

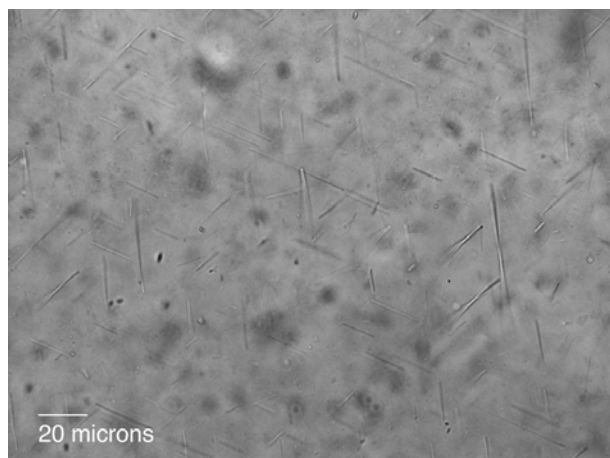


Figure 6a. Oriented fibres with granular and pinpoint inclusions in blue rose quartz.

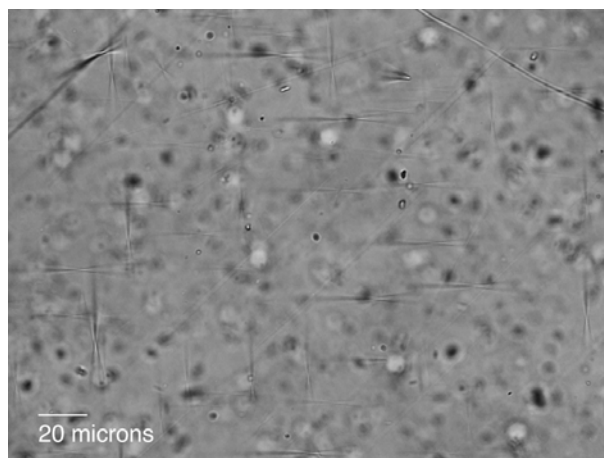


Figure 6b. Oriented fibres and granular inclusions in pink rose quartz.

Discussion and further investigation

The presence of oriented fibres in this blue rose quartz, along with hints of pink colour in some specimens, prompts comparison with the dumortierite-like fibres identified by Ma *et al.* (2002) as the cause of colour in pink rose quartz. The overall hazy blue colour with a transmitted yellow colour, plus the Tyndall scattering effect that makes the laser beam visible (Kerker, 1991), combined with the observed high density of almost invisible pinpoint inclusions, all suggest that the dominant blue is due to scattering phenomena as postulated by Seifert *et al.* (2011) for other examples of blue quartz. The similarity of the dichroism shown by the blue rose quartz to that in the standard rose quartz is consistent with reports that “even rather

pale rose quartz may show medium to strong dichroism” (Liddicoat, 1989). However, since scattering phenomena polarize light (Strutt, 1919), the relative contribution of dichroic fibres versus scattering to the dichroism requires clarification. The muddy colour of this blue rose quartz may be linked to the yet-to-be-determined size distribution of pinpoint and granular particles present. Particles below and at the resolution of the microscope may contribute the blue, while larger ones may scatter white light or even act as absorbers. Beyond colour, the eye-visible fracturing and the nature of the quartz microstructures observed in the thin sections opens avenues for understanding temperatures and strain states associated with the deformational history of the deposits that sourced these quartz specimens.

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Gem-quality pink fluorite from Planggenstock, Switzerland: chemical and physical properties and a comparison with pink fluorites from other localities

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Introduction

In this study, first results of trace element analyses and various spectrometric investigations on pink fluorites from alpine deposits (Aar and Montblanc Massifs) as well as from the skarn de-posit near Huanggang, China, will be presented.

Pink fluorite is a typical mineral from Alpine extensional fissures, especially from the granitic central massifs of the Alps Aar Massif (Switzerland) and Mont Blanc Massif (France) (Stalder *et al.*, 1998). It occurs exclusively in the form of octahedral crystals of usually less than 3 cm edge length, but very large crystals of over 15 cm are rarely found. The colour ranges from mostly pale pink to pinkish red and rarely intense raspberry red.

The formation of alpine pink fluorites occurs in connection with the formation of the minerals in alpine crystal clefts. The formation of these alpine fissures was caused by tectonic processes during the Alpine orogeny (Gnos *et al.*, 2021). At a depth of about 13 to 18 km within crustal rocks and at temperatures of 450 to 550 °C, the tectonic stress about 20 My ago led to the for-mation of extensional fissures in the rocks, with cavities reaching up to several metres in size. These cavities were subsequently filled with supercritical hydrothermal aqueous fluids, which dissolved minerals from the rock of the fissure walls. The cooling associated with the uplift of the Alps led to supersaturation of the dissolved complexes and thus to the hydrothermal for-mation of minerals in these fissures. Quartz, the most common mineral species in these Alpine clefts, started crystal formation at temperatures between about 450 and

400°C, which represents the earliest stage of crystallisation. In contrast, the crystallisation of fluorite and chlorite began at lower temperatures in the final stage of growth of the rock crystals. This is why fluorites always are found grown on the surface of quartz crystals.

Probably the largest find of alpine pink fluorite to date was made in the fissure system of the Planggenstock mountain, canton of Uri, central Switzerland. This deposit is located in the so-called Central Aar Granite, which, with an extension of about 90 km from west to east, is the largest body of granitic rock in the Aar Massif and in Switzerland. The crystal deposit was dis-covered in 1993 and contained several very large fissure cavities which yielded mainly rock crystal in crystals up to one metre in size. In 2019, another cavity was discovered nearby containing numerous intensely coloured and large fluorite crystals (Fig. 1 and Fig. 2) together with large amounts of green chlorite sand.

Trace elements analysis of pink fluorite

At the time of submission of this abstract, trace elements of pink fluorites from Planggenstock and Huanggang, China have been analysed using Gem-TOF (LA-ICP-TOF-MS) at SSEF. Pink fluorite samples from the Montblanc Massif (France) are currently prepared and their data integrated later.

The most abundant trace element in pink fluorites from both Planggenstock and China is yttrium, which was measured in concentrations of 170 to 278 ppm in the alpine samples and between 205 and 273 ppm in the fluorites



Figure 1: Octahedral crystals of pink fluorite on granite from the Planggenstock mountain, Switzerland. Total height approx. 30 cm. Photo: M. Hügi

from Huanggang. Based on our samples, a clear distinction between the alpine and Chinese pink fluorites is exhibited by the varying concentrations of Sr and Lu. For both elements, the alpine samples exhibited higher mean concentrations, with factors of 3 for Sr and 8 for Lu.

Despite the divergent geochemical formation conditions, the REE patterns of both deposits demonstrate remarkable similarity in terms of concentration ranges and the ratio of light REE (LREE) to heavy REE (HREE) (Fig. 3). The REE pattern exhibited by the Chinese samples deviates from that observed in the alpine samples, and is characterised by a negative Eu - anomaly. This discrepancy can be attributed to the divergent redox conditions that prevail in the respective geochemical growth environments, as generally described by Armbruster *et al.*, (1996).

The LREE / HREE - ratio measured in the alpine fluorites also aligns with the observation by Armbruster *et al.*, (1996). These authors reported that the REE patterns of alpine fluorites and the host rock (Central Aar Granite) are contrasting, which the granite showing a notably higher LREE/HREE ratio and a slight negative Eu anomaly (Schaltegger and Krähenbühl, 1990) compared to the fluorites. The enrichment of HREE in pink fluorites can be attributed to a post-magmatic fluid-rock interaction with F-bearing fluids, which have the capacity to form complexes of HREE, in conjunction with Na, Nb, Y, Zr, Hf, Ta and U (Webb *et al.*, 1985).



