegmatites are of greater importance for the gems they contain. Many of the well mineralized pegmatites are characterized by pinch and swell structures. The swellings are generally very albitized and frequently correspond with centres of rich ore mineralization.

The younger granites of Nigeria, emplaced during the period 190 to 144 m.y. ago, form ring complexes typically 2-25 km in diameter and are composed of 95 per cent acid rocks. These Mesozoic ring complexes of Nigeria form part of a larger province of alkaline magmatism. They occur in a zone 200 km wide and 1600 km long, extending from northern Niger to south central Nigeria. This zone is related to the breakup of the Gondwana supercontinent during the Mesozoic era.

According to Dickin et al. (1991), the ring complexes of the Jos Plateau are the product of mantle-derived magmas which have suffered a significant amount of crustal contamination during their ascent and differentiation in the crustal basement. There are three main granite types: hornblende biotite granite; biotite granite and alkali granite with Na-Fe pyroxenes and amphiboles.

In the ring complexes, a series of hydrothermal alteration processes with related mineralization can be recognized (Kinnaird, 1979). Early sodic metasomatism (albitization) may have modified all the granite types. The effects of the later processes beginning with potassic metasomatism are only well documented for the biotite granites. Each hydrothermal process is characterized by a distinct assemblage of silicate minerals and there is a clearly defined sequence of ore deposition associated with each of the hydrothermal processes. It is with albitization processes that the gem varieties of beryl are associated. The temperature interval of the emerald and green beryl crystallization is about 500-400 °C.

Beryl-bearing basement pegmatites

Pegmatites are common throughout the Kabba Province and, in addition to quartz and microcline, contain both biotite and muscovite and in many cases considerable amounts of albite with beryl and
Tourmaline. Beryl crystals of the 'Basement' pegmatites can reach enormous sizes. To the east of the Niger River and north of the Benue River, there is an important pegmatite belt which is apparently not related to major granite intrusions. The pegmatites occur in an area bounded to the east by the Jos Plateau, to the south by the Afu Hills, to the west by Nassarawa and to the north by Kafanchan (Figure 2). These pegmatites vary considerably in their mineralogy. The bulk of those which are metal-bearing are complex albitized pegmatites with important gem potential. Typical minerals of these pegmatites are microcline, albite, perthite, quartz and muscovite. Accessory minerals are, in addition to beryl, tourmaline varying from black to blue and green, apatite, gem-quality blue gahnite spinel, garnet, nigerite.
(Zn,Mg,Fe$^{2+}$)(Sn,Zn)(Al,Fe$^{3+}$)$_2$O$_{22}$(OH)$_2$, chrysoberyl, cassiterite and columbotantalite.

In the Wamba and Akwanga regions (Figure 2), the microcline-quartz-muscovite pegmatites are strongly albitized, dyke-shaped bodies. The albite contains abundant small crystals of alkali-tourmaline and other accessories, including cassiterite, garnet, apatite, beryl and a little columbo-tantalite. In the Egometi district, where pegmatite dykes are mainly of microcline-quartz-muscovite varieties, accessory minerals are biotite, tourmaline, albite, garnet, apatite, beryl and cassiterite.

Gem-quality aquamarines have been collected in the Nassarawa Eggon area from Sabon Wana and Tundun Delli and emeralds are known from Kwafam Gwari.

These are likely to be the most important sources of gem quality beryls from the basement pegmatites, although there may be other localities not divulged by local gem dealers.

**Beryl-bearing Younger Granites**

In the Mesozoic ring complexes, gem quality beryl, topaz and smoky quartz are located in the 'roof' and contact zones of the latest intrusive stage of biotite alkali-feldspar granites, especially where the granite is in contact with the Basement Complex.

Gem quality beryl occurs as pale blue, greenish-blue or occasionally as pale green short stumpy hexagonal crystals often singly terminated. The gems occur in two settings, both formed in similar geological conditions.
Fig. 4. Growth structures in a Nigerian emerald or green beryl (view perpendicular to the c-axis). The most dominant features run parallel and perpendicular to the c-axis. $\times 70$

(i) The aquamarine and emerald may occur in small pegmatitic pockets in association with quartz, feldspar and topaz. These pegmatite pockets, which can be up to 8 cm in size, are found at the contact of a granite with basement rocks and represent cavities created by gas loss from the cooling magma. Localities of this type include the well known Timber Creek area near Rukuba, and the Barakin William occurrence in the Sha Kaleri Complex. In these pegmatitic pods crystals of more than 5 cm have been recorded. The stones from Barakin William, in particular, show distinctive colour variations within the crystal, either grading from colourless to deep blue or green along the c-axis of the crystal or with very dark colour bands, perpendicular to the c-axis, within otherwise pale crystals.

(ii) In the Janta area to the east of the Afu complex, 35 km SSW of Keffi, aquamarine and emerald have been found in soft decomposed granites. The crystals are pale blue, greenish or yellowish and less elongated than those from pegmatitic occurrences. Crystals have grown in small miarolitic cavities formed by gas loss in the ‘roof’ of the granite, within a zone of <20 m from its upper contact with the overlying rock type (Figures 3a and 3b). Fluid inclusion studies indicate crystallization at relatively high temperatures of 400–500 °C.
Observations and results

Crystal growth and trapping of inclusions

Growth structures are extremely common in Nigerian emeralds (compare Figures 4 and 5). Strongly developed growth bands parallel to basal-pinacoid, prism, and pyramidal faces are the most frequent kind (Figures 4 and 5), and are consistent with the descriptions given by Lind et al. (1984, 1986) and by Kiefert and Schmetzer (1990).

During the growth of most crystals, numerous kinds of mineral and fluid inclusions have been incorporated. Also, interruptions of the growth process are commonly indicated by ‘growth faces’ on which different types of mineral inclusions have been deposited, and this results in phantom-like phenomena. These growth faces are generally parallel to the crystal faces observed macroscopically for each crystal. Those crystals that contain a distinctly visible ‘core crystal’ surrounded by an ‘envelope’ which has a different colour are considered to consist of different beryl generations (Figure 6). During the hiatus between the different growth phases, corrosion phenomena have commonly developed on the faces of the early-formed crystal. When the crystal growth re-started, large fluid inclusions were trapped along planes principally parallel to the prism faces. Another indication that crystal growth was not steady and continuous but took place in several phases (‘pulses’) is the distinct colour zoning visible when looking down the c-axis of the crystals.

An additional indication of relatively fast crystal growth is the presence of distinct anomalous birefringence. This is caused by internal stresses and can be observed under crossed polars in sections of emerald and green beryl perpendicular to the c-axis. What is rare in the Nigerian beryls compared with those from other localities is the so-called ‘rain-effect’ which is caused by the presence of minute growth tubes oriented parallel to the c-axis of the host crystal. Fractures are common. Many have been partially healed by growth nutrient-bearing fluids as indicated by the presence of secondary fluid inclusions, while others are not healed. Most of the fractures are normally un-oriented with respect to the host crystal but some occur parallel to crystal faces. Epigenetic fracture fillings consist of Fe and Mn oxides.

Mineral inclusions

Mineral inclusions in Nigerian emeralds or green beryls have not been documented previously. This is understandable because they are not common; and only about 180-200 of the 1000 Nigerian emeralds/green beryls examined in this
study contain mineral inclusions. These have been studied in order to determine whether any inclusion features might be diagnostic in identifying the source of these emeralds or green beryls and might help in distinguishing them from synthetics.

The most common mineral inclusions in the Nigerian emeralds and green beryls are fluorite and albite which were identified in 80–100 samples (which is less than 10 per cent of the analyzed material). Tourmaline and K-feldspar inclusions only occur in a few samples, which means that their abundance is only a few tenths of a per cent, whilst other inclusion minerals have abundances of a few per cent.

The mineral inclusions vary in size with the largest up to 1–2 mm. Normally, there is no visible crystallographic orientation or control on the mineral inclusion distribution within the beryl host crystals. Only the minerals trapped on crystal faces during a period of growth interruption show a spatial distribution related to the emerald crystallography. Most of the mineral inclusions probably nucleated and grew only a short time before they were enclosed by the growing beryl crystals. Many are identifiable optically but some of the fluorides are not and these have been identified by microanalysis. The following list gives details of the mineral inclusions in approximate order of abundance.

(i) Albite
Albite, together with fluorite, is the most common mineral inclusion in Nigerian emeralds and green beryls. The albite crystals form colourless, transparent, angular to irregularly-shaped grains (Figure 7). In reflected light under a microscope, they generally appear translucent white because of their own numerous fine inclusions.

Feldspar inclusions of varying chemical composition have been described in African emeralds from different localities (Schwarz, 1994). They have also been encountered in emeralds from Colombia (Schwarz, 1995), Brazil (Hänni et al., 1987; Schwarz and Eidt, 1989; Schwarz et al., 1990), the Ural Mountains (Schwarz, 1991b), the Habachtal (Grundmann, 1991) and the Swat region in Pakistan (Schwarz, 1995). Consequently the identity and abundance of feldspar inclusions are of little diagnostic value when determining the origin of an emerald (Table II).
<table>
<thead>
<tr>
<th>Inclusion</th>
<th>Locality of emerald host</th>
<th>Reported by</th>
</tr>
</thead>
<tbody>
<tr>
<td>Albite</td>
<td>Mananjary, Madagascar</td>
<td>Schwarz, 1994</td>
</tr>
<tr>
<td></td>
<td>Cordillera Oriental, Colombia</td>
<td>Schwarz, 1995</td>
</tr>
<tr>
<td></td>
<td>Sandawana, Zimbabwe</td>
<td>Fillmann, 1987</td>
</tr>
<tr>
<td></td>
<td>Lake Manyara, Tanzania</td>
<td>Delé-Lasir and Poirot (pers. comm., 1991)</td>
</tr>
<tr>
<td></td>
<td>Carnaíba/BA, Brazil</td>
<td>Schwarz and Eidt, 1989</td>
</tr>
<tr>
<td></td>
<td>Socotó/BA, Brazil</td>
<td>Schwarz, 1987</td>
</tr>
<tr>
<td></td>
<td>Khaltaro, Pakistan</td>
<td>Gubelin, in Kazmi and Snee, 1989</td>
</tr>
<tr>
<td></td>
<td>Mingora-Swat, Pakistan</td>
<td>Schwarz, 1995</td>
</tr>
<tr>
<td></td>
<td>Panjshir, Afghanistan</td>
<td>Schwarz, 1995</td>
</tr>
<tr>
<td>Fluorite</td>
<td>Colombia (?)</td>
<td>Roulet, 1956</td>
</tr>
<tr>
<td></td>
<td>Madagascar</td>
<td>Delé-Lasir and Poirot (pers. comm., 1991)</td>
</tr>
<tr>
<td></td>
<td>Pakistan</td>
<td>Eppler, 1984</td>
</tr>
<tr>
<td></td>
<td>Ural Mountains</td>
<td>Shernakow and Laskowenkov, 1991</td>
</tr>
<tr>
<td></td>
<td>Emmaville/NSW</td>
<td>Stevens, 1980</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Brown, 1984</td>
</tr>
<tr>
<td>Fe-rich mica</td>
<td>not reported</td>
<td>Schwarz, 1995</td>
</tr>
<tr>
<td>Mg-biotite/phlogopite</td>
<td>most occurrences worldwide, e.g. Belmont Mine/MG</td>
<td>Hänni et al., 1987</td>
</tr>
<tr>
<td></td>
<td>Capoeirana-Nova Era/MG</td>
<td>Schwarz et al., 1988a</td>
</tr>
<tr>
<td></td>
<td>Tauá/CE</td>
<td>Schwarz et al., 1988b</td>
</tr>
<tr>
<td></td>
<td>Carnaíba/BA</td>
<td>Schwarz and Eidt, 1989</td>
</tr>
<tr>
<td></td>
<td>Socotó/BA</td>
<td>Schwarz et al., 1990</td>
</tr>
<tr>
<td></td>
<td>Santa Terezinha/GO</td>
<td>Schwarz, 1990</td>
</tr>
<tr>
<td></td>
<td>Goiás State, Brazil</td>
<td>Barros and Kinnaird, 1987</td>
</tr>
<tr>
<td></td>
<td>Western Australia</td>
<td>Schwarz, 1991a</td>
</tr>
<tr>
<td></td>
<td>(Poona, Menzies)</td>
<td>Schwarz, 1991b</td>
</tr>
<tr>
<td></td>
<td>Ural Mountains</td>
<td>Grundmann, 1991</td>
</tr>
<tr>
<td></td>
<td>Habachtal</td>
<td></td>
</tr>
<tr>
<td>Inclusion</td>
<td>Locality of emerald host</td>
<td>Reported by</td>
</tr>
<tr>
<td>-----------------</td>
<td>------------------------------------------------------------------------------------------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>Ilmenite</td>
<td>Mananjary, Madagascar</td>
<td>Schwarz, 1995</td>
</tr>
<tr>
<td></td>
<td>other African localities (Zambia, Zimbabwe, Tanzania, South Africa, Egypt)</td>
<td>Schwarz, 1995</td>
</tr>
<tr>
<td></td>
<td>Mananjary, Madagascar</td>
<td>Hänni and Klein, 1982</td>
</tr>
<tr>
<td></td>
<td>Sandawana/Machingwe</td>
<td>Schwarz, 1995</td>
</tr>
<tr>
<td></td>
<td>Santa Terezinha/GO, Brazil</td>
<td>Cassedanne and Sauer, 1984</td>
</tr>
<tr>
<td></td>
<td>Ural Mountains</td>
<td>Barros and Kinnaird, 1987</td>
</tr>
<tr>
<td></td>
<td>Habachtal, Austria</td>
<td>Shernakov and Laskovenkov, 1991</td>
</tr>
<tr>
<td>Boldyrevite</td>
<td>not reported</td>
<td></td>
</tr>
<tr>
<td>Ralstonite</td>
<td>Belmont Mine/MG, Brazil (?)</td>
<td>Souza, 1988</td>
</tr>
<tr>
<td>Monazite</td>
<td>Habachtal</td>
<td>Grundmann, 1991</td>
</tr>
<tr>
<td></td>
<td>Itaberaí/GO</td>
<td>Schwarz, 1995</td>
</tr>
<tr>
<td>Quartz</td>
<td>Many localities worldwide</td>
<td>Kleyenstüber, 1991</td>
</tr>
<tr>
<td></td>
<td>Mananjary, Madagascar</td>
<td>Schwarz, 1994</td>
</tr>
<tr>
<td></td>
<td>Lake Manyara, Tanzania</td>
<td>Bank and Gübelin, 1976</td>
</tr>
<tr>
<td>K-feldspar</td>
<td>Mananjary, Madagascar</td>
<td>Schwarz, 1987</td>
</tr>
<tr>
<td></td>
<td>Mananjary, Madagascar</td>
<td>Herrmann, 1991</td>
</tr>
<tr>
<td></td>
<td>Mananjary, Madagascar</td>
<td>Graziani and Lucchesi, 1979</td>
</tr>
<tr>
<td></td>
<td>Lake Manyara, Tanzania</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Socotó/BA, Brazil</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Capoeirana/MG, Brazil</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Salininha/BA, Brazil</td>
<td></td>
</tr>
<tr>
<td>Tourmaline</td>
<td>many occurrences worldwide, e.g. Zambia</td>
<td>Koivula, 1982, 1984</td>
</tr>
<tr>
<td></td>
<td>Mananjary, Madagascar</td>
<td>Schwarz, 1987</td>
</tr>
<tr>
<td></td>
<td>Zimbabwe</td>
<td>Anderson, 1976</td>
</tr>
<tr>
<td></td>
<td>Mananjary, Madagascar</td>
<td>Schwarz, 1995</td>
</tr>
<tr>
<td></td>
<td>Ural Mountains</td>
<td>Hänni and Klein, 1982</td>
</tr>
<tr>
<td></td>
<td>Habachtal</td>
<td>Schwarz, 1994</td>
</tr>
<tr>
<td></td>
<td>Habachtal</td>
<td>Grundmann, 1991</td>
</tr>
<tr>
<td></td>
<td>Carnaiba/BA</td>
<td>Schwarz and Eidt, 1989</td>
</tr>
<tr>
<td></td>
<td>Socotó/BA</td>
<td>Schwarz et al., 1990</td>
</tr>
<tr>
<td></td>
<td>Panjshir, Afghanistan</td>
<td>Schwarz, 1995</td>
</tr>
<tr>
<td>Emerald/beryl</td>
<td>Many occurrences worldwide</td>
<td></td>
</tr>
<tr>
<td>Mg-Ca-Al-Na-Fluosilicate</td>
<td>not reported</td>
<td></td>
</tr>
</tbody>
</table>
(ii) **Fluorite**