Figure 2: Intensely coloured fluorite of 97.18 ct from the Planggenstock mountain, cut by Philipp Munsteiner. Photo: Atelier Munsteiner

Cause of colour of pink fluorites

According to Bill and Calas, (1978), the attractive pink colour of fluorite is due to the presence of a YO^{2+} colour centre. This assertion is supported by UV-Vis spectrophotometry, which confirms the main absorption at 485 nm, and by the high Y concentrations that have been measured on samples both from Planggenstock and Huanggang. A weaker absorption band at 365 nm is attributed to Yb^{2+} (Armbruster *et al.*, 1996).

Some fluorite crystals from the Planggenstock show a distinct inhomogeneous colour distribution gradually ranging from intense pink to colourless. Based on our GemTOF analyses, this colour distribution is not related to variations in Y concentration. So it can be assumed that the YO^{2+} colour centres may have been activated differently due to variable exposure to natural radi-oactive irradiation.

Inclusions in alpine pink fluorite.

So far, microscopic inclusion analysis was conducted on the polished pink fluorites from the Planggenstock deposit. The inclusion pattern demonstrates the presence of two-phase fluid inclusions, which are characteristic of hydrothermal mineral formations found in the alpine fissures of the Aar massif (Mullis, 1996). These inclusions are euhedral tetrahedral negative crystals in shape and are characterised by an aqueous filling with a water vapour bubble (Fig. 4). The most prevalent solid inclusions observed on or in close proximity to the crystal faces of the fluorite crystals were layers of fine-grained chlorite, which crystallised as the final phase of mineralisation within the alpine fissure.

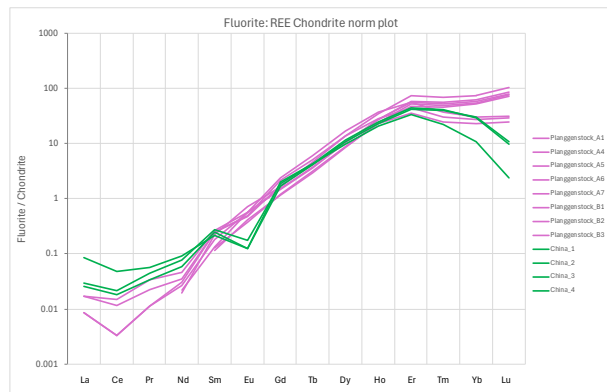


Figure 3: Chondrite-normed REE-pattern of pink fluorite of the Planggenstock mountain (red lines) resp. from the Huanggang Mine (green lines).

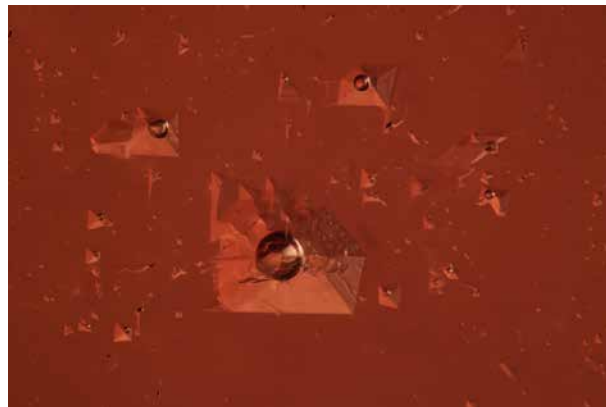


Figure 4: Two-phase inclusions in tetrahedral negative crystals in pink fluorite from Planggenstock mountain, Switzerland. Width of field 1.2 mm. Photo M. Hügi

Conclusions:

Although not a gemstone of great prevalence and value in the trade, fluorite, and especially attractive pink fluorite of gem-quality is considered an attractive collector stone and especially also sought after, even more so as rough crystals with or without matrix, due to their perfect octahedral shape.

With the discovery of the huge cavity with numerous pink fluorites at Planggenstock (Switzerland), and new pink flu-

orites emerging from Huanggang in China, it is the aim of this study to better characterise this new material from Planggenstock and to compare it with pink fluorites from Huanggang and the Montblanc Massif. Our preliminary results show that despite quite different geological setting and formation (Planggenstock: hydrothermal; Huanggang: skarn-related) their trace element concentrations and patterns only differ slightly.

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Typomorphic features of tourmalines of the Malkhan pegmatite field from the Irkutyanka vein

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This abstract presents the examination results of 5 tourmaline samples found in the Irkutyanka vein of the Malkhan pegmatite field in 2022 (Figure 1). The samples were studied using optical and scanning electron microscopy as well as LA-ICP-MS analyses. Conclusions are made on the chemical composition of the samples, noting the anomalous content of bismuth and lead in the studied samples.

The Malkhan pegmatite district is among the today active tourmaline deposits, which is the largest source of tourmaline in Russia. The deposit is located in the Krasny Chikoy area of the Zabaikalsky Krai, Transbaikalia, Russia. Geology, mineralogy, and geochemistry of the pegmatite field are described in detail by Zagorsky and Peretyazhko (1992). There are more than 15 tourmaline-bearing veins in the Malkhan pegmatite field including Danburitovaya, Geologicheskaya, Irkutianka, Karkadilovaya, Mokhovaya, Novaya, Oktyabrskaya, Oreshnaya, Skakunya, Solnechnaya, Sosedka, Svetlaya, Tabornaya, Zapadnaya-1 and Zimoveynaya (Zagorskiy *et al.*, 2008).

The Irkutyanka vein has been known since 2018. It is a wedge-shaped body up to 390 m in length and from 5 to 12 m in diameter. The productive part with gem quality tourmaline is represented by an albite and quartz-albite pegmatite. The sizes of miaroles with tourmaline vary widely and have a variety of shapes. The distribution of miaroles and tourmaline in the productive zone is extremely uneven. Tourmaline crystals vary from several mm to 5 cm in size. In addition to the traditional colored varieties of elbaite from Malkhan, we found monochromatic translucent crystals of a light green (mint) color.

The chemical composition of the crystals was analyzed on a TESCAN VEGA 3 scanning electron microscope at an accel-



Figure 1. Samples of the studied tourmalines from the Irkutyanka vein of the Malkhan pegmatite field. FOV 6.5 cm.

erating voltage of 15 kV. Average Li and B contents in each sample were determined using LA-ICP-MS measurements, carried out with an Elan DRC-e quadrupole inductively coupled plasma mass spectrometer. Laser ablation: NWR-213. Calibration was carried out using solid standard samples NIST SRM 610 and NIST SRM 620. The calculation of the elemental content was carried out with normalization to 100% of the sum for oxides.

The tourmaline supergroup has the generalized chemical formula $[^{9}X^{6}Y_3]^{6}Z_6 ([^{14}T_6O_{18}) ([^{3}BO_3)_3 V_3 W$ (Henry *et al.*, 2011) and can be classified into primary groups based on the dominant occupancy of the X-site. Twenty-one members of the tourmaline supergroup contain Na as the dominant cation on the X-site, seven Ca, six □ (vacancy) and only one K.