Fluorite is one of the most frequent mineral inclusions; most are developed as perfect octahedra (Figure 8)

![Fig. 7. Colourless, transparent, angular to irregularly shaped albite grains. x 100](image)

**Fig. 8.** Perfectly developed fluorite octahedra. x 100

but some are cubes and others have a more or less rounded or irregular shape (Figure 9), sometimes occurring as aggregates. Many fluorite crystals show a strong surface structure which may be an etching phenomenon. The fluorite inclusions are colourless but because of their low refractive index they show a very strong relief in the beryl host crystal and consequently appear very dark when immersed in fluid and examined under the gem microscope (Figures 8 and 9). In dark field illumination, they appear colourless-transparent or greyish-white.

Fluorite has been identified as an inclusion in emeralds from several localities. It was noted in Colombian emeralds by Roulet (1956), although it is not known by which method the inclusion mineral was identified and the identification has not been confirmed in subsequent studies on other Colombian emeralds. Fluorite has been noted in emeralds from Madagascar and Pakistan (Delé-Lasir and Poirot, pers. comm., 1991), the Ural Mountains (Eppler, 1984; Sherbakow and Laskowenkow, 1991; Schwarz, 1995) and Emmaville, NSW, Australia (Stevens, 1980; Brown, 1984; Schwarz, 1995).

(iii) **Mica**

The mica inclusions in the Nigerian emeralds and green beryls are Fe-rich, most likely in the compositional range annite/siderophyllite or zinnwaldite. Energy dispersive X-ray spectrometry is insensitive to lithium and until this element content is measured it is not possible to fully identify the mica. The mica crystals are flaky or book-like in appearance, dark brown or reddish in colour with a strong pleochroism and basal cleavage (Figure 10).

![Fig. 9. Irregularly shaped fluorite crystals with very strong relief. x 140](image)
Mica inclusions are common in emeralds from other localities but are usually of Mg-biotite or phlogopite composition. The occurrence of a Fe-rich mica in the Nigerian emeralds and green beryls is a distinctive feature and is a strong indication that these gems are from the alkali biotite granites of the ring complexes where Li-Fe biotites are the major mafic mineral (Kinnaird, 1985).

(iv) Ilmenite
Ilmenite occurs as opaque-black platelets, sometimes forming aggregates, or as irregularly shaped grains with a metallic lustre. This inclusion mineral has been identified in emeralds from several localities (see Table II).

(v) Boldyrevite
The occurrence of boldyrevite (NaCaMgAl₃F₁₄ · 4 H₂O) was identified by chemical microanalysis. It cannot be distinguished optically from fluorite because their crystal habits and refractive indices are very similar. The association of different fluoride inclusions in a Nigerian emerald is shown in Figure 11.

(vi) Ralstonite
The identification of the fluoride ralstonite, Naₓ(Alₓ₋ₓMgₓ) (FₓOₓH₉₋ₓ) · H₂O is also based on microanalysis. It also cannot be distinguished optically from fluorite on the basis of crystal form or refractive index.

(vii) Monazite
Monazite, (Ce,La,Th)PO₄, inclusions in Nigerian emeralds and green beryls show distinct relief as transparent prismatic crystals or aggregates which are transparent, colourless or of a weak yellowish colour. Analyses show that these monazites are very Th-rich.

Generally, monazite is a very rare mineral inclusion in emerald, having been found previously only in emeralds from the Habachtal (Grundmann, 1991) and the small occurrence at Itaberai in the Goiás State of Brazil (Schwarz, 1995). The composition of the monazite at both these localities differs from that of the inclusions in the Nigerian stones, those of the Habachtal-emeralds being more Ce-rich (Grundmann, 1991), and the Itaberai-emeralds being Ce/La-rich varieties (Schwarz, 1995).
Emeralds containing tourmaline inclusions. Tourmaline inclusions have been identified in emeralds from widely different occurrences (Table II).

(xii) Fluorosilicate
It was not possible to identify precisely a Mg-Ca-Al-Na-fluorosilicate that was observed in one of the samples examined. The existence of fluorosilicates as daughter crystals in fluid inclusions in aquamarines from the Younger alkali granites of the Afu complex has been noted by Kinnaird (1985).

Comparing the association of the inclusion minerals in the emeralds and green beryls with the mineral associations in the different granite types of central Nigeria, there is a strong similarity to the mineral associations described by Kinnaird (1984) for the earliest mineralization stage in the alkali granites. In these, the most important accessories are U-pyrochlore, cryolite (Na-Al-fluoride), fluorite, thomsonite (hydrated Na-Ca-Al-fluoride), zircon, Th-rich monazite and ilmenite. The minerals identified as inclusions in the Nigerian emeralds/green beryls are several fluorides, ilmenite, Th-rich monazite, quartz, feldspar (normally albite) and a Fe-rich representative of the mica group (probably annite/siderophyllite). All these minerals can be linked in terms of their origin to alkali granites. The micas of the alkali granites are characterized by very high Fe-contents (MacLeod et al., 1971) and this is valid also for mica inclusions.
in the emerald or green beryl. Potential inclusion minerals that may be found during further investigations of the emeralds are U-pyrochlore and zircon.

**Fluid inclusions**

Nigerian emeralds and green beryls show a large variety of fluid inclusions, some of which may exceed more than 1 mm in size. The primary cavities which contain the fluid inclusions show tubular, partly spiky, or irregularly shaped forms (Figure 12). Secondary cavities are irregularly shaped, normally with a rounded outline, sometimes, however, showing a 'jagged' outline (Figure 13). There is an enrichment of inclusions in certain more or less well-defined areas of the emerald host crystal, and this is considered to be a growth phenomenon. Such features include the more frequent appearance of fluid inclusions in the border zones of crystals or the common enrichment of growth tubes in the core of a crystal.

The fluid inclusion population of aquamarines, together with inclusions in topaz, fluorite and quartz from different localities in the younger granites of central Nigeria, have been described by Kinnaird (1985) who identified six different fluid inclusion-types, some of which have been found in the present study.

Most of the fluid inclusions in the Nigerian beryls have very strong relief and appear almost opaque in transmitted light; this is due either to the difference between the refractive index of the cavity filling and the surrounding emerald or to the irregularity of the cavity walls. Both of these features enhance reflection rather than transmission light and result in a dark image. The most abundant type of inclusion in emeralds and aquamarines from central Nigeria is one containing liquid, a vapour bubble and several solids (multiphase 's-lg' inclusions' with several s-phases; Figures 12, 14, 15). Such multiphase inclusions are also known in

![Fig. 14. Primary multiphase fluid inclusion of the 's-lg' type with a large isotropic cube (probably halite). × 100](image1)

![Fig. 15. Planar arrangement of multiphase fluid inclusions. × 70](image2)
Table III: Chemical compositions of emeralds and beryls from the Kaduna and Plateau States in Central Nigeria

<table>
<thead>
<tr>
<th></th>
<th>1</th>
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<td>(a)</td>
<td>(b)</td>
<td>(a)</td>
<td>(b)</td>
<td>(a)</td>
<td>(b)</td>
<td>(c)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SiO₂</td>
<td>66.10–68.40</td>
<td>68.82/68.61</td>
<td>68.87/68.39</td>
<td>69.11/69.09/68.74</td>
<td>65.17</td>
<td>65.45</td>
<td>65.83</td>
<td>65.53</td>
<td>67.31</td>
<td>61.35–68.81</td>
<td>65.74–67.54</td>
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<td>Al₂O₃</td>
<td>18.02–19.21</td>
<td>18.61/17.88</td>
<td>18.48/17.86</td>
<td>18.57/18.63/18.04</td>
<td>18.84</td>
<td>18.24</td>
<td>18.70</td>
<td>18.39</td>
<td>18.40</td>
<td>14.86–18.49</td>
<td>18.01–18.18</td>
</tr>
<tr>
<td>FeO_{tot}</td>
<td>0.11–0.95</td>
<td>0.28/0.87</td>
<td>0.23/1.10</td>
<td>0.14/0.35/0.89</td>
<td>0.46</td>
<td>0.89</td>
<td>1.13</td>
<td>0.31</td>
<td>0.25</td>
<td>940*–12170*</td>
<td>0.06–0.91</td>
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<tr>
<td>Cr₂O₃</td>
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<td>&lt; / 0.06</td>
<td>&lt; / 0.06</td>
<td>&lt; / 0.02/0.21</td>
<td>0.01</td>
<td>&lt;</td>
<td>0.02</td>
<td>0.08</td>
<td>0.07</td>
<td>&lt; −480*</td>
<td>0.06–0.11</td>
</tr>
<tr>
<td>V₂O₅</td>
<td>&lt; −0.06</td>
<td>0.02/0.06</td>
<td>&lt; / 0.09</td>
<td>&lt; / 0.07/0.06</td>
<td>0.03</td>
<td>&lt;</td>
<td>0.01</td>
<td>0.06</td>
<td>0.03</td>
<td>&lt; −280*</td>
<td>0.05–0.07</td>
</tr>
<tr>
<td>TiO₂</td>
<td>&lt; −0.02</td>
<td>&lt;</td>
<td>&lt;</td>
<td>&lt; &lt;</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>&lt; −0.04</td>
<td>0.07–0.63</td>
</tr>
<tr>
<td>MgO</td>
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<td>0.02/0.06</td>
<td>0.03/0.06</td>
<td>0.08/0.03/0.05</td>
<td>0.03</td>
<td>0.04</td>
<td>0.02</td>
<td>0.20</td>
<td>0.03</td>
<td>0.03–0.05</td>
<td>0.04–0.06</td>
</tr>
<tr>
<td>Na₂O</td>
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<td>0.12/0.15</td>
<td>0.13/0.16</td>
<td>0.13/0.06/0.12</td>
<td>0.10</td>
<td>0.17</td>
<td>0.14</td>
<td>0.15</td>
<td>0.07</td>
<td>0.08–0.57</td>
<td>0.09–0.12</td>
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<tr>
<td>CaO</td>
<td>&lt; −0.01</td>
<td>&lt;</td>
<td>&lt;</td>
<td>&lt; / 0.01</td>
<td>–</td>
<td>–</td>
<td>–</td>
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<td>–</td>
<td>0.01</td>
<td>0.02–0.05</td>
</tr>
<tr>
<td>K₂O</td>
<td>&lt; −0.05</td>
<td>0.01/</td>
<td>&lt;</td>
<td>&lt; / 0.02</td>
<td>&lt; / 0.01/0.05</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.04–0.21</td>
<td>0.01–0.02</td>
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<tr>
<td>CS₂O</td>
<td>&lt; −0.05</td>
<td>0.01/</td>
<td>0.02/0.02</td>
<td>0.01/0.02/0.03</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>MnO</td>
<td>&lt; −0.02</td>
<td>&lt; / 0.01</td>
<td>0.01/</td>
<td>&lt; &lt;</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.01–0.05</td>
<td>–</td>
</tr>
<tr>
<td>Sc₂O₃</td>
<td>&lt; −0.03</td>
<td>&lt;</td>
<td>0.01/</td>
<td>&lt; / 0.01</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Li₂O</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.02</td>
<td>0.02</td>
<td>0.02</td>
<td>0.01</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

Key:
- < contents below detection limit
- = not analyzed or range
* = element concentration in ppm
tr = traces

1: light to medium green and bluish-green (variation observed in 13 samples)
2–4: crystals with colour zoning; colour intensity c>b>a
5–8: Lind et al., 1986;
5 = intense blue
6 = light blue
7 = light green
8 = intense green
9: Schrader, 1987
10: Henn et al., 1984; range of 6 samples
11: Henn and Bank, 1991
12–14: J. Zang, pers. comm., 1992;
crystals with colour zoning; colour intensity b>a
emeralds from other localities: for example, those from Colombia normally are three phase ‘slg’ inclusions with an isotropic daughter phase, usually halite.

Healed fractures are common in Nigerian emeralds and green beryls. Some are parallel to the hexagonal prism faces or to the basal pinacoid of the emerald host crystal (Figure 13) and are practically planar. More commonly, the healed fractures are undulating and without crystallographic orientation, often appearing as veil-like formations; some show a delicate net-like structure or contain larger flattened cavities whilst others contain numerous small rounded cavities resembling strings of pearls. Larger pseudosecondary inclusions along healed fractures vary in type, some are multi-phase inclusions with or without birefringent solid phases. Three-phase inclusions of the ‘Colombian type’ (Figure 14) seem to be less frequent in the healed fractures than in primary cavities.

Chemistry
The chemical compositions of 32 beryls from central Nigeria are given in Table III. Since colourless, blue, blue-green and emerald-green beryls with varying intensities of colour may occur within the same paragenesis, their chemical characteristics will be discussed together.

The representative points of more than 100 emeralds from nine different localities in Africa are plotted in terms of their Al₂O₃ and MgO contents in Figure 16 (from Schwarz, 1995). The localities are:

Fig. 16. Weight % Al₂O₃/MgO correlation diagram with representative points for emeralds from different African localities (1 = Ndola Rural District, Zambia; 2 = Alto Ligonha, Mozambique; 3 = Sandawana/Machingwe, Zimbabwe; 4 = Mananjary, Madagascar; 5 = ‘Egypt’; 6 = Kaduna/Plateau States, Nigeria; 7 = Lake Manyara, Tanzania; 8 = ‘South Africa’; 9 = Cobra Mine, South Africa.)
(1) Ndola Rural District (Kafube Belt), Zambia
(2) Alto Ligonha, Mozambique (including three pegmatite beryls)
(3) Sandawana-Machingwe mining area, Zimbabwe
(4) Mananjary mining region, Madagascar
(5) Eastern desert, Egypt
(6) Kaduna-Plateau State, Central Nigeria
(7) Lake Manyara, Tanzania
(8) ‘South Africa’ (exact locality not known)
(9) Cobra Mine, Transvaal, South Africa.

The emerald occurrences in Zambia, Zimbabwe, Mozambique, Madagascar, Egypt and South Africa all belong to ‘schist-type’ deposits; that is, they are associated with various schist rocks containing biotite or phlogopite, actinolite or tremolite and variable quantities of talc, chlorite or carbonates, probably derived essentially from a mafic-ultramafic protolith. Because of this environment, the MgO concentrations of emeralds from these rocks are generally between 1 and 3 wt%. The negative, approximately linear correlation between Al₂O₃ and MgO indicates that magnesium enters the beryl structure principally by substituting for aluminium. Such a substitution of Al³⁺ by bivalent ions such as Mg and/or Fe requires a coupled substitution to maintain a neutral charge balance, normally with Na in the following way:

\[ \text{Al}^{3+} = \text{Mg}^{2+} + \text{Na}^{1+} \] (channel)

or

\[ \text{Al}^{3+} = \text{Fe}^{2+} + \text{Na}^{1+} \] (channel)

In contrast to the emeralds from schist deposits, the Nigerian emeralds and green beryls are characterized by extremely low magnesium and sodium contents. This suggests that they are largely from the alkali granites of the Younger Granite suite which have extremely low contents of Mg. The suggestion made earlier that the majority of the Nigerian emeralds and green beryls containing fluorite and Fe-rich mica are probably of Younger Granite origin is consistent with this low Mg content.

Ferromagnesian minerals in the Nigerian alkali granites are always Fe or Na-Fe end members, so these elements must have been available during the albitization stage of alteration. Nevertheless, despite the availability of Na, the Na-contents of the Nigerian beryls are consistently low (see Table III and Figure 17). This is in accordance with observations on emeralds from other localities, especially in the Cordillera Oriental, Colombia where Schwarz (1995) has shown that Na is incorporated in the channels of the emerald structure only if there is some need for charge compensation caused by the entrance of bivalent cations \( \text{Me}^{2+} \) in the \( \text{Al}^{3+} \) positions. If there is no substitution of \( \text{Al}^{3+} \) by \( \text{Me}^{2+} \), the emeralds show only low Na-contents, even if the mineralizing fluids are sodium-rich.

The iron-concentrations of the Nigerian emeralds and beryls show a strong variation, reaching \( \text{FeO}_{\text{tot}} \) values of up to 1.2 wt% (Figure 17, from Schwarz, 1995) and the high contents are consistent with iron being available during the beryl growth. As is shown by the absorption spectra (see below), at least part of the iron of the Nigerian emeralds or green beryls is present as Fe³⁺ (substituting for Al³⁺) and therefore the beryl needs no sodium for charge compensation. The correlation diagram (Figure 17) shows that there is no correlation between the sodium and the iron contents for these emeralds.

In some of the analyzed samples that show colour-zoning, the concentration of the chromophore elements, especially iron, increases with the intensity of the bluish-green colour. Such colour variation is not related directly to the elements Mg and Na although they may be involved with chromophores in a charge-balancing role.

Comparing the values of optical data of emeralds from the schist-type deposits with those of the Nigerian emeralds and green beryls, the latter are low. The same is
Fig. 17. Weight % Na$_2$O/FeO$_{tot}$ correlation diagram with representative points for emeralds from different African localities (1–9 see Figure 16)

Table IV: Optical data of beryls/emeralds from the Kaduna/Plateau States, Central Nigeria.

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<th>$n_d$</th>
<th>Birefringence</th>
<th>Reference</th>
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<td>1.566–1.568</td>
<td>1.572–1.575</td>
<td>0.006–0.007</td>
<td>Bank, 1984</td>
</tr>
<tr>
<td>1.565–1.567</td>
<td>1.570–1.572</td>
<td>0.005</td>
<td>Henn et al., 1984</td>
</tr>
<tr>
<td>1.564–1.568</td>
<td>1.570–1.574</td>
<td>0.006</td>
<td>Lind et al., 1986</td>
</tr>
<tr>
<td>1.569–1.570</td>
<td>1.574–1.576</td>
<td>0.005–0.006</td>
<td>Henn and Bank, 1991</td>
</tr>
<tr>
<td>1.560–1.569</td>
<td>1.566–1.574</td>
<td>0.005–0.006</td>
<td>This paper</td>
</tr>
</tbody>
</table>
Fig. 18. (a,b,c,d) Absorption spectra of emeralds and green beryls from the Kaduna and Plateau States in Central Nigeria. The spectra show different combinations and intensity ratios for the absorption bands of Fe\(^{2+}\), Fe\(^{3+}\), Fe\(^{2+}/\text{Fe}^{3+}\), V\(^{3+}\) and Cr\(^{3+}\).
true for the specific gravity values: 2.63-2.65 (Henn et al., 1984), 2.66-2.68 (Lind et al., 1986), 2.65-2.67 (Henn and Bank, 1991) and 2.64-2.68 (this paper). After Henn and Bank (1991) and Lind et al. (1984), the absorption spectra of the Nigerian emeralds represent ‘mixed types’ with absorption patterns relating to an emerald component (Cr/V) and an aquamarine component (Fe$^{2+}$ and Fe$^{2+}$/Fe$^{3+}$). Similar spectra have been reported for emeralds from Zambia (Schmetzer and Bank, 1980) and Madagascar (Hänni and Klein, 1982; Schwarz and Henn, 1992). The spectra of Figures 18a to 18d indicate that the Nigerian emeralds exhibit no consistent pattern. Almost all ratios of intensity between the different absorption bands of Cr, Fe$^{2+}$, Fe$^{3+}$ and Fe$^{2+}$/Fe$^{3+}$ have been seen in different Nigerian specimens (see also Schwarz, 1995), and the intense Cr$^{3+}$ absorption bands observed in some samples show that the designation ‘emerald’ is justified at least for some of the beryls from the Kaduna and Plateau States.

**Discussion and conclusions**

There are two provenances for emeralds and green beryls in central Nigeria, one from pegmatites intrusive into the Basement complex, the other from Mesozoic alkali granite ring complexes. From their solid inclusions and chemistry, the large majority of the emeralds and green beryls in this study appear to have a granitic origin.

The very few beryls with tourmaline inclusions most likely originate from Basement pegmatites, whereas those which contain fluorite and Fe-rich mica originate from the Younger Granites.

The particular geological setting and the special genetic conditions result in a unique set of mineralogical-gemmological properties of the emeralds and green beryls from central Nigeria. These distinguish central Nigerian emeralds from all other main emerald deposits in the world. The association of mineral inclusions, comprising such ‘exotic’ representatives as boldyrevite-ralstonite-monazite-Fe-rich mica, together with the presence of a large variety of fluid inclusions, has no counterpart in beryls from other occurrences.

The presence of fluorite as a mineral inclusion in emeralds is consistent with the newest models postulated to explain emerald occurrences (Schwarz, 1995). Although it is likely that beryllium is transported in most cases as a complex with fluorine, fluorite (CaF$_2$) is apparently only a common mineral inclusion in emeralds if the emerald crystallization is related to greisenization processes (Schwarz, 1995). Th-rich monazite and the Fe-rich micas, also found in Nigerian beryls, are typical minerals of greisen associations. The occurrence of other fluorides (boldyrevite and ralstonite) and fluorosilicates indicates a specific genetic relationship for the emerald and green beryl mineralization in Nigeria. Boldyrevite and fluorosilicates have not been described as inclusions in emeralds from other localities, and the identification of ralstonite in one emerald from the Belmont Mine in Minas Gerais (Souza, 1988) has not been confirmed.

In the Younger Granites, emeralds and green beryls were probably formed during the late magmatic and early subsolidus hydrothermal stage, based on solid mineral inclusions and the nature of the fluid inclusion evidence. Early hydrothermal fluids associated with the granite magma were saline and of high temperature (about 400-500 °C) and the fluid inclusions in the emeralds and beryls normally contain several daughter minerals.

The unusual chemical properties, especially the very low Mg and Na concentrations of the emeralds and green beryls from central Nigeria, can also be explained considering the (geo-)chemical features of their host rocks. These are characterized by low Mg values. The incorporation of Na in the channels of the
emerald structure does not depend on the availability of this element in the mineralizing fluids. The controlling factor is the need for charge compensation caused by the entrance of bivalent cations. Consequently, the Na-concentration is low because (Mg+Fe)$_2^{2+}$ in the emeralds and green beryls is also low. The low Cr and V concentrations of the host rocks is reflected by low contents of these elements in the beryls. The large variation of the Fe-contents in the emeralds and green beryls is due to the possibility of incorporating iron as Fe$^{2+}$ and/or Fe$^{3+}$ in different lattice positions (including channel sites).