Occupation of structural positions in chemical formulas was carried out step by step in accordance with the procedure specified in the current nomenclature of tourmalines (Henry *et al.*, 2011). First, Na and Ca were placed in the

X-position. The fraction of a vacancy ($\square$) in this position was calculated from the stoichiometric ratio based on the equation $\text{Na} + \text{Ca} + \square = 1$ apfu. The assignment of a particular composition to a specific mineral species was carried out in accordance with the rules on the dominant valence approved by the KNMNK MMA (Bosi *et al.*, 2019). First, based on the predominant component in the X-position, the mineral was determined to belong to the group of alkaline tourmalines ($\text{Na} > \text{Ca}$ and $\text{Na} > \square$) or tourmalines with a dominant vacancy ($\square > \text{Na}$ and $\square > \text{Ca}$) (Figure 2). After that, the dominant anion in the W– F[–], OH[–] or O^{2–} position was taken into account. At the last stage, the dominant end-member determining the formula of the end-member of the corresponding mineral species in the

tourmaline supergroup was calculated based on the results of the Y-position occupancy.

All the examined tourmalines contain Li (0.76–0.94 apfu) and Al (7.76–8.11 apfu) in octahedral positions. Other octahedral cations (Mn, Fe, Ti) contain less than 1 apfu. Tourmalines do not show a deficiency of Si (5.90–6.00 apfu) and are represented by species with a predominance of F (0.78–0.83 apfu).

As a result, we have calculated the compositions of tourmalines that fall into the fields of the following two mineral species:

Fluor-elbaite $\text{Na}(\text{Li}_{1.5}\text{Al}_{1.5})\text{Al}_6(\text{Si}_6\text{O}_{18})(\text{BO}_3)_3(\text{OH})_3\text{F}$

Fluor-rossmanite $\square(\text{LiAl}_2)\text{Al}_6(\text{Si}_6\text{O}_{18})(\text{BO}_3)_3(\text{OH})_3\text{F}$

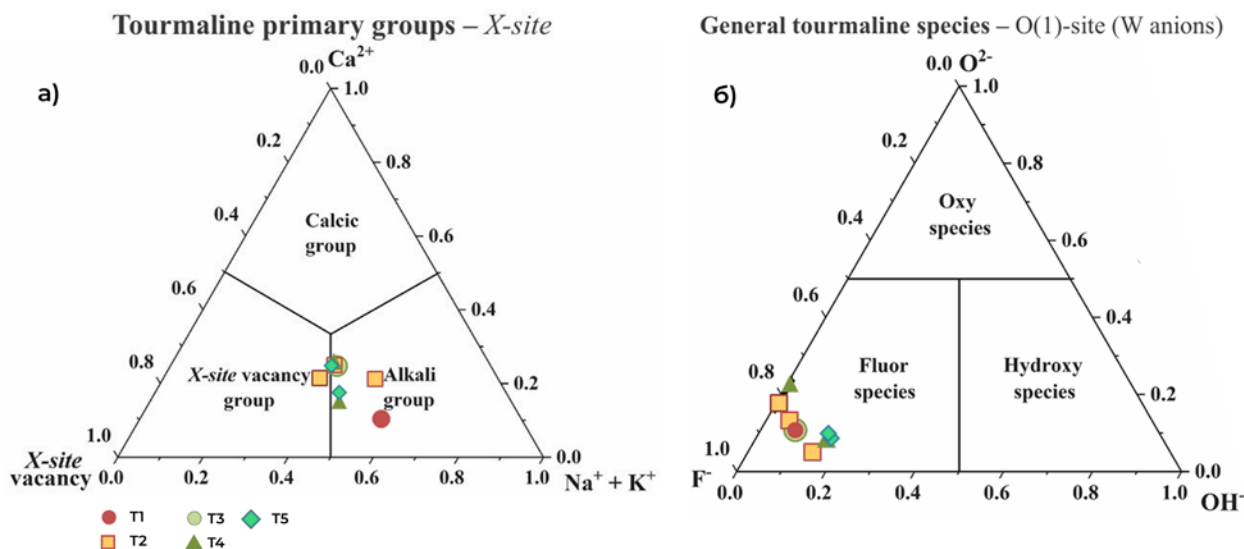


Figure 2.

a) The ratio of cations and vacancies ($\square$) in the X position in tourmaline supergroup minerals from the Irkutyanka vein,
b) The ratio of anions occupying the W position in tourmaline supergroup minerals from the Irkutyanka vein.

It is worth noting that elbaite and fluor-elbaite are common mineral species not only in the Malkhan pegmatite field, but also in other tourmaline deposits. Fluor-rossmanite is endemic to Malkhan and was approved as a new mineral species in 2022. The studied tourmaline samples were found to contain higher lead (up to 0.5 wt.%) and bismuth (up to 0.7 wt.%) contents (figure 3). The largest amount of lead was found in sample T1, and the largest amount of bismuth was found in samples T4 and T5. Bismuth and bismuth-containing minerals have already been described in separate veins on the Malkhan Ridge. The bismuth-columbite mineral was discovered in the Danburitovaya albite-pegmatite vein, and Bi- and Pb-rich tourmalines were found in the Zapadnaya-1 pegmatite vein (Peretyazhko

et al., 1989). The lead and bismuth content in our samples may be a typomorphic property. Vereshchagin *et al.* (2022) suggested in their article that trivalent cations can also occupy the X-position of tourmaline. It can be predicted that the general classification scheme for tourmalines based on the occupancy of the X-position will include not only vacancies, monovalent (e.g. Na, K) and divalent (e.g. Ca, Pb) cations, but also trivalent (e.g. Bi) cations. (Vereshchagin *et al.*, 2022)

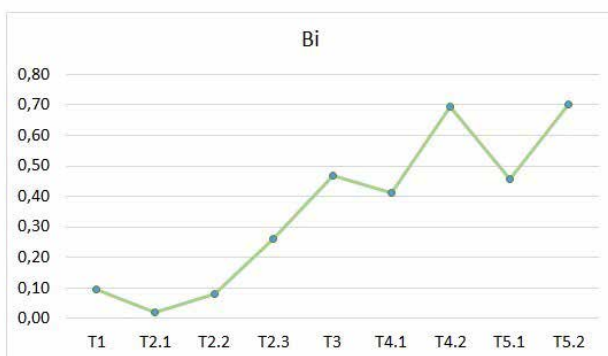


Figure 3. Bismuth content in the studied tourmaline samples in mass percent.

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Colored gemstones of metamorphic origin: Progress report in understanding how they crystallize at depth in East Africa, the Hindu Kush, Pamir Mountains, and the Middle-Urals Ring Structure

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Two problems

It is difficult to reconcile the existence of deposits of transparent gems of metamorphic origin with the observation attributed to Robert Boyle (1627–1691) that the best crystals grow in cavities. A problem arises for the gem-rich areas in East Africa, the Urals, the Hindu Kush, Pamirs and elsewhere because cavities do not exist at the depths where regional metamorphism occurs.

In the absence of actual cavities, we might seek regions at depth where the constraining pressure is somehow regionally reduced. But such circumstances are also absent from metamorphic terrains.

To be clear, isolated 3–D spots or groups of spots with low pressure do exist and may indeed contribute to gem-formation. Low pressure spots at Merelani where tanzanite crystallized in association with boudins are a clear example (Olivier, 2008). But localized spots such as furnished by boudins or at the apexes of tight folds are only contributing factors. As a matter of scale — cm vs. km — they throw no light on the existence of extended metamorphic terrains (in East Africa, the Urals...) that host multiple gem deposits formed at depth.

There is another problem too: all the colored gemstones in a given region crystallized within one particular short-lived window of time. In East Africa, for example, all the gems crystallized approximately 550 million years ago.

Elements of a solution

The pressure at any depth in the Earth is essentially fixed, but this is not necessarily true for temperature.