Both beryllium and fluorine are enriched in the alkali granites of Nigeria compared to 'average' values in granites worldwide, and both elements may occur in a range of minerals. The source of the chromophore elements, Cr and V, in the Nigerian beryls is problematic because both elements are scarce (0–10 ppm) in Nigerian granites. However, most beryl crystallization occurs close to the 'roof' of a granite in the region where granite fluids may react with surrounding country rocks. Thus Cr and V may have been incorporated from the basement schists or younger volcanic rocks both of which have higher Cr and V contents than many of the granites. The model calculations of Kozlowski et al. (1988) for the emerald occurrences of the Cordillera Oriental in Colombia, show that extremely low chromophore contents in the host rocks are sufficient to provide enough Cr and V for the crystallization of significantly coloured green beryls (emeralds). This suggests that it is not necessary to have exceptional enrichment of these elements in the country rocks of the Nigerian beryls.

The optical data for the beryls and emeralds from the Kaduna and Plateau States can be interpreted bearing in mind their chemical characteristics. The 'mixed type' spectra with almost all ratios of intensity between the different absorption bands of Cr, Fe$^{2+}$, Fe$^{3+}$ and Fe$^{2+}$/Fe$^{3+}$ are in accordance with the (crystal-) chemistry of the beryls.

Gemologists should be able to distinguish between emeralds from central Nigeria and those from other localities. For those Nigerian beryls that contain solid inclusions, the mineral association of albite + fluorides + Fe-rich mica is unique. For those Nigerian beryls which are free of solid inclusions, the inclusion pattern may be similar to that of certain Colombian emeralds. However, the 'classic' fluid inclusions of the Colombian emeralds are 'slg'-three-phase-inclusions with a cubic halite crystal, whereas the emeralds and green beryls from central Nigeria are more commonly 's,lg'-multiphase inclusions which, in addition, contain one or more (anisotropic) s-phases. Absorption spectra can also aid in origin identification. The 'mixed-type' spectra of the Nigerian beryls are quite different from the spectra of the Colombian emeralds which, as a rule, are free of iron bands (see also Schwarz, 1995).

Acknowledgements

G. Becker of Fa. Friedrich August Becker, Stefan Kehl, F. Gebr. Herrn - all of Idar-Oberstein, Germany, provided samples. Bettina Becker and Dr Joachim Zang of the Department of Gemstone Research, Johannes Gutenberg University, Mainz, did some SEM and microprobe analyses. The studies of D. Schwarz were part of a research project sponsored by the German Research Association (DFG) and carried out at the Department of Gemstone Research, Johannes Gutenberg University of Mainz, Germany.

All photomicrographs are by D. Schwarz.

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Smaragdvorkommen in Brasilien entdeckt: Capoeirana bei Nova Era, Minas Gerais. Zeitschrift der Deutschen Gemmologischen Gesellschaft, 37(3-4), 146-7


Note 1. Examples for the fluid-inclusion nomenclature used in this paper:

- lg - liquid-gas two-phase inclusion
- llg - liquid-liquid-gas three-phase inclusion
- slg - solid-liquid-gas three-phase inclusion
- sllg - multi-phase inclusion with several different solid phases, one liquid phase and one gas phase

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Abstracts

Diamonds

Cathodoluminescence properties of CO$_2$-bearing and CO$_2$-free diamonds from the George Creek K1 kimberlite dike.

CL properties of diamonds from the George Creek K1 kimberlite dike, in the State Line kimberlite district of Colorado-Wyoming, USA, are discussed. Diamonds shown by IR spectroscopy to contain sub-microscopic, high-pressure CO$_2$ inclusions show anomalous CL colours, including pink, pale violet, orange and brown, while CO$_2$-free diamonds show typically blue.

M.O'D.

Lab Notes.

A very light green marquise diamond contained a row of included crystals most of which appeared to be diamond although at least one crystal was prismatic and yellow green suggesting pale diopside or enstatite. Testing showed that this inclusion was the sole cause of the colour. A visible spectrum showed absorption at 741 nm, the primary line of the GRI centre, showing that the stone had not been exposed to ionising radiation. Small dark spots around some of the included crystals were graphite.

An eye-visible brownish-orange inclusion breaking the table surface of a fancy yellow diamond was shown to be almandine-spessartine garnet by EDXRF.

A yellow round brilliant diamond showed a natural iridescence in a fracture which had also been filled. In the iridescent area colours were seen simultaneously whereas the flash-effect colours were seen one at a time; green in brightfield and red in darkfield illumination. At high magnification the filling in the fracture showed small bubbles and flow lines.

R.J.P.

Gem news.

Firms in Ramat Gan, Israel, have fracture-filled 30,000 ct of diamond in the last 30 months. All sizes and shapes of diamonds are being treated and the greatest demand is for those with an apparent clarity of SI after fracture filling.

Synthetic diamonds were misrepresented recently as Canadian rough. All the stones were about the same size, colour and shape (very atypical for a mine run sample) and showed evidence of metallic flux inclusions.

R.J.P.

ABSTRACTORS

For further information on many of the topics referred to consult Mineralogical Abstracts or Industrial Diamond Review.
A slightly brownish-grey diamond crystal contained a cavity with a mobile triangular inclusion which was of the same type (la) as the host. The original material filling the cavity was thought to be gas which had escaped during the cutting of a window.

A visit to the kimberlite pipes in the Siberian Sakha (Yakutia) Republic showed that flooding was a serious problem. There was very little activity at two of the sites and both the international pit as well as the quarry-like Zarnitza prospect were inactive. Udachnaya seemed to be the most productive and its large processing plant was in full production servicing not only its own but also the Aikhal and Jubilee sites. When winter temperatures reach -45°C work stops because of an inversion layer which traps potentially deadly methane in the mine. R.J.R.

Diamonds are reported from the Akuluâk dyke in the Central Churchill Province, Gibson Lake area, Northwest Territories. N.D. MacRae, A.E. Armitage, A.L. Jones and A.R. Miller. *International geology review*, 37, 1995, pp 212-29.

Diamonds are reported from the Akuluâk dyke in the Central Churchill Province, Gibson Lake area, Northwest Territories of Canada. The dyke is a narrow lamprophyre intrusion of ultra-potassic composition and has yielded a large number of diamonds, few of which are larger than 0.5 mm. Crystal forms are primarily octahedra and tetraxahedroids though some macles, cubes, aggregates and fragments also occur. The dyke is situated approx. 120 km NW of Rankin Inlet. M.O'D.

A study of diamonds formed in the Popigai and Ries impact craters has shown that they are similar in their mode of occurrence, morphology, physical properties, crystalline structure, etc. They occur as polycrystalline aggregates of cubic and hexagonal phases and inherit the shape and some other features of the original graphite. Authigenic impact diamonds were established to exist in both craters in strongly shocked and partially fused fragments of graphite-bearing crystalline target rocks. After the solid transition from graphite at ~ 40-45 GPa, impact diamonds were included in hot media and subjected to oxidation and graphitization. R.A.H.

**Carbon isotopic composition and origin of SiC from kimberlites of Yakutia, Russia.**


Analyses are presented of carbon isotopic compositions of moissanite (SiC) from the Mir and Aikhal kimberlites. Yakutia Province. The mineralogy of moissanite is described and illustrated with an SEM image. Silicon metal is the most common inclusion and inclusions of ferro-silicate, Fe-Ti silicides, REE silicate and sinito have been found. δ¹³C ranges -22 to -29‰, and δ¹⁵N +9.7 ± 4.0 to +5.6 ± 2.0‰. Moissanite may be a relic of a reduced, primordial Earth and now present as a trace phase in an otherwise oxidized mantle. Alternatively there may be present-day global regions that are highly reduced with light C isotopic compositions. Other possibilities are discussed, including the formation of moissanite by metamorphism of reduced, carbonaceous sediments during subduction. R.K.H.

**Distribution of the luminescent centres in Yakutian diamonds.**


The laser-luminescence tomography method for revealing crystal zoning is described and has been used to examine diamonds collected from the Malobotubinskoie field in the Mirnyi region of Yakutia, Russia. This non-destructive method shows the distribution of luminescence centres in the crystals, usually arranged in zones formed during successive stages of crystal growth. 'Phantoms' inside crystals, i.e. luminescent outlines of the growth zones, even from the earliest stages, provide information on changes in diamond crystallization conditions in the parent rock matrix. The method has been developed for use in the classification of diamonds from both primary kimberlite and secondary alluvial deposits in the Mirnyi region. The paper is illustrated with nine colour photographs showing yellow, green or blue UV luminescence. R.A.H.
The Luanda diamond fields. Part II.
Continuing account of a visit to the diamond-producing area of Luanda with some details of recovery methods.

Heart and arrow shaped pattern observed in round brilliant-cut diamond.] (In Japanese with English abstract)
A recent claim that gained popularity on the Japanese diamond market was that if a brilliant cut diamond exhibits a heart and arrow shaped pattern when illuminated under certain conditions, this proves that it has the highest grade of cut. Even an instrument to illuminate and record the pattern has appeared on the market. This paper reports the results of a mathematical analysis of the reliability of such a claim. It has been demonstrated that (1) such a pattern can be seen when a stone is illuminated in a range of inclination of 9 to 15 degrees to the axis, (2) the pattern is mainly due to the arrangements of pavillion facets and not to the crown facets, and so (3) this pattern does not automatically represent the highest cut grade. The appearance of the pattern is a necessary condition but not a sufficient condition to prove the cut grade.

Gems and Minerals

Detrital origin of fuchsite-bearing quartzites in the western Dharwar craton, Karnataka, India.
Quartzites in the Nuggihalli, Holenarsipur, Javanhalli, Bababudan and Chitradurga schist belts all originated as clastic sandstones rather than as precipitated cherts. Detrital chromite occurs in the oldest quartzites and this gave rise to fuchsite during metamorphism; there is no evidence for significant contributions of hydrothermal Cr. There are secular trends in median Cr/Al₂O₃, Cr/Fe₂O₃, Cr/Zn and K₂O/Al₂O₃ ratios which are consistent with the gradual evolution of mafic and silicic sedimentary source terrains over the interval from ~3500-2500 m.y. ago. The compositions of the Javanhalli and Bababudan quartzites are sufficiently different to suspect, but not prove, that they are from different sources.

Charoite: a unique mineral from a unique occurrence.
M.D. EVDOKIMOV. World of Stones, 7, 1995, pp 3-11, illus. in colour.
Details of the properties and occurrence of charoite are given, with notes on the geology of the occurrence and a list of references. The charoite-producing source (Sirenevyi Kamen [lilac stone]) is found on the meridian passing from the town of Olekminsk on the Lena River to Khani Station on the Baikal-Amur railway line, Russia.

Kazakhstan and Middle Asia. A brief mineralogical guide.
A.A. EVSEEV. World of Stones, 8, 1995, pp 24-30, illus. in colour, 1 map.
List of mineral species found in Kazakhstan and Middle Asia, arranged in alphabetical order of location and including some gem minerals.

Recent colored gem stones from Russia. [In Japanese with English abstract]
Diamond, ruby, alexandrite, demantoid, topaz, emerald, aquamarine, tourmaline, diopside, jadeite and lapis-lazuli from Russia which have recently been encountered in the Japanese gem market are described. Chemical and analytical data obtained by electron dispersive spectroscopy, FT-IR charts, and XRD are given. Possible marketing routes of Russian emeralds are explained.

Gem identification by FT-IR-emerald, jadeite, sillimanite, etc. [In Japanese with English abstract]
Fourier transform infrared (FT-IR) spectroscopy has been applied extensively to identify resin-impregnated emeralds and jadeites. FT-IR spectra show different features characteristic of natural, flux-grown, hydrothermally-grown emeralds and also resin impregnated emeralds, and jadeites, that can be used for diagnostic purposes. FT-IR spectroscopy is also useful to differentiate between gemstones whose refractive indices are similar, such as sillimanite, kornerupine, diopside and enstatite.
Rubis et saphirs de République Populaire de Chine.
M.O'D.

Gem-quality grossular-andradite: a new garnet from Mali.
Thousands of carats of these fashioned stones from the Republic of Mali are poised to enter the marketplace. In the trade these garnets are termed 'Mali garnet' or 'Mali grossular garnet'. The composition of this garnet is intermediate between grossular and andradite garnets and has been described by the GIA Gem Trade Laboratory as grossular-andradite. Quantitative chemical analysis using an electron microprobe and X-ray powder techniques revealed a composition 78% grossular and 20% andradite. Refractive indices for yellow-green and green stones are in the range of 1.752–1.769, whilst those of orange brown stones are 1.773–1.779 (RIs for grossular 1.73–1.76 and andradite 1.880–1.895).

The absorption spectra of yellow-green, and orange brown stones show an absorption band at 440 nm. The stacked parallel planes of growth zoning are always visible between crossed polars and together with moderate to strong ADR are diagnostic of grossular-andradite garnet.

Most of the gem quality grossular-andradite garnets from Mali are reminiscent of the pale greenish yellow to yellow-green grossular garnets from Tanzania, with some rare stones resembling peridot. Numerous brown to brownish-orange stones can be quite brilliant due to higher RIs and dispersion and may serve as inexpensive substitutes for 'cognac' diamonds.  
R.J.P.

Mosaic geographical map – a masterpiece of stonecutting art.
M.N. Kalinina. World of Stones, 8, 1995, pp 31–2, illus. in colour.
Brief account of a mosaic map in the Georgian Hall of the Hermitage Museum in St Petersburg, Russia. The map is constructed from hard ornamental stones.  
M.O'D.

Lab notes.
A stone showing a colour change from dark bluish green in daylight to dark reddish purple in incandescent light was identified as natural alexandrite. It had atypically high RI values of 1.753–1.761 and was found to have unusually high concentrations of chromium, titanium and iron. Several surface-reaching fractures contained a transparent filler which easily melted.

Three purplish-red mixed-cut stones were identified as corundum but lacked distinctive absorption features. The facets had uneven surfaces with some slight rippling and many minute cracks. Immersion in di-iodomethane revealed concentration of the colour along the facet edges indicating that they were diffusion-treated corundums.

A translucent mottled-green carving in a closed pendant setting gave a spot RI reading of 1.66 and showed a series of 'chrome lines' in the red end of the spectrum which are consistent with natural-colour jadeite. However, the carving consisted of a thin hollow jadeite shell filled with a soft transparent colourless filler containing gas bubbles. FTIR spectroscopy of the filler indicated polymers known to have been used for jadeite impregnation. The thickness of the jadeite shell at the edge was only 0.05 to 0.10 mm.  
R.J.P.

Gem news.
Rare collector stones from the former Soviet Union were shown in Tucson in February 1995. These included small colourless crystals of leucophanite, colour-change clinochlore together with görgeyite, kaliborite and preobrazhenskite. The full constants of these were quoted.

Chatoyant moonstones with an atypical darker orange colour originated from Madras, India. Unusual opal carvings incorporating both opal and its matrix were cut from the back to produce a three-dimensional effect. Sapphires from Madagascar were well described. These had been heat treated and the blue colour was due to iron-titanium charge transfer. Details
Gem news.


A blue green apatite cabochon from Malagasy weighing 11.45 ct showed pronounced chatoyancy and an unusual colour. The chatoyancy was caused by barely visible white needles. 'Eilat stone' not from Israel but from Peru is now on the market. It is a by-product of copper mining. At 10.03 ct round cabochon-cut emerald from Brazil showed a well defined six-ray star. It showed strong pleochroism with intense blue-green colour parallel and a light yellowish-green colour perpendicular to the c-axis attributed to a distinct iron-bearing (aquamarine) component which was found in the spectrum together with chromium bands of emerald. Although there were channel-like inclusions orientated parallel to the c-axis, the star was caused by three sets of unidentified small elongated particles orientated perpendicular to the c-axis. A transparent purplish red-brown stone of 0.33 ct had RI 1.742, singly refractive, SG 3.68-3.70, inert to both long and short wave UV radiation. Unusually it showed a typical almandine-rhodonite visible-light absorption spectrum. It was confirmed as genthelvite, Zn₄Be₃(SiO₄)₆S, a member of the helvite group. X-ray powder diffraction analysis and energy dispersive X-ray fluorescence (EDXRF) spectrometry were used to identify the stone. A 'drusy' hausmannite cabochon showed an imposing architectural appearance with its natural crystalline surface. Green opal from Serbia appeared green through the Chelsea filter. Spot RI 1.46 and SG 2.10 values were obtained but were difficult to measure because the stone was porous. The stone was inert to short-wave UV radiation but fluoresced an even faint chalky green to long-wave UV. The spectrum showed cut-off edges at 450 and 620 nm. Some healed fractures and veining were visible under magnification. EDXRF analysis revealed the presence of nickel which was the probable cause of the colour. An Egyptian green quartz reportedly collected in the Eastern desert has a pseudo-fibrous structure. EDXRF analysis revealed a fairly large concentration of chromium which probably was the cause of the colour as finely disseminated chromium-coloured mineral inclusions. A spangled quartz is coloured by bright orange red flakes which are probably hematite. Although randomly orientated throughout the cabochon the flakes are concentrated in one layer parallel to a rhombohedral face and are unusual in that each grew outward from one corner in an elongated fashion to give an 'hour-glass' or 'cello' shape. An aromatic resin necklace sold as myrrh was thought not to be from the somewhat rare Commiphora myrrha of biblical fame. It showed RI 1.41(spot) SG 1.27, hardness less than 2.5 on Mohs' scale and on the approach of a hot needle easily melted emitting a characteristic sweet spicy odour. Purple scapolite from Tajikistan reputedly found in the Pamir mountains is mined at an altitude of 5000 m and miners wear oxygen masks when working. Specimens showed uniaxial negative character, RI 1.539 and 1.550, SG 2.61, inert to long wave UV and fluorescing a very weak pinkish orange to short wave UV. Another stone from this region was a spinel of 123 ct described as virtually flawless. An opticon-treated malachite showed enhanced lustre and depth of colour. The impregnation process consisted of repeated coatings and oven treatment at each stage so that finally the whole of the product was impregnated. R.J.P.

Spectrophotometric identification of emeralds. [In Japanese with English abstract]


UV-VIS and NIR spectra have been obtained on some hundreds of emerald samples from different origins, both natural and synthetic. In UV-VIS spectra, dominant absorption due to Cr can be seen in natural Colombian emeralds, while the absorption peaks due to both Cr and Fe are seen in natural emeralds from other localities. Hydrothermally-grown synthetic emeralds show absorption due to Cr and high transmission of UV. In Russian synthetic emeralds, an absorption related to Ni is seen. In flux-grown synthetic emeralds, only Cr-related absorption can be recognized, and the transmission peak in the UV lies at a higher wavelength than that of Colombian samples. From NIR (1200-2700 nm) spectra, the origin of emeralds and the conditions of their growth can be conjectured from the absorption peaks, since they show the presence or absence and the nature of H₂O. I.S.

[A few unusual gemstones recently encountered.] (In Japanese)

Y. Kitawaki. *Journal of the Gemmological Society of*
Natural bi-colour topaz, natural euclase, natural pink clinozoisite which were encountered in gem testing are described.

[A trial for grading black pearls.]


Based on the quality elements of colour and brilliance, efforts have been made to establish a method to quantitatively grade the quality of black cultured pearls produced by *Pinctada margaritifera*. The method of grading, inherent mechanism of colour emission, and the difference in the emission mechanisms between black and white pearls are described and discussed. Reflectance is measured and RGB analysis is done by image analyser. The results showed that the measured values of four quality elements (blackness, body colour, interference colour and its intensity) on the image analyser, were almost identical to the grades obtained by visual grading.

Native copper in agates from Rudno near Krzeszowice.


Copper is reported in agates from Rudno near Krzeszowice in Poland. Notes on the occurrence and on the agates are given and samples were taken from more than 300 agate specimens. Native copper was found to occur in secondary microfissures crossing agate layers obliquely, at the boundaries of individual agate layers in a tree-like form, and as dispersed grains independent of agate layers.

What's new in minerals?


Gem-quality minerals and fashioned stones are reported from mineral and gem shows held in the United States. Species seen at the Denver show include fine rhodochrosite from the Sweet Home mine, Alma, Colorado; smoky quartz from the Calumet iron mine in the same state; twinned crystals of chrysoberyl from Santa Tereza, Espirito Santo, Brazil; very deep pink octahedral fluorite crystals from Mont Blanc, France; emerald from the Jos region of Nigeria; brown to green vesuvianite from Sandare, Diakon Arrondissement, Nioro du Sahel, Mali (this area also produces brown-green grossular); sceptre amethyst from Obman, Yakutia, Russia; blue topaz from Nura-Taldy, central Kazakhstan; emerald crystals from the mica schists of the Malysheva mine, central Urals, Russia (the crystals are not of gem quality); fine elbaite showing green, colourless and blue sections perpendicular to the c-axis, from the Asthor mine, near Shigar, Pakistan.

From the show held in Pretoria, South Africa, David Minster reports fine green tourmaline crystals from Mozambique; rhodochrosite crystals up to 3 cm long from the Kalahari manganese field, South Africa; gem-quality yellow orthoclase and rhodizite from the Malagasy Republic.

La saga peregrina, [first part].


Enquiry, with some proposals, into the present whereabouts of the Peregrina, a celebrated pearl once in the possession of the Kings of Spain.

The co-precipitation of Fe$^{2+}$ and SiO$_2$ and its role in agate genesis.


Detailed experimental work studying the use of Fe$^{2+}$ and Mg$^{2+}$ in removing H$_4$SiO$_4$ and colloidal silica from separate dilute silica solutions and sols over periods of 45 min to 14 years is reported. It is demonstrated that Mg$^{2+}$-SiO$_2$ gels in an alkaline saline environment at room temperature can develop signs of transformation into opal-CT after 14 years. The concentrations of Fe$^{2+}$ found in some agates is considered to be coincidental and too low to precipitate silica from any silica sol or solution.

Ein trapiche-rubin aus Myanmar (Burma).


A ruby crystal of 6 ct and 12 mm thick showed six sections of ruby divided from one another by spokes of a light-coloured carbonate material tentatively identified as either calcite or ankerite. The piece closely resembled the trapiche emerald long known from the Muzo mine in Colombia.
The degree of metamictization in zircons: a Raman spectroscopic study.

A series of natural zircons representing various degrees of metamictization were investigated by Raman microprobe analysis. Systematic changes in wavenumbers and half-widths of the Raman bands were shown to be caused by increasing irregularities of bond-lengths and bond-angles and a general breaking-up of the structure as a result of metamictization. The half-width of the antisymmetric stretching vibration band \( B_{lg} \) of the SiO\(_4\) tetrahedra, which has a frequency of \( \sim 1007 \text{ cm}^{-1} \) in well-crystallized and \( 1000 \text{ to } 995 \text{ cm}^{-1} \) in metamict zircons, is most suitable for estimating the degree of metamictization: its value increases from \( \sim 5 \text{ cm}^{-1} \) in well-crystallized samples up to \( 30-55 \text{ cm}^{-1} \) in highly metamict X-ray amorphous zircons, and is strongly dependent on the degree of lattice destruction by metamictization. In contrast, the Raman parameters seem to be almost uninfluenced by chemical variations. The potential value of such measurements, especially in radiometric age determinations, is discussed.

R.A.H.

Antique cameos.

Descriptive paper introducing the classical motifs found in cameos, some of which are illustrated. Examples are taken from the collections of the Hermitage Museum, St Petersburg, Russia.

M.O'D.

Optical and infrared spectroscopic studies of epidote group minerals from the Rhodope region.

Crystals of various colours, including Fe, Mn-bearing clinozoisites, Mn-bearing zoisite (thulite) and epidote from skarns of various Bulgarian deposits, have been studied. Oriented single-crystal platelets have been used to record optically polarized, \( \alpha, \beta \) and \( \gamma \) spectra. The green colour of epidote is associated with \( \text{Fe}^{3+} \) occupying the \( M(3) \) sites in the crystal structure and is determined by a short-wave \( \text{O}^2+ \rightarrow \text{Fe}^{3+} \) transfer. The pink colour of Mn-clinozoisite and thulite is caused by Mn\(^{3+}\) in \( M(3) \) structural sites. The IR spectra of the two minerals show definite differences. The OH-group valency oscillation band at \( 210 \text{ cm}^{-1} \) is most important for identification of zoisite. Mn-bearing specimens show three maxima in the \( 500-600 \text{ cm}^{-1} \) interval whereas Fe-bearing ones show a single broad band which is explained with bond oscillations in the AlO\(_2\)-octahedra. A comparative analysis of infrared spectra of clinozoisite before and after its treatment with deuterium has shown that the OH-group band at \( 3370 \text{ cm}^{-1} \) and the band at \( 1045 \text{ cm}^{-1} \) (caused by Al-OH deformational oscillations) disappear. They are replaced by new bands at \( 2490 \) and \( 780 \text{ cm}^{-1} \). The absorption bands of the diorthogroups lined by \( M(3) \) octahedra are the most variable. This corroborates the assumption of preferable substitution of Al by Fe in the \( M(3) \) sites. A linear regression equation based on the valency frequency oscillations of diorthogroups is proposed for determining the Fe content.

R.A.

Epitaxial overgrowths of marcasite on pyrite from the Tunnel and Reservoir Project, Chicago, Illinois, USA; implications for marcasite growth.

The marcasites occur as thin tabular crystals \( \approx 300 \mu\text{m} \) on edge, dominated by \{010\}, and have grown around the \{001\} zone on pyrite octahedra. The marcasite has apparently grown around the pyrite by a layer growth mechanism as shown by macrosteps terminating partially developed layers on the marcasite \{010\}. Other forms on the marcasite are described and illustrated with SEM micrographs. A proposed reconstruction of the pyrite surface into a marcasite-like arrangement may be due to interactions of a protonated disulphide species with the pyrite during growth.

R.K.H.

Luminescence in minerals.
J. RAKOVAN AND G. WAYCHUNAS. Mineralogical Record, 27(1), 1996, pp 7-19, illus. in colour.

The property of luminescence is explained with reference to mineral structure and chemistry, drawing examples from some of the commoner species.

M.O'D.

Ruby and sapphire from Vietnam. (In Japanese)

Over 2400 samples of ruby and sapphire col-
lected from Luc Yen and Quy Chau, Vietnam were investigated. Crystal habits, inclusions, and internal features such as growth sectors and banding are described.

A guide to mineral localities in the former Soviet Union.