An increase in temperature at depth would allow certain minerals to crystallize higher in the Earth, hence under lower constraining pressure, hence, in some cases, as gems.

A rise in temperature leading to the formation of gems under metamorphic conditions cannot, however, be achieved by familiar sources of heat such as volcanism or igneous intrusions. For whereas volcanism and igneous intrusions are common, gem-producing regions are rare.

Thus, for the gem-rich areas in East Africa, for example, it is necessary to identify a source of heat that was active. c.550 million years ago *but not before that time and not after*.

A candidate-source of heat available for the formation of metamorphic gems is furnished by thrust faults whereby thick sheets of rock, thrust one over another, produce temperature inversions that place hot rocks from deep in the Earth on top of cooler rocks from closer to the surface; Figure 1.

Such temperature inversions are locally augmented by transient frictional heat whose magnitude is determined by the types of rocks and the thickness and temperature of the thrust sheets (Fritz, *et al.*, 2009).

Heat was dispersed *upward* into overlying rocks *though only temporarily*, producing elevated temperatures at unusually shallow depths. Certain minerals were then “tricked” into crystallizing higher in the Earth than usual, hence with better quality crystallization. And in East Africa, the Urals, Himalayas, Hindu Kush, Pamirs, and presumably elsewhere, the time of thrusting and the time of gem formation are correlated.

Such scenarios, along with secondary contributing factors such as faults, the formation of boudins, or the production of temperature-lowering fluxes, provide circumstances that might allow high-quality crystallization at depth throughout a sizable region.

But we are confronted with the fact that large thrusts, which characterize continent-to-continent collisions, are not uncommon. And, to repeat, gem-rich regions are rare. So there must be differences between large-scale thrusting in general and those particular large-scale thrusts (in East Africa, the Urals...) that actually lead to the formation of gems.

One difference may be the thickness of individual thrust sheets, and another difference would be their initial temperature (see Fritz, *et al.*, 2009).

Another difference, potentially affecting heat production, is the distance inland from the collision zone. For, as has been shown for the India–Eurasia (Hindu Kush, Pamirs) convergence (Figure 2), the initial subduction of coastal sediments causes a “temporary acceleration in subduction rates” by a factor of >2, a phenomenon responsible for additional frictional heat, and thought to be “a common feature at the final stage of continental assembly” (Zhou *et al.* 2024). The incoming plate sprints at the end of its marathon!

Yet this is not the last word for, as can be observed (Figure 2), gems are preferentially formed at the passive edges of the “target” continent in places where the incoming plate encounters a circular plug of resistant rock, which is the case, as I have long argued in East Africa, the Hindu Kush, Pamirs and the Middle Urals Ring Structure (Saul, 2014, 2016, 2017, 2022, 2023; Burba, 1991, 2003a, 2003b). In such places thrusting over the resistant plug is much like that of sandpapering over a knot.

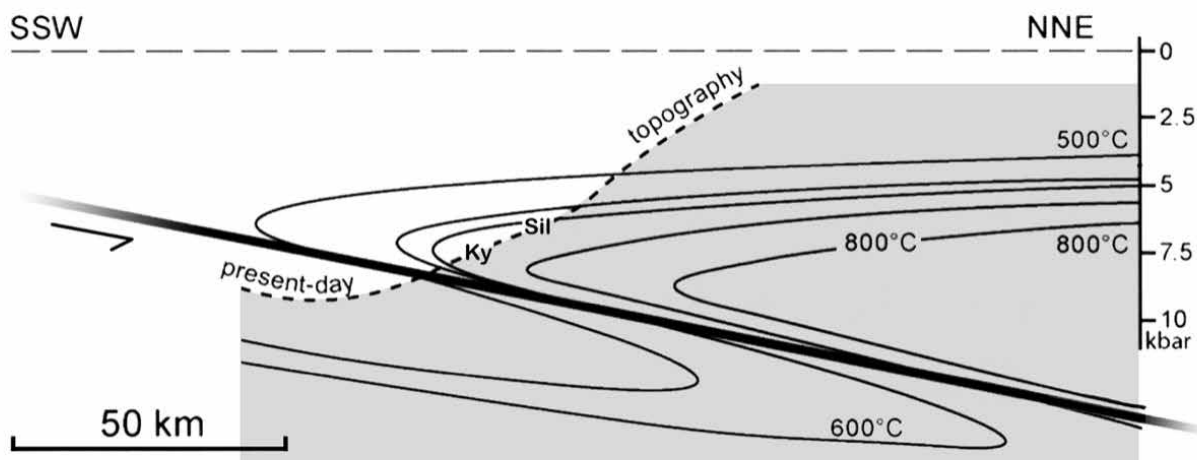


Figure 1: Thermal profile along the Himalayas in central Nepal. A thrust fault (thick line) caused overlying rocks to be metamorphosed at higher temperatures than those below. This allowed minerals to crystallize higher in the Earth than is usual,

hence with less constraining pressure. “Sil” indicates sillimanite and “Ky”, below it, indicates kyanite, which normally crystallizes above sillimanite. Adapted from Le Fort (1975). Similar profiles for East Africa or the Urals were not available for this study.

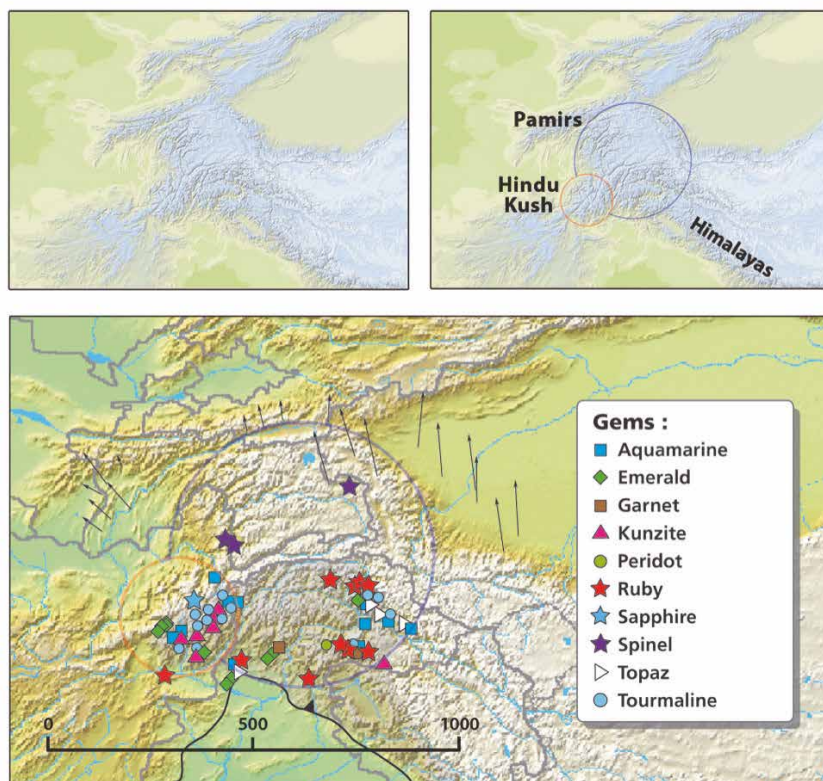


Figure 2: Gemstone locality map of the Hindu Kush and the Pamirs, both of which have faint circular outlines.

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Hessonite garnet from Angul district of Odisha, India

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Keywords: Hessonite garnet, inclusions, Angul, Odisha

Introduction

This study describes hessonite (Fe-bearing variety of grossular garnet), found in the Birlamunda village, located in Palalahada tehsil of Angul district (N 20° 50' 56.508", E 85° 3' 38.844") of Odisha state. Rocks in this region are a part of the Precambrian khondalite-charnockite-granite gneiss. Hessonite occurs along the contact zone of garnetiferous-gneiss and the amphibolite. Gemmological properties have been determined. Semi-quantitative analyses carried out using EDXRF show presence of iron, titanium and chromium. Material shows distinct turbidity or roiled structure like treacle commonly known as heat-wave effect. Crystalline inclusions identified with Raman were apatite, calcite, diopside, ilmenite, quartz, rutile and zircon.

Materials and methods:

Hessonite garnet from Birlamunda village with prominent inclusions were selected. Using gemmological methods their optical properties, refractive index, specific gravity, UV fluorescence etc. were determined. Inclusions crystallizing near the surface were identified by Raman spectroscopy. Semi-quantitative analyses were carried out using EDXRF and electron microprobe.

Results

Visual Appearance: Hessonite garnet from Birlamunda village of Angul District of Odisha were found to be having turbidity, treacly appearance with a brownish yellow to bright orangish yellow body color and a large number of crystalline inclusions. The coloration in these hessonite garnets is due to Cr and Fe content.

Gemmological Properties: Specific gravity was found to be in the range of 3.69 to 3.72, refractive indices fell in the range of 1.743 to 1.746. All samples showed no fluorescence under LWUV as well as under SWUV.

Microscopic Observations: Birlamunda hessonite garnet display a characteristic turbidity or roiled appearance which looks more like heat wave effect (Fig 1&2). Most samples were included with very tiny transparent crystals of apatite and some other crystals (Fig 3). Calcite rhombs were also seen in some specimen (Fig 4). In many of these hessonite garnets there were euhedral apatite crystal inclusion (Fig 5). These apatite crystals were of prismatic shape. some were grouped together with quartz crystals, which manifest as stumpy crystals (Fig 6). Some samples contained rutile in the form of fine needles arranged in three directions (Fig 7). In four specimen there were ilmenite crystals as inclusions (Fig 8). Diopside crystals were prominently observed in many of the specimen (Fig 9). Some diopside crystals were grouped together in a straight line (Fig 10). The yellowish-brown specimen had more of zircon halo type inclusions (Fig 11). In one unique case there were 'fused' crystals of apatite looking like a 'giant' crystal inclusion (Fig 12).

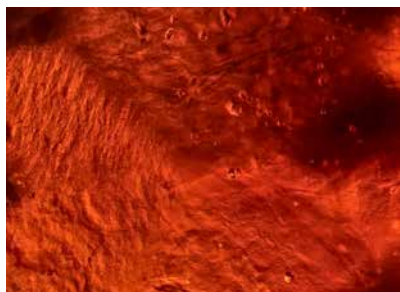


Fig 1: Heat wave like appearance



Fig 2: Roiled treacle-like appearance

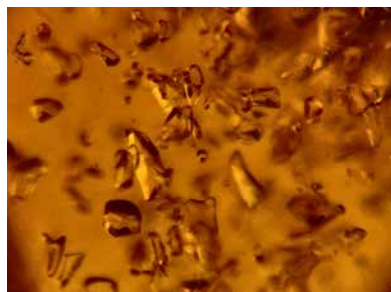


Fig 3: Variety of crystals observed



Fig 4: Calcite crystal inclusion

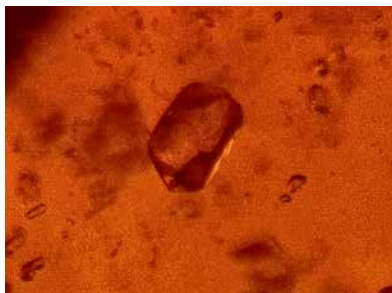


Fig 5: Apatite crystal inclusion

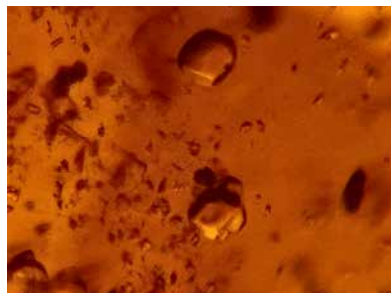


Fig 6: Stumpy quartz crystals



Fig 7: Fine rutile needles



Fig 8: Ilmenite crystal inclusion



Fig 9: Diopside crystal inclusion



Fig 10: Diopside crystals

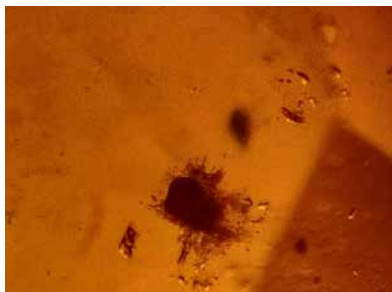


Fig 11: Zircon inclusion with halo

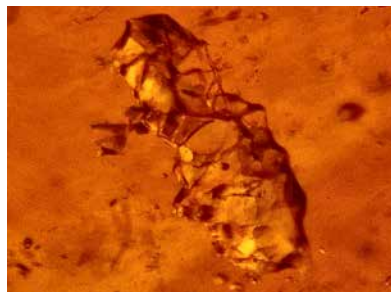


Fig 12: Fused crystals of apatite

Chemical Analyses: Semi-quantitative chemical analyses by EDXRF for hessonite garnet gave the oxide weight % for SiO_2 =37.98 to 38.82, Al_2O_3 =20.47–20.05, for CaO =38.98–39.01 and for FeO = 1.12-1.15, whereas the oxide weight % for Cr_2O_3 and TiO_2 was determined as 1.08 and 0.37 respectively. No manganese was detected in our samples by EDXRF.

Discussion: The inclusions in hessonite described by Kanis & Redmann (1994) from four different localities, viz. Ghatpara, Burubura, Budhido and Dahikbala (Fig13 & 14) in Orissa (now Odisha) are somewhat similar to the ones observed in the present study. All hessonites described by them from these four occurrences contain traces of MnO based on their chemical analyses, in contrast to this we did not analyze any manganese in our hessonite samples from Birlamunda. Besides, the Ghatpara and Burubura hessonite display two and three phase inclusions. Whereas the hessonite from Dahikbala contain needlelike inclusions. Budhido hessonite garnets have orthoclase mineral inclusions. None of the specimen investigated in this study displayed neither orthoclase inclusions nor two or three-phase inclusions. They contain, however, other inclusions such as zircon, apatite, diopside and calcite crystals.