An alphabetical list of places where minerals have been found has been prepared for territories of the former Soviet Union. Each entry gives name in English with variants where appropriate, the name in Cyrillic characters, geographical co-ordinates, type of location (region, town, stream, village, bay, mountains and so on), location of the place of higher order (similar to state capital, province [Russian name *oblast*], local seat of government) and list of species found. Useful notes are given on how the table was compiled and the difficulties encountered, on how to transliterate and what scheme to use, and on the nature of Russian and non-Russian names in this specialized context. There is a table of useful terms and a transliteration table.

M.O'D.

Sapphires from Southern Vietnam.

Gem quality blue to bluish-green sapphires from Southern Vietnam came on the market in the late 1980s. The sapphires occur in alkali basalts in the mining areas of Phan Thiet and Di Linh. Mining was carried out by hand at the former but relatively sophisticated equipment was used at Di Linh.

The properties and characteristics of these sapphires are similar to those from other basalt sources such as Australia, Cambodia, China, Nigeria and Thailand. The crystals show prominent growth structures and colour zoning in geometrical patterns. Sharply bordered blue bands alternate with narrow colourless or yellowish to brownish bands parallel to the crystal planes r and z; many stones showed a colourless 'core'.

The most common inclusions are various types of cloud with cross-hatch or lath-like patterns which have not been recorded from other sapphires. Although occasional dust-like

'snow-flake' inclusions have been noted in Kashmir sapphires, the very fine whitish needles presumed to be rutile or ilmenite lacked the well formed strongly iridescent appearance of rutile needles and plates observed in sapphires from metamorphic environments such as Sri Lanka and Burma. Inclusions of plagioclase feldspar, zircon, uranpyrochlore, columbite, ilmenite, pyrrhotite, chromite-hercynite, magnetite-hercynite and goethite were identified. The UV-VIS-NIR absorption spectra are typical of basaltic sapphires with absorption bands towards the near-infrared region due to Fe$^{2+}$ → Fe$^{3+}$ intervalence charge-transfer. Spectral curves in the 280-880 nm region might be used to identify the country of origin of sapphires but should be interpreted with caution. The high iron content of Southern Vietnamese sapphires allowed easy distinction from sapphires from a metamorphic source. Separation from other basaltic sapphires was more difficult but the cross-hatch and lath-like 'cloud' patterns coupled with distinctive colour zoning and a colourless 'core' are a unique combination.

R.J.P.

Estructura cristallina, composiciö quimica I propietats fisiques de les gemmes.5. La forma dels cristalls. (In Catalan and Spanish)

The crystal structure of beryl is explained with diagrams showing the ring formation.

M.O'D.

Brief chronicle of the formation of the Vernadsky State Geological Museum.

Account of the events preceding and accompanying the establishment of the Vernadsky State Geological Museum, Moscow, Russia, with illustrations of some notable specimens. M.O'D.

Larimar: blauer Pektolith aus der Dominikanischen Republik.

The name Larimar has been given to an opaque blue ornamental variety of pectolite found in the Dominican Republic. The occurrence is on the south-west coast of the country.

M.O'D.
Die türkisgrünen Steine' von Nagar in Nord-
Pakistan.
W. TOROSIAN-BRIGASKY AND V.M.F. HAMMER. Mineralien Welt, 7(1), 1996, pp 47-51, 1 map.
Quartz crystals with a turquoise-green colour have been found at 3400 m in the Nagar area of northern Pakistan. M.O'D.

Among specimens shown at the 1995 Münchner Mineralientage in 1995 were emerald crystals from Mengdong, Yunnan Province, China and peridot crystals from Sopat, Kohistan, Pakistan. Fine orange-red rhodochrosite from southern Kazakhstan was also available, one specimen measuring 3.26 cm in width. While these specimens were illustrated, the text also mentions chrome cerussite from the Dundas Extended Mine, Tasmania; brazilianite crystals up to 4 cm long, from Minas Gerais and chrysoberyl trillings from Colatina, Espirito Santa, Brazil; these crystals are up to 2 cm. Pakistan has also produced morganite crystals up to 10-12 cm long. M.O'D.

Remarkable pyrite pseudomorphs after ammonites.
E.Y. ZAKREVSKAYA. World of Stones, 7, 1995, pp 24-5, illus. in colour.
Fine ornamental specimens of pyrite pseud­morphs after ammonite are described from the collections of the Vernadsky State Geological Museum, Moscow, Russia. M.O'D.

Fluorescence and luminescence in minerals.
Short papers on luminescence in minerals are abstracted. Titles covered are luminescence zoning in minerals; fluorescent minerals - some answers, questions and enigmas; single-colour and multi-colour fluorescence in calcite; luminescence mechanisms and activators in Terlingua-type calcite and in benitoite and related silica minerals; the influence of heating on the luminescence of calcite; uranium-activated fluorescence in minerals of the Franklin-Sterling Hill area, Sussex County, New Jersey; fluorescent minerals of the Karnes uranium district, south Texas; cathodoluminescence in the study of rocks and minerals; activators for cathodoluminescence of charoite and associated minerals; illuminating gems with visible monochromatic lasers - what can it do for the gemmologist? a combined micro­scope/fluorescence spectrometer system for mineralogical and gemmological applications; demonstration of mineral fluorescence excited by a 193-nm ArF excimer laser; the fluorescent characteristics of coal. M.O'D.

What's new in minerals?
A number of gem minerals have been seen at different mineral shows held in the United States during 1995. Examples include watermelon elbaite from the Otjua mine, Namibia, brown zircon from Lovozero, Kola Peninsula and golden-brown scheelite from Tenkergin, Chukotka, both Russia: fine green fluorite is coming in crystal sizes up to 13 cm on an edge from the Westmoreland mine, New Hampshire, USA, and peridot from the newly-discovered Pakistan location. Thumbnail-sized crystals of aquamarine and yellow beryl are coming from Cherlovaya Gora, Chita, Siberia and China continues to produce very high quality crystals of beryl, cassiterite and scheelite. It is reported that the violet red beryl claims in the Wah Wah mountains of Utah have been sold to a major mining company. A matrix group of crystals measuring 17.9 cm across, with about 25 crystals on it, has recently been recovered. Fine zoned elbaites in pastel blue, green and pink are coming from Pech, Kunar, Afghanistan.

Madagascar is producing more tourmaline: colourless crystals with ruby-red bands at each end have come from a small pocket south of the Mania river and liddicoatite crystals are coming from a site at Alakamisy, Anjanabonoina and from Ibity. Grass-green slabs with a perfectly symmetrical red-orange triangle in the centre have been found at old deposits in the Laondany area. Crystals of rhodizite are available as are hambergites: the export of both these species, with orthoclase, is forbidden. Some supposed hambergite on offer has turned out to be quartz or phenakite.

Crystals of rose- and peach-coloured morganite have been found in the past at the Bennet quarry, Buckfield, Maine, USA: New England beryl formed the theme of this year's Springfield show and a collection of Bennet beryls was shown by the Harvard Mineralogical Museum.
Very fine amethyst crystals are being found in opium-growing areas of Veracruz state, Mexico: this area is not recommended for visiting.

Gemmologie.

This issue comprises a booklet by U. Henn entitled ‘Practical Gemmology’ (Edelsteinkundliches Prakticum). It is a survey of methods of differentiation between natural and synthetic diamonds, corundums, spinels, beryls (emeralds), chrysoberyls (alexandrites), quartzes (amethysts and citrines) and opals. The introductory part deals with a short history, describes in general the characteristics of gemstones and their imitations, and discusses their microscopic characteristics. In the specific, second section each of the gems mentioned is dealt with fully, listing origin of name, physical and chemical properties, inclusions, going on to list the various methods of synthesis and their characteristics. Each chapter has an extensive bibliography and many illustrations (145 illustrations in all) as well as numerous tables and graphs.


This Golden Jubilee issue celebrates the 50th Anniversary of the formation of the GAA which was formed on the 29 October 1945. Fifty years on the aspirations of the founders have been amply fulfilled. This expanded reference edition traces the 50 year history of the association at both Federal and State Division levels as well as cataloguing the gemstone resources of each State in Australia. The survey includes biological gem materials such as pearl production in Western Australia and the Coburg Peninsula in the Northern Territory. The compact and comprehensive nature of this edition does not lend itself to conventional abstraction, and the reader is recommended to consult the original.

Instruments and Techniques
El conoscopi figueras 93.

Evaluation of a new model conoscopic computer program as an instrument in research.
I. Vesselinov. Geochemistry, Mineralogy & Petrology (Bulgarian Academy of Sciences), 29, 1994, pp 97-105.

The commercially available program of E. Dowty is shown to be a useful tool in solving morphological problems in mineralogy. Three groups of questions are discussed and exemplified: (1) identifying crystal forms on photographs of real crystals (e.g. zircon, monohydrocalcite and pyrite); (2) reproducing crystal cross-sections (e.g. pyrite), and (3) simulating crystal growth by successive growth forms (e.g. zoned arsenopyrite).

Jewellery
Joaillerie indienne. 5000 ans de tradition.

Several important pieces of Indian jewellery are illustrated in a short survey of the craft over the last 5000 years.

Synthetics and Simulants
'Ti-sapphire': Czochralski-pulled synthetic pink sapphire from Union Carbide.

The pink 'Ti-sapphire' commercially available from Union Carbide by the Czochralski-pulling technique is used in laser applications as it produces a high-energy light over a wide range of frequencies in and near the visible range. It has also been made by flame fusion and melt techniques. Its gemmological properties are compared with natural and with other synthetic pink sapphires.

Separation from natural sapphire may be difficult but differences include no luminescence to long-wave UV radiation, chalky blue luminescence to short wave UV radiation (due to octahedral Ti\(^{4+}\)O\(_6\) impurities, an interstitial oxidised titanium) and pinpoint inclusions some of which were resolvable as gas bubbles. However reliable separation can be done by means of its chemistry or UV-visible absorption spectrum. Natural pink sapphires owe their colour to...
chromium, but with the 'Ti-sapphires' the colouring agent is trivalent titanium and a faint but different spectrum is seen with a spectro­scope (the main difference is the lack of chromium lines in the red). R.J.P.

**Lab notes.**


An imitation emerald necklace gave a chalky yellow reaction to long wave UV radiation (not unexpected from a heavily oiled stone) and appeared dullish red through the Chelsea colour filter; in the drill hole of one bead a dark green ring was observed containing flattened gas bubbles and it proved to be a natural beryl bead with a green plastic coating. A fine green carving gave a spot RI of 1.56, a weak mottled greenish blue fluorescence to short wave UV radiation and a spectrum showing general absorptions from 400 to 480 nm and 650 to 700 nm. The presence of gas bubbles, flow lines, mould marks and minute conchoidal fractures indicated a glass. An imitation lapis lazuli which looked more like sodalite was shown to be dyed feldspar by tests which included X-ray diffraction. A red stone with RI 1.552 and SG 2.64 and an orange stone with RI 1.480 and SG 2.39 imitating Mexican opal were examined. Both were singly refractive but gave anomalous effects in crossed polars in the form of snake-like bands. Both were inert to long wave radiation but gave a faint chalky yellow to short wave UV radiation. Magnification revealed small gas bubbles in both stones. EDXRF analysis revealed the presence of both selenium and cadmium, already known as constituents in orange and red glasses. A strand of silvery dark grey to black pearls was shown to consist of lead glass beads coated with bismoclite, a bismuth oxychloride which formed silvery layers (with elastic properties) and an outer layer of a transparent rubber-like substance containing gas bubbles.

A large Czochralski-pulled ruby weighing over 100 ct was transparent with a strong red colour with no purple or brown components. Some fine curved striae were seen but apart from these it was free from inclusions and colour variations. The manufacturer is reported to be aiming at pure crystals above 500 ct. R.J.P.

**Gem news.**


An unusual colour-zoned synthetic amethyst from Russia showed exceptionally uneven ‘speckled’ or ‘leopard’ spot colour distribution attributed to the fact that the crystal grew on a seed plane cut perpendicular to the c-axis with darker-colour regions forming expanding rhombic pyramids. Tanzanite-coloured synthetic sapphire has been sold in India as tanzanite. Typical constants for corundum and obvious curved colour banding and pin point inclusions which were probably gas bubbles enabled easy identification. A constructed quartz containing both natural and synthetic quartz formed a multi-coloured gem and included a rutilated quartz ‘termination’. R.J.P.

**Synthetic red beryl.**


Internal features and spectra in the range from UV to visible and the near infrared region are reported for a purplish red synthetic beryl. Absorption due to Mn is remarkable. I.S.

**So-called AGEE emeralds.** (In Japanese)


Internal features and spectra in the range from UV to visible and the near infrared region are reported for a purplish red synthetic beryl. Absorption due to Mn is remarkable. I.S.
Synthetic emeralds appeared on the market a few years ago under the name AGEE, and were supplied from AG. Japan Ltd. Gemmological investigations indicated that there were several types; one type has the same gemmological properties as those of Biron synthetic emerald, other types showed similarity to Russian synthetics, flux-grown synthetics, and vanadium beryl.

I.S.

Fraud at the source.

Useful account of various substances offered as natural gem materials from Namibia, Zambia and Zimbabwe. Most substitutes are glass of one form or another. M.O'D.