The Birlamunda hessonite garnet occurs in a variety of color shades of orangish- brown to yellowish brown tending towards honey color. The coloration is mainly due to Fe with possibly minor contributions by Cr. The Birlamunda hessonite garnet contain a large number of solid inclusions in a distinct roiled or treacle like ‘background’ in which the crystalline inclusions are present. According Gübelin and Koivula (1986) the treacle effect is due to the colloidal separation of calcite. The authors are of the opinion that this roiled treacle-like appearance is mainly due to the drastic changes in temperature and pressure during the crystallization of the hessonite garnet. These variations in pressure and temperature had created disturbances in



Fig 13: Position of Odisha state

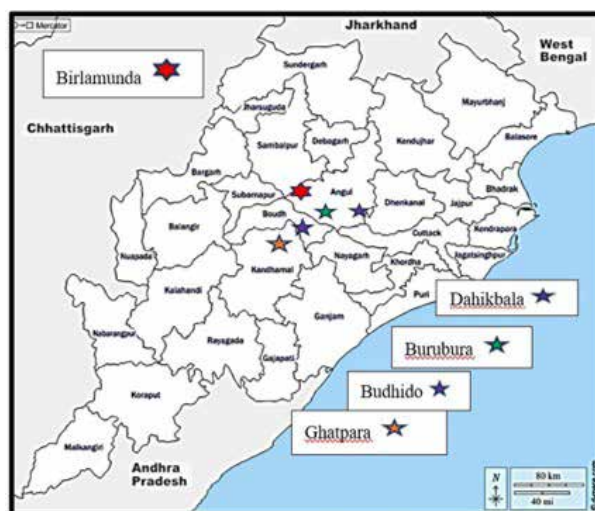


Fig 14: Localities described in Kanis & Redmann and Birlamunda

the solid solutions and given rise to polycrystalline aggregate of small hessonite garnet. The small RI differences between the single grains of the polycrystalline structure are responsible for the roiled type effect. The large number of crystalline inclusions of apatite, diopside, zircon, calcite, rutile, quartz and ilmenite indicate the mother-liquor was having a characteristic metamorphic assemblage. Some of the inclusions of apatite and diopside appear to be fused together to form large giant crystals.

Conclusion

The Birlamunda hessonite garnets have some inclusions similar to those studied by Kanis & Redmann, (1994), indicating that the emplacements of these garnets may have been in a similar type of metamorphic rock mentioned in that study. The geology of the Indian subcontinent is extremely complicated with numerous metamorphic events, so it is indeed difficult to indicate whether the geological period was also similar to that in the study done by

Kanis & Redmann (1994), it could be similar. There are crystals of rutile and ilmenite in some of the specimens which were not described in the hessonites studied by Kanis & Redmann or by Gübelin and Koivula (1986). Rutile needles seem to be perfectly oriented. Oriented acicular inclusions of the Birlamunda hessonite garnet could be the result of crystallization at ultrahigh temperature of above 1000°C and pressure above 3 GPa as stated in Axler & Ague (2015).

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Acknowledgements

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Hurlbutite from Myanmar

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A crystal of the very rare mineral hurlbutite $\text{CaBe}_2(\text{PO}_4)_2$ originating from Myanmar (without further locality details) was studied and faceted afterwards in two gems, which we also studied. This specimen is of particular interest because the species rarely produces well-formed crystals, but rather masses or globules (spherulites; Mindat, 2025). The crystal is a near-colorless, short lozengic prism, measuring approximately 17 x 10 x 7 mm and weighing about 2 grams. It is naturally etched, with few inclusions visible through the frosted surface, and a dissolved dislocation near the broken base. The ultraviolet (UV) luminescence is weak, greenish white in longwave, and weak orange in shortwave UV. The rough was faceted into two stones, a 0.64 ct round brilliant cut and a 2.56 ct rectangular scissor cut (RSC; Figure 1).

The refractive indices measured on the faceted stones are $n_\alpha = 1.593$, $n_\beta = 1.600$ and $n_\gamma = 1.606$, close to values provided in the original description $n_\alpha = 1.595(3)$ $n_\beta = 1.601(3)$ $n_\gamma = 1.604(3)$ (Mrose, 1952). It is optically biaxial negative. The specific gravity measured by the hydrostatic method is 2.88 compared to 2.877(5) (measured: Mrose, 1952) and 2.90 (calculated) (Mindat).

To verify the exact nature of this rare gem, on which very limited literature is available, we combined Raman scattering with X-ray diffraction and chemical analysis using Energy Dispersive X-ray Fluorescence (EDXRF).

The Raman scattering was obtained on a Renishaw InVia Raman microscope with a 514 nm laser and a 4 cm^{-1} resolution. The main peak is that of phosphates, at about 1017 cm^{-1} (Figure 2). The patterns obtained on the rough and the two faceted gems match both RRUFF references R070612 (from Finland) and R090048 (from the type locality in the USA).



Figure 1: The rough hurlbutite crystal (height about 17 mm), and the two fashioned gems, a 2.56 ct rectangular scissor cut and a 0.64 ct round brilliant cut. Photos LFG

Powder X-ray diffraction was performed on a D6 Phaser de Bruker, with a copper anode, a 30 kV voltage and a 18 mA current, on ground fragments remaining from the cutting process. The diffractogram obtained in the 5 to 90° 2 θ angle range matches hurlbutite references in RRUFF and the Crystallography Open Database (COD).

The semi quantitative chemistry on the rough and afterwards on the faceted stones using EDXRF was measured on a ThermoScientific QUANTX. As expected from the chemical formula, calcium and phosphorus are responsible for the strongest signals, confirming identification indirectly. The main impurity is strontium (Sr) known to substitute for Ca. Additionally minor Mn, Pb, Al, Si and S were detected. The beryllophosphate hurlbutite is typical of the pegma-

titic environment; Two of its components, beryllium and phosphorus, are generally found concentrated in pegmatitic fluids. In its original occurrence, Chandlers Mill Quarry, Newport, New Hampshire, USA, hurlbutite appears in a complex granitic pegmatite (Mrose, 1952).

Photoluminescence (PL) excited at 514 nm (RSC 2.56 ct) shows a broad band centered about 580 nm (greenish yellow) extending from about 500 to 650 nm, and a more intense one with apparent maximum about 737 nm in the red and near infrared (Figure 3). The 580 nm band may cor-

respond to the greenish white emission observed in LWUV. There are in addition a few sharp lines at 543 nm, as well as a group at 854, 859, 869, 874, 886 nm, possibly caused by trivalent rare earth elements ions (REE^{3+}).

To our knowledge, there was no existing description of hurlbutite as a gem material in the literature before our contribution. We were able to confirm properties from the mineralogical references and gather some gemological properties, which are awaiting confirmation if another gem-quality hurlbutite is found.

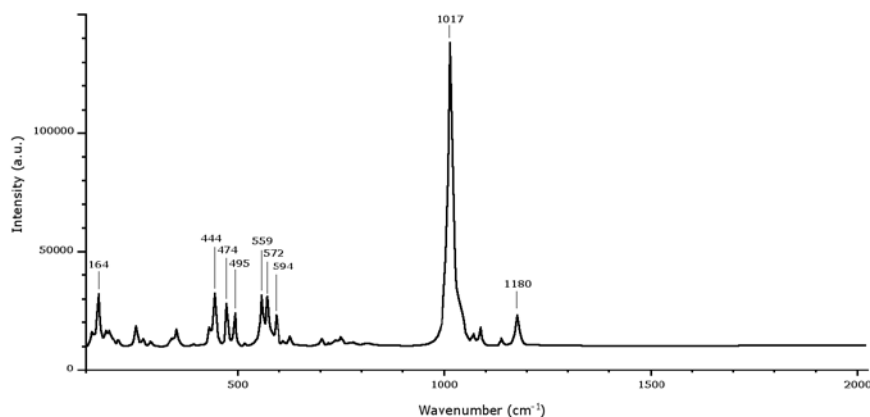


Figure 2: Raman spectra of the hurlbutite crystal. Vertical scale arbitrary.

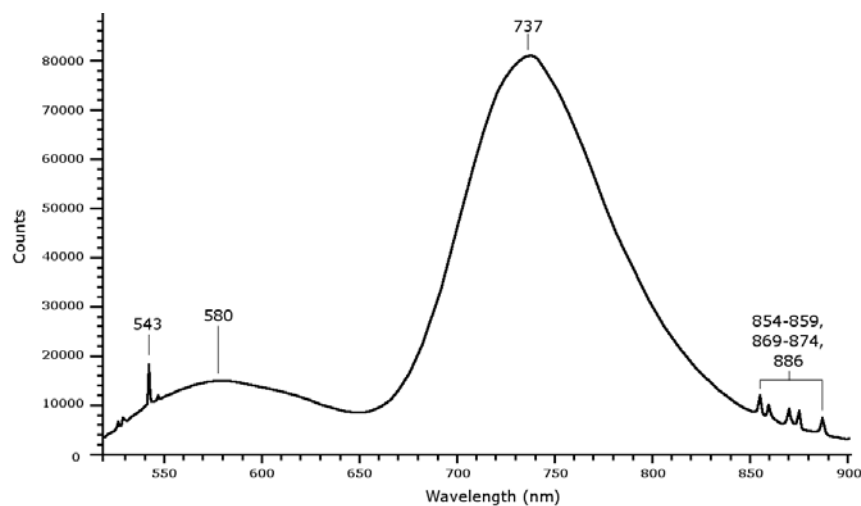


Figure 3: Photoluminescence spectrum of the 2.56 ct hurlbutite excited with a 514 nm laser at room temperature, from 500 to 900 nm. Vertical scale arbitrary.

References:

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A convincing glass imitation of larimar (pectolite)

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Larimar, a massive blue variety of the mineral pectolite ($\text{NaCa}_2\text{Si}_3\text{O}_8(\text{OH})$), belonging to the wollastonite group of minerals is known in the trade for few decades now. However, in recent times it has gained popularity amongst jewellery designers and consumers, as suggested by high frequency of submissions at the laboratory for certification. Originating only from the Dominican Republic, its striking blue colour with light blue to white waves or web like patterns caused by the spherulitic growth has made larimar a sought-after gemstone.

With increasing popularity, larimar has been subject to imitation by various materials, including ceramics (Kiefert and Groenenboom 2013) and naturally dyed stones such as chalcedony (Miura, 2020). Similar man-made ornamental materials were produced as early as the 1950s, including the well-known 'Victoria Stone' (Renfro, 2017).

The present author has examined numerous specimens of translucent to opaque partly devitrified glass submitted to the laboratory, many of which closely resemble larimar in both appearance and colour—making them convincing imitations. These materials are reportedly being presented as larimar at both online and offline platforms. This study documents the gemmological characteristics of this glass imitation of larimar encountered at the laboratory over the past one and a half years, sourced from a local gem dealer.

Results and Discussion

Visual appearance

Rough samples, slices, and polished cabochons of both larimar and the larimar imitation were selected for this study. Majority of larimar as well as imitation samples were opaque, however, using strong fibre optic light few sections appeared translucent. Larimar specimens exhibited colours ranging from blue to greenish blue, with characteristic white

web-like patterns (Figure 1, left). In comparison, the larimar imitation (partly devitrified glass) samples ranged from greenish blue to bluish green and displayed similar white webbing, particularly evident in sliced sections (Figure 1, right). In cabochon form, the imitation samples often featured a darker central core surrounded by a radiating fibrous and whitish zone. This fibrous zone also exhibited a noticeable sheen or chatoyant effect.

Cross-section of rough imitation samples displayed radiating fibrous structure typically associated with spherulitic growth, while the outer surface (crust) had a transparent glassy layer with botryoidal pattern (Figure 2, left and right).

Properties

Gemmological properties of the studied samples of larimar and larimar imitation are summarized in Table 1.

Raman spectroscopy was performed on larimar and larimar imitation samples using a 532 nm excitation laser, covering the spectral range of 100–2000 cm^{-1} (Figure 3). The larimar samples exhibited a rising background absorption from 200 to 2000 cm^{-1} , attributed to fluorescence, with major peaks observed at approximately 651 and 1025 cm^{-1} , along with smaller features at ~317, 375, 413 and 970 cm^{-1} .

The opaque or crystalline areas of the imitation samples displayed a consistent spectral pattern across multiple spots, characterized by major peaks at approximately 635 and 970 cm^{-1} , and additional smaller or broader features at ~331, 413 and 1042 cm^{-1} . Interestingly, few features such as ~413 and 970 cm^{-1} were present in spectra of both, larimar and crystalline areas of imitation samples suggesting a similarity in composition and structure. Raman spectra of crystalline areas of the imitation were consistent with that of wollastonite given in the RRUFF database and elsewhere (e.g., Hänni *et al.*, 2001). Peak at ~970 cm^{-1} is present as

the major feature in glass, while present only as a weak feature in pectolite. Further, the peak at $\sim 635\text{cm}^{-1}$ present in glass shifts to peak at 651cm^{-1} in larimar, while the peak at 1025cm^{-1} present in larimar shifts to 1042cm^{-1} in the

devitrified glass. In contrast, the transparent glass crust of the larimar imitation exhibited broad bands in the ranges of approximately $370\text{--}620\text{ cm}^{-1}$ and $800\text{--}1100\text{ cm}^{-1}$, consistent with the spectral signature typical of amorphous glass.



Figure 1: Representative cabochon and sliced rough samples of larimar (left) and glass imitation (right), displaying a close resemblance in colour—ranging from greenish blue to bluish green—and white web-like patterns. The left cabochon in glass (right image) also exhibits a notable sheen effect due to fibrous inclusions. Photos by G. Choudhary.



Figure 2: Rough sample of devitrified glass showing an opaque radiating pattern originating from a central core and terminating in circular edges, indicative of spherulitic growth (left). In contrast, the surface or crust of the same sample exhibits a transparent, glassy layer with a botryoidal pattern (right). Photos by G. Choudhary.

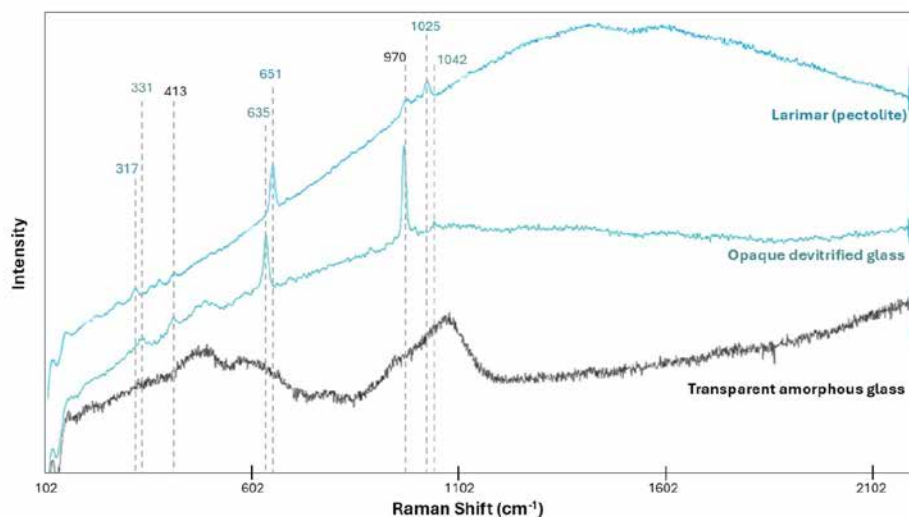


Figure 3: Raman spectra of larimar (blue), opaque devitrified glass (blue-green), and amorphous glass matrix (black).