Crystal chemistry of non-stoichiometric Mg-Al synthetic spinels.

Twenty-seven single-crystal structural refinements were performed on synthetic spinels along the MgAlO2-Al2O3-3O2 join. The number of vacancies (from 0.09 to 0.22) was determined by EPMA. Vacancies are limited to octahedral sites, with a vacancy-to-oxygen distance refined to 2.113(3) Å. Mg is ordered in the tetrahedral site, only for a > 7.99 Å are the amounts of octahedral Mg significant. The decrease in a for vacancy-rich spinels is essentially due to the decrease of the T-O distance because of large amounts of tetrahedral Al. A linear relation between a and vacancy content was derived fitting literature data, including end-members. K.v.G.

[Observations on synthetic emeralds by the scanning cathodoluminescence (SCL) method]. (in Japanese with English abstract)

The internal inhomogeneities of ten faceted synthetic emeralds were investigated using scanning cathodoluminescence (SCL) methods. The origin of each specimen was established before SCL investigation by ordinary gemmological identification and by measurement of its NIR spectrum. The samples are from Lechleitner, Regency, Gilson, Saramandor and Chatham and include both flux and hydrothermal growths. Clear SCL images showed complex characteristics related to and depending on the methods of synthesis. Flux-grown emeralds are characterized by straight parallel growth banding or ‘aurora-like’ growth banding. Hydrothermally grown emeralds are characterized by the development of ‘comet-like’ luminaries arranged in one direction or unique rhombic patterns. SCL investigations are useful not only to distinguish between natural and synthetic emeralds but also between different synthetic methods. I.S.

Synthetische, nach dem Hydrothermalverfahren hergestellte Smaragde.

A hydrothermally-grown emerald is described and its inclusions illustrated. The specimen was grown at a temperature of 380°C.

M.O'D.

[Growth of emerald crystals by cooling the high-temperature solutions using PbO.V2O3 flux.] (In Japanese with English abstract)

Transparent green emerald crystals up to 2.2 mm long were synthesized from PbO.V2O3 solution, using no seed crystals. The nutrient was heated at 1200°C for 10 hours, followed by cooling to 600°C at rates varying between 5 and 240°C/h to find the optimum conditions. The size of crystals decreased with increase in the cooling rate. The optimum cooling rate for obtaining larger crystals was 5°C/h.

I.S.

[The present status of synthetic rubies.] (In Japanese with English abstract)

General characteristics, UV spectra, internal features, inclusions, growth sector patterns and growth banding of 'Rubiate' synthesized by a flux method by Culture Crystal Inc., USA, 'Douros' synthesized by a flux method by O.E. Created Gems, Greece, and rubies supposed to be synthesized by hydrothermal methods in Russia are described. Growth banding seen in flux grown rubies can be mistaken for banding seen in natural rubies. Hydrothermally synthesized rubies are generally dark red in colour and contain many flaws. Their refractive indices are significantly higher than those of natural rubies. Internal features revealed by laser-beam tomography are also described.

I.S.
**BOOK REVIEWS**

**Blood stones [a tale].**

Books on 'how the diamond market can be overturned' are not too uncommon and here is quite a readable tale whose thinly-disguised protagonist's battle with Russian diamond magnates over a possible move of Russian sales. Featuring as a prop to the complicated intrigues is a necklace of blood-red diamonds: whether or not they have been colour-enhanced forms an important part of the plot but I did not find out the answer! M.O'D.

**Norsk Steinbok.**

Norway has a rich mineralogy and a number of the species found have an ornamental application: these include emerald, peridot, sunstone and other gem-quality feldspar group members and some attractive colour-change and dark red garnets. The present book is arranged in traditional chemical order (beginning with elements), each entry including chemical composition, crystal system, common form and habit, mode of occurrence, and details of the particular site of recovery. Both mineral and rock species described are illustrated in colour, with welcome size details, and the quality of reproduction is good. There is a useful glossary, a bibliography and an identification table. While not everyone may be able to read Norwegian, the text is not too difficult for English-speaking students and in any case the photographs are sharp and chosen to be instructive. M.O'D.

**Cornish mineral reference manual.**

Many gemmologists are mineral collectors too and this concise and well-produced guide to one of the world's best-known mineralized areas will be useful to them and also to those beginning to perceive a world a little away from faceted gemstones. After a short introduction to the mining history of Cornwall and a bibliography, minerals are described in alphabetical order with composition, details of type specimens where known and of major analytical work performed on the species. Blank sheets at the end allow for additions. M.O'D.

**Fancy-color diamonds.**

'Fancy-coloured diamonds were virtually unknown to the general public and ... I wanted to say "wow"!' Harvey Harris is a diamond dealer specializing only in coloured diamonds and through his book, containing around 150 coloured photographs (many by leading gem photographer Tino Hammid), seeks to share his passion with the reader. The book is intended as a connoisseurs' guide to fancy-coloured diamonds, and is meant as much for enjoyment as for guidance. The diamonds illustrated are not simply fancy-coloured stones - they are the most intensely coloured prize-specimens from collections around the world and one's reaction is indeed 'wow'!

Before 1970 fancy-coloured diamonds were of little importance in the gem trade but, by the end of 1991, 19 of the top 20 price-per-carat records were held by fancy-coloured specimens. In the Introduction and in Chapter 9, the author chronicles some of these auction records, including the US$800 000 paid for a 0.95 carat round fancy purplish-red diamond in 1987.

The first two chapters of the book are largely historical, dealing with the evolution of the fancy-coloured diamond market and coloured diamond production through the ages - the deep blues and pastel pinks from India and Brazil, the Cape yellow stones from South Africa and, lately, the pink and brown diamonds from Australia. Harris also believes that Brazil was the first significant source of green diamonds including, perhaps, the Dresden Green. The histories of other fancy-coloured diamonds are mentioned briefly and it is emphasized just how
rare these strongly coloured diamonds are. Some are, indeed, the 'Old Master' of the gem world and, in an era when collectors are prepared to pay huge sums for unique works of art and unique works of Nature, this explains why prices of coloured diamonds have escalated a hundred-fold or more over the last 30 years.

Chapter 3 outlines the understanding and causes of colour in diamond and contains some excellent photographs. It clearly emerges that, while some colours (blue, green, yellow) are well-understood, there are no detailed explanations yet available for many other colours (brown, orange, pink). The specialist reader will find the treatment in this chapter, whilst mostly correct, rather shallow, whereas the general reader will encounter many unfamiliar concepts inadequately explained (especially the details and meaning of the various absorption spectra presented). However, an important point to emerge is that there are laboratories able to analyse the colours of diamonds and which are continually striving to understand in greater depth the origins of these colours. This information allows gem-testing laboratories to differentiate between naturally and artificially coloured diamonds (described in chapter 5) and between natural and synthetic diamond (described in chapter 6).

Chapter 4 presents interesting material on the cutting of fancy-coloured diamonds. A brilliant-cut, for example, washes out light-to-medium fancy-colours, while other shapes are devised to accentuate the colours of the stones.

The author is highly critical (Chapter 7) of the methods used by the GIA for colour-grading fancy-coloured diamonds. They use a subjective comparison technique and the lack of an adequate range of master stones can lead to diamonds with quite different colours being given the same description. Harris argues strongly for coloured diamonds to be measured objectively on equipment like that currently used by the Gübelin Gemmological Laboratory; it is claimed that this gives reproducible numerical values for the hue, tone and saturation which are the three key parameters in describing the colour of a gemstone.

The book has a Preface by François Curiel of Christie’s and a Foreword by George Bosshart of the Gübelin Gemmological Laboratory. It has a very readable style, although the eye is continually diverted from the text by the magnificent illustrations, and there are relatively few misprints or layout errors in this first edition. Harris is not a scientist and, although he has consulted several diamond experts, the text contains a number of minor technical errors. Furthermore, readers who are not diamond dealers will probably find the author’s obsession with the price-per-carat, of each stone described, rather tedious by the end of Chapter 10. Notwithstanding these criticisms, this is a book which anyone interested in coloured diamonds should have on the bookshelf.

A.T.C.

Rare earth minerals. Chemistry, origin and ore deposits. (Mineralogical society series.)

While gemmologists will inevitably associate rare earths with the attractive colours and absorption spectra produced in synthetic crystals when RE are used as dopants, these interesting and important minerals have a life outside gemmology. The present work consists of papers selected from those presented at the conference Rare Earth Minerals: chemistry, origin and ore deposits, held at the Natural History Museum (London) in April 1993.

The first chapter, introducing and reviewing the rare earth elements, should be required reading for serious gemmologists. For those who may not have a chance of reading the book, the rare earth elements [REE] form Group IIIa of the Periodic Table, that is scandium followed by yttrium and lanthanum through to lutetium. The name lanthanides is used for the 14 elements from cerium to lutetium. These names often crop up in mineralogy texts read by gemmologists who should establish their meaning.

After the introduction the chapters are devoted to RE minerals, their composition, formation and properties. Many of the species form crystals which are attractive to the mineral collector and readers will note that the papers are grouped into deposit types: these include carbonate-rich melts, hydrothermal systems, syenite pegmatites, peralkaline systems, carbonatites, laterites, karst-bauxites and karstic nickel deposits; other chapters cover RE deposits in China, in the Kola Peninsula, Russia, an analysis of RE minerals, and a revised classification scheme for an important RE host in alkaline rocks. There is a glossary (in this case a list of minerals with their composition and arranged in
chemical order). Chapters carry their own lists of references which are comprehensive in all cases.

With first-class illustrations in colour and well-presented tables and diagrams the book provides as useful an introduction to RE minerals as can be found.

Cameos old and new.

Opening with an account of the history of the glyptic arts, followed by a discussion of the commoner cameo subjects, the book goes on to describe how cameos are made and how the collector may distinguish old from new artefacts. The various cameo imitations are described and there are suggestions on how a collection may be formed. Major collections are cited and there are useful sections comprising a bibliography, addresses of major auction houses, retail outlets and booksellers (confined to the United States). There is also a ‘chronology of engraved stones and cameos in Europe’. The illustrations are placed together in the centre of the book.

While this is a perfectly reasonable structure and on the whole produced quite well, I found it hard to decide whether the book aimed at serious scholarship or whether it was merely a compilation of hastily-assembled facts. Some statements in the materials section are questionable: chalcedony will not disclose its cryptocrystalline structure under the microscope. Although some facts need investigation, as a general guide the book succeeds quite well; it is easy to read and well-illustrated. The writer’s interest comes through strongly and perhaps she could be described in her own words for Maxwell Sommerville as ‘a curious and trusting scholar’.

Jade.

It seems impossible for so much information to be packed into so small a compass, especially when the extensive and excellent picture coverage is included, but Fred Ward has done it again, this time with a notoriously difficult topic. A superbly coloured cabochon of Imperial jade arrests the eye as soon as it settles on the front cover and the same standard persists throughout the book which must be required reading for all sorts of gemmological and general readers.

The use of the jade minerals in different civilizations is covered first and the book then looks at the location and recovery of specimens, passing then to methods of working and to brief notes on valuation. Look out for the pictures of lavender jadeite, of jade from New Zealand and from Myanmar. The book is a delight and can stand with the already extensive jade literature without any loss of face.

Antwerp gemmological update.

Topics discussed include an exhibition celebrating the opening of the ‘Treasury’ in the Royal Museum of Arts and History and the Royal Museum for Central-Africa, Brussels, the exhibition including gemstones: inclusions in gemstones: orange ‘mandarin’ garnet from Namibia: details of diamond jewellery courses scheduled to be held during 1996.

Christie’s jewellery review 1995.

Major items sold by the firm during 1995 are illustrated in colour and described, with notes on prices paid. A number of yellow diamonds are included and the review also contains a short account of the work of the De Beers group of companies and an introduction to blue diamonds.
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OBITUARY

Ted Thomson, Chairman of E.A. Thomson Gems Limited, died on 2 December 1995 aged 80. He will be remembered as the man who put more into the Gem Trade than he took out.

Ted, born on 2 March 1915, started his career by working for an insurance company after graduating from the Stationers School in North London. He decided that insurance was not for him and found a job with George Lindley & Co., gem dealers, a company known at the time as the Jade Dragon. It was there that Ted gained an interest in gemstones until the war interrupted his career. He joined the Royal Artillery in 1939 and rose up through the ranks to Captain. On demobilization he returned to his job with George Lindley where he worked until 1958 when he left to start up his own gem dealing company. His co-director, Eric Allen, joined shortly afterwards and together they built up the international business which constitutes E.A. Thomson Gems Ltd.

In 1971 E.A. Thomson Gems Ltd acquired the Birmingham stone dealing company of Morris Goldman Ltd and, predicting Great Britain’s involvement in the EEC, an office was opened in Paris.

Not content with building a successful gemstone business, Ted devoted himself to enhancing the reputation of the trade. He served on the committee of the Diamond, Precious Stone and Pearl section of the London Chamber of Commerce and Industry, becoming Chairman of that section in 1984. Apart from dealing with many trade issues, liaising with the Treasury and HM Customs and Excise, the section also had the responsibility of running the British Gem Testing Laboratory, which subsequently merged with the Gemmological Association to form the GAGTL.

During his period of office, Ted was energetic in his promotion of the Laboratory and can claim credit for securing a contract with the State of Bahrain. The British Gem
FORTHCOMING MEETINGS

London
Unless otherwise stated, meetings will be held at the GAGTL Gem Tutorial Centre, 27 Greville Street (Saffron Hill entrance), London EC1N 8SU. Entry will be by ticket only at £3.50 for a member (£5.00 for non-members) available from the GAGTL.

8 May  Identifying inclusions within gemstones          Jamie Nelson
5 June  3-D slide presentation of gemstones and minerals  David Minster
10 June Annual General Meeting, Reunion of Members and Bring and Buy
         (There will be no charge for this event)
         The speaker will be Nairn Attallah, former Chief Executive of Asprey.

Midlands Branch
Monthly meetings will be held at the Discovery Centre, 77 Vyse Street, Birmingham 18 (for directions to the Sunday Gem Club venue contact Gwyn Green on 0121-445 5359). Further details from Mandy MacKinnon on 0121-624 3225.

26 April Annual General Meeting followed by a talk by D.H. Ariyaratna
28 April Preliminary Gemmology Seminar
5 May Diploma Gemmology Seminar
19 May Gem Club - visit to Dolgellau Gold Mine

North West Branch
Meetings will be held at Church House, Hanover Street, Liverpool 1. Further details from Joe Azzopardi on 01270 628251.

15 May Jade, past and present         Rosamond Clayton
19 June Jewellery in the auction world     David Lancaster

Scottish Branch
For details of Scottish Branch meetings contact Ruth Cunningham on 0131–225 4105.

19–21 April A weekend of Scottish gemmology     Alan Hodgkinson
A three-day event to be held in Perth to include talks and practical workshops by Alan Hodgkinson, the Branch’s first Annual General Meeting and a field trip to Ethiebeaton and Ardownie quarries near Dundee.
Testing Laboratory provided the equipment, trained the staff and seconded a senior staff member to set up what was to be the first laboratory in the Middle East working to international standards.

In addition to his duties at the LCCI, Ted Thomson was one of the founder representatives to CIBJO (the International Confederation of Jewellery, Silverware, Diamonds and Precious Stones). He was elected President of the Coloured Stone Commission in 1979 and served two terms in office. During this time Ted worked tirelessly on the production of the CIBJO Gemstone Book - the first definitive manual containing nomenclature and rules of application for the gem trade which, when combined with the Diamond and Pearl Books, became known as the Blue Book. Ted applied himself selflessly to the task and was not averse to telephoning Alec Farn, or latterly Ken Scarratt, at some unearthly late hour to clarify some arcane point regarding nomenclature. His charming approach never brought down the wrath of the awakened respondent. Ted’s charm stood him in good stead when it came to the CIBJO Annual Conference. He would lobby all the major participants prior to the meeting to iron out any difficulties, and made sure of a clear path for his carefully prepared resolutions.

Ted prided himself on his knowledge of the Statute Book – an asset which would have an unprepared opponent slinking away in shame when confronted by chapter and verse. It was largely through Ted’s drive and conference skills that the important second edition of the Blue Book was produced in 1982.

Apart from his skill in business and trade politics Ted also had an intense interest in gemmology on a personal level. As a tribute to his enthusiasm for gemmology and in recognition of his achievements in the trade, Ted was awarded an honorary FGA, one of only two non-academics to be granted this honour. He was invited to present the diplomas at the annual reunion of members of the Gemmological Association on 15 November 1982. By happy coincidence, one of the students to receive her award that evening was his daughter-in-law, Jenny Thomson.

Tragedy struck Ted when his wife of 44 years, Beryl, died after a long illness in 1983, after which Ted gradually handed over the running of the business to his two sons.

Although he was the head of a large company and an important official of several trade bodies, Ted Thomson never lost that common touch which earned him the respect and affection of his family, his loyal staff, his associates in the gem trade and all those people who had the good fortune to know him. He is survived by his second wife, Tuu, and his two sons, Ian and Peter.

George Lindley, FGA (D. 1937), died on 27 November 1995 aged 90. Born in Melitopol in 1905 of Russian-American parents he saw the early days of the 1917 Revolution while at school in Russia. At the age of 12 he made his own way, via Beijing, to the USA where he completed his education.

His early years in China inspired the love and knowledge of jade which became his life’s work, accentuated by his appointment in 1926 as the buyer in Beijing for a major New York jade merchant. By 1929 he had resolved to open a business in London dealing in jade for jewellery, and as carvings and objets d’art. This flourished, despite the difficult trading conditions of the time, and he was ably partnered from 1935 by his late wife, Carol (née Melville) (D. 1937).

The company was joined in 1936/37 by E.A. Thomson and W.C. Buckingham but business was inevitably slowed in the years immediately prior to the second World War. George Lindley served in the Auxiliary Fire Service in London, 1940/42, then the RAF Volunteer Reserve (Pathfinder Squadron) until 1945, when he restarted the business, being joined again by E.A. Thomson and W.C. Buckingham.

By now the work of the company had turned to coloured gemstones, in which George Lindley was a pioneer in careful
grading and calibration of original parcels, specifically for a demand generated by the centrifugal casting process. This gave the company a recognized place in the UK and international markets.

There was a steady flow of staff through the company in these years. Almost all attained their FGA. Some are still active with their own successful companies in the trade and would acknowledge the stern, but detailed guidance they had from George Lindley.

W.C. Buckingham

Mrs Deidre M.H. Inches Carr, MA FGA (D. 1982), Edinburgh, died on 12 December 1995.

Mr Juan C. Paredes Quevedo, FGA (D. 1994), Madrid, Spain, died in 1995.

GIFTS TO THE ASSOCIATION
The Association is most grateful to the following for their gifts of gems and gem materials for research and teaching purposes:

Dr J.L. Emmett, Pleasanton, California, for an illustrated summary of a lecture delivered to the GAGTL entitled 'The heat treatment of corundum: a view from the laboratory'.

Mrs Theresa Khairallah, London, for samples of amber, cornelian and coral.

Mrs Eynat Matza, London, for a specimen of eilat stone.

Professor E. Ralph Segnit, Melbourne, Australia, for a specimen of Opalus, an imitation opal from Melbourne.

D. Swarovski & Co., Wattens, Austria, for over 900 cut gemstones of various natural and synthetic material.

MEMBERS' MEETINGS
London
On 4 January 1996 at 27 Greville Street, London EC1N 8SU, Dr Grahame Brown from Albany Creek, Queensland, Australia, gave an evening lecture entitled 'Gems stones - Australia's national treasure'. On the following day Dr Brown led a half-day seminar entitled 'Hand lens characteristics of biological gem materials'.

On 28 February at 27 Greville Street Rosamond Clayton gave a lecture entitled 'The art and science of valuation'.

Midlands Branch
On 26 January at the Discovery Centre, 77 Vyse Street, Birmingham 18, members enjoyed a Bring and Buy, Quiz Night and Raffle.

On 23 February at the Discovery Centre E. Alan Jobbins gave an illustrated talk on jades.

On 4 March a visit was arranged for members to the Birmingham Assay Office.

On 29 March at the Discovery Centre Edgar Taylor gave a talk entitled 'Lavriotike: treasure house of Attika'.

Gem Clubs were held on 21 January, 18 February and 24 March.

North West Branch
On 20 March at Church House, Hanover
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Street, Liverpool 1, Deanna Brady gave a talk entitled ‘Introduction to inclusions’.

Scottish Branch
On 7 February a Lab Night was held at Telford College, Edinburgh.
On 22 March at Newliston House, Newbridge, Edinburgh, members enjoyed a quiz and Bring and Buy sale.

GEM DIAMOND EXAMINATIONS
In January 1996 12 candidates sat the Gem Diamond Examinations in the UK of whom 8 qualified. The names of the successful candidates are listed below:

Atkin, Marie, Sheffield.
Cauchi, Savona, Louis, Malta.
Crowder, Michael J., London.
Gartery, Tania M., Christchurch.
Hill, Katia, Glasgow.
Johnson, Christopher W.R., Bickley, Nr Bromley.
Ryder, Peter Ian, London.

The overseas candidates who qualified will be published in the July issue of the Journal.

EXAMINATIONS IN GEMMOLOGY
In the Examinations in Gemmology held in January 1996 183 candidates sat for the Preliminary examination of whom 147 qualified; 116 candidates sat for the Diploma examination; 50 have qualified so far. The additional overseas candidates who qualified will be published in the July issue of the Journal.

The names of the successful candidates are as follows:

Qualified with Distinction
Grondin, Daniel, Geneva, Switzerland.

Qualified
Abramian, Levon, London.
Ahamed, Omar Faiz, Gillingham.
Ang, Zen Yi, Singapore.
Ayyar, Rajeshwari Narayan, Bombay, India.
Brangulis, Peters, Riga, Latvia.
Close, Helen Jane, Selby.
Cunningham, DeeDee Cecilia, Toronto, Ont., Canada.
Dempster, Stuart, Glasgow.
Fleischner-Zois, Krista, Athens, Greece.
Golad, Tanya, London.
Grondin, Magali, Geneva, Switzerland.
Horniblow, Kathy, Sherbourne.
Hyo-Jin, Choi, Tae-gu, Korea.
Jayasendil, J., Salem, India.
Kim, Yun-Hwa, Kyung Buk, Korea.
Kimber, Sarah, London.
Kuulman, Heli, Tallinn, Estonia.
Lake, Richard John, Gorey Village, Jersey.
Limaye, M.D., Calcutta, India.
McFarlane, Iain, Chesham.
MacNish Porter, Holly, Edinburgh.
Marr, Peter, Torquay.
Mathon, A.P.T., Oosterhout, The Netherlands.
McFarlane, Iain, Chesham.
Melego Canet, Merce, Valencia, Spain.
Michelson, Max J., Lincoln.
Mihama, Saki, London.
Naing, Aye Myo, Yangon, Myanmar.
Oja, Tonis, Tallinn, Estonia.
Parekh-Guastella, Rita, Pontedera, Italy.
Perkins, David J., Ayr.
Pierce, Jason Robert, Truro.
Pierce, Jill Fiona, Manchester.
Plomp, Jean Victor, Rotterdam, The Netherlands.
Popov, Vladimir V., Tallinn, Estonia.
Prabhu, Miss Vaishali, Bombay, India.
Rai, Ashish, Jaipur, India.
Rege, Sonal, Bombay, India.
Shepherd, Louis, Nottingham.
Song, Gyu Bong, Ulsan, Korea.
Stopelli, Christopher J.D., Oakville, Ont., Canada.
Torrent, Denise, Geneva, Switzerland.
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van der Molen, W.N., Schoonhoven, The Netherlands.
Van Duijnen, A.M., Bergen aan Zee, The Netherlands.
Vermi, Hitesh, Jaipur, India.
Vummidi, Nagaraj, Madras, India.
Withington, Terry, London.

Preliminary

Amrute, Rahul M., Bombay, India.
Andaluz Sanchez, Maria, London.
Anderson, Elizabeth Anne, Lewes.
Ayyar, Rajeshwari Narayan, Bombay, India.
Bagadia, Pavan Madanlal, Bombay, India.
Barbas, Constantinos, Athens, Greece.
Burden, Lorraine, Sawbridgeworth.
Cao, Shumin, Wuhan, China.
Chan, Hor Wai Pierre, Kowloon, Hong Kong.
Chan, Vivian S.F., N.T., Hong Kong.
Chan, Lam Ming, Hong Kong.
Chen, Kai-Wen, Taipei, Taiwan.
Chong, Yung Yung Sylvia, Toronto, Ont., Canada.
Chung, Wen-Hao, Taipei, Taiwan.
Coker, Olive Admira, London.
Cowley, Jacelyn Gloria, London.
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Davies, Darryn Lewis, London.
Dismore, Frances Jane, London.
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- Chu-Li, Wu, Edgbaston. 1995
- Dang, Xiao-Ying, Beijing, China. 1995
- Heyes, Alan John, Stourbridge. 1970
- Hui, Wai Yee Wendy, Hong Kong. 1994
- Lounghi, Jagdish, Jaipur, India. 1995
- Maheshwari, Ashoo, Jaipur, India. 1995
- Marathe, Tanuja, Pune, India. 1995
- Salm-Reifferscheidt, Sophie, London. 1995
- Stu, Ming Wa, Kowloon, Hong Kong. 1995
- Tang, Zhen Yi, Kowloon, Hong Kong. 1995
- Rossiter, John Timothy, Weston Super Mare. 1995

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Transfers from Ordinary Membership to FGA
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Yen, C.K., Taipei, Taiwan.
Corrigenda

In the *Journal of Gemmology*, 1995, 24(8), pp 543 and 544, the author was unfortunately inconsistent when drawing Figures 3 and 4 in that in Figure 3 he showed the pearl in cross section, while in Figure 4 the pearl is spherical, although shell and mollusc are drawn in cross section. The following Figures 3a and 4a now display the correct structure in the pearl consisting of the conchin nucleus, enveloped by the primary layer of calcite, coated by calotte shaped layers of aragonite platelets – all in cross section.

![Fig. 3a](image1)

![Fig. 4a](image2)
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