Property	Larimar	Larimar imitation
Colour	Blue to greenish blue; white webbing	Greenish blue to bluish green; white webbing
Diaphaneity	Opaque under standard lighting; few sections translucent under fibre optic light	Opaque under standard lighting; few sections translucent under fibre optic light. Top surface transparent.
RI	~1.60 (spot); distinct birefringence blink ranging from ~1.58 – 1.62	~ 1.49 (spot); no birefringence blink
SG (hydrostatic)	2.83 – 2.87	2.54 – 2.58
UV Fluorescence	Moderate to distinct yellowish green in shortwave; inert under longwave	Weak yellowish green in shortwave; inert under longwave
EDXRF analyses	Major: Ca, Si, Al, Na, Mg Minor: Cu, Fe, Mn, K Traces: V, Sr, Zn, Ti, Ni	Major: Si, Ca, Al, Na, K, Fe, Mg Minor: Ti, Zr, Traces: Sr, Rb, W, Mn, Sn, Zn, Pb
Raman analyses	Major peaks: ~651 and 1025 cm ⁻¹ Weaker features: ~317, 375, 413 and 970 cm ⁻¹	Crystalline areas Major peaks: ~635 and 970 cm ⁻¹ Weaker features: ~331, 413 and 1042 cm ⁻¹ Amorphous areas (glass crust) Broad bands in the range ~370-620 and 800-1100 cm ⁻¹

Microscopic examination

The larimar imitation samples exhibited fine fibrous inclusions (Figure 4) arranged in a radiating pattern originating from distinct cores. In several specimens, multiple such cores were present, resulting in a botryoidal or spherulitic appearance. These fibrous inclusions often produced a sheen effect, contributing to the visual similarity with natural fibrous minerals. The inclusions themselves were colourless, showed well-defined growth striations along their lengths, and displayed fine cracks reminiscent of cleav-



Figure 4: Fine, tightly packed, intergrown fibrous crystals arranged in a radial to subparallel pattern in opaque areas of glass-imitation of larimar. Photomicrograph by G. Choudhary; image width 5.08mm

age features commonly observed in minerals with prominent cleavage perpendicular to the crystallographic c-axis.

Similar lath-like crystals in a glass matrix have previously been reported to be wollastonite (Hänni *et al.*, 2001) or apatite (Renfro, 2017), tanohataite and cristobalite (Fayed *et al.*, 2025). However, Raman spectroscopic analysis of the inclusions in the present study was consistent with that of wollastonite.

Conclusions

The described partly devitrified glass serves as a convincing imitation of larimar (pectolite) due to its similar webbing pattern and visual appearance, which may potentially mislead gemmologists and gem dealers. However, standard gemmological properties, including RI and SG, allow for easy distinction between partly devitrified glass and larimar, supplemented by Raman spectroscopy and chemical analysis. These imitations typically show higher concentrations of Si and Al, and lower Ca content compared to larimar. Additionally, the presence of coarse, needle-like microscopic structures resulting from devitrification provides further diagnostic evidence for separation. Although the identification and separation of this partly devitrified glass are not particularly challenging, raising awareness about its existence is crucial to prevent misidentification in the gem trade.

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Bertrandite in Emeralds

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Over the past few years we have been seeing more and more lower quality stones in the lab. The reasons for this may be related to lack of supply of finer goods, rising prices, the global pandemic or others. Among these lower quality stones have been large numbers of heavily included emeralds (Figure 1).

We began to notice that a number of these emeralds were highly fractured with the fractures being very low relief. They appeared to be filled with a clarity enhancing substance and for a while were treated as such. However, we noticed that there were differences between these fractures and ones we knew to be filled with polymers to enhance the clarity of emeralds.

When testing for clarity enhancement, we look for certain clues. These include flash effect, high relief unfilled areas, gas bubbles, UV fluorescence, sweating of filler out of the fractures, polymers in the FTIR spectra and more. We use these indications to support the conclusion that there is a filler present and it is affecting the appearance of the stone.



Figure 1: An example of a heavily included emerald that contains bertrandite filled fractures. 1.82 carats.

However, it is possible that many of these things may not be present in a given stone. So the lack of these indications in suspicious looking fractures can make it challenging to decide if a stone is clarity enhanced or not.

For the emeralds in question what looked like a filler in many of them did not show bubbles or unfilled areas, did not have a flash effect or UV fluorescence of any indication of polymer in the FTIR. The fractures sometimes appeared brownish when viewed down the length of the fracture (Figure 2) and also sometimes showed interference colors when viewed under crossed polaroids. Sometimes there was an irregular structure within the fractures that resembled a roiled effect.

Many of these fractures are very narrow but some get wide enough that the material in them can be seen at the surface. The luster of this material is somewhat different than the host emerald and often slightly undercut, indicating that it is softer than the emerald (Figure 3). Raman analysis came up with a match for two minerals – some phlogopite but mostly a beryllium mineral called bertrandite.

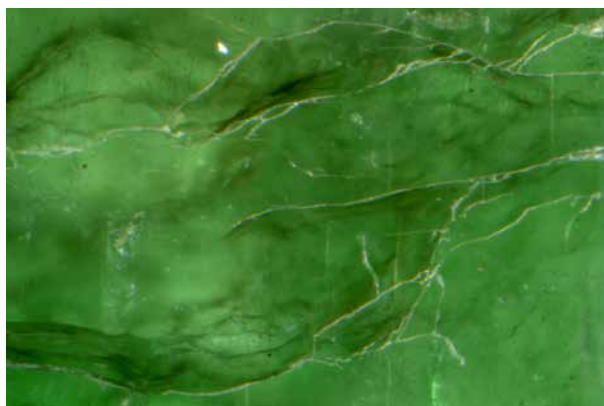


Figure 2: Fractures filled with bertrandite sometimes look brownish when viewed down the length of the fractures. Photomicrograph by Chandler Powers, field of view 1.83 mm

Phlogopite is fairly common in emeralds from a number of localities but typically as an inclusion, not as a filler in fractures. Bertrandite, while commonly associated with beryl in some environments, has been mentioned a few times as an inclusion in emerald, but we could only find one reference of it filling fractures.

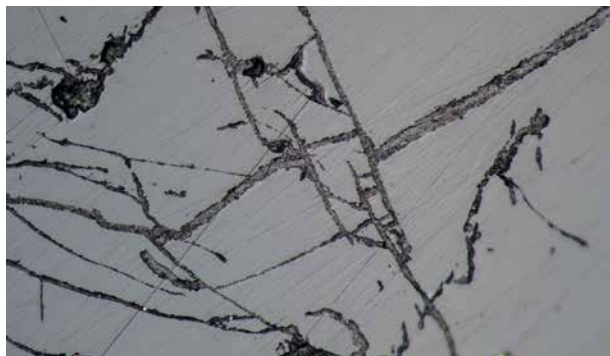


Figure 3: Extensive fracture system with bertrandite showing different surface luster and slight undercutting. Photomicrograph by Nathan Renfro, field of view 0.72mm

A search of the literature revealed a note by Zellagui (2022). In this article he documented an instance where they identified a wide fracture system in an emerald with bertrandite and phlogopite filling it. He showed the bertrandite along the edges of the fractures with the phlogopite filling the middle.

In our experience we have not yet seen an example where there was a clear difference in surface luster in a single fracture which might indicate both minerals being present. This could possibly be attributed to the width of the fractures we have observed being narrower than the example mentioned in the note.

In an attempt to see if we could identify the presence of bertrandite in these fractures in the FTIR spectra we found that in some cases the two strongest peaks in the FTIR spectrum of bertrandite showed up as tiny broad peaks at approximately 6981 and 6930 cm^{-1} within the emerald spectrum (Figure 4). Unfortunately these peaks are often too weak to be visible even though spectra were run in multiple directions.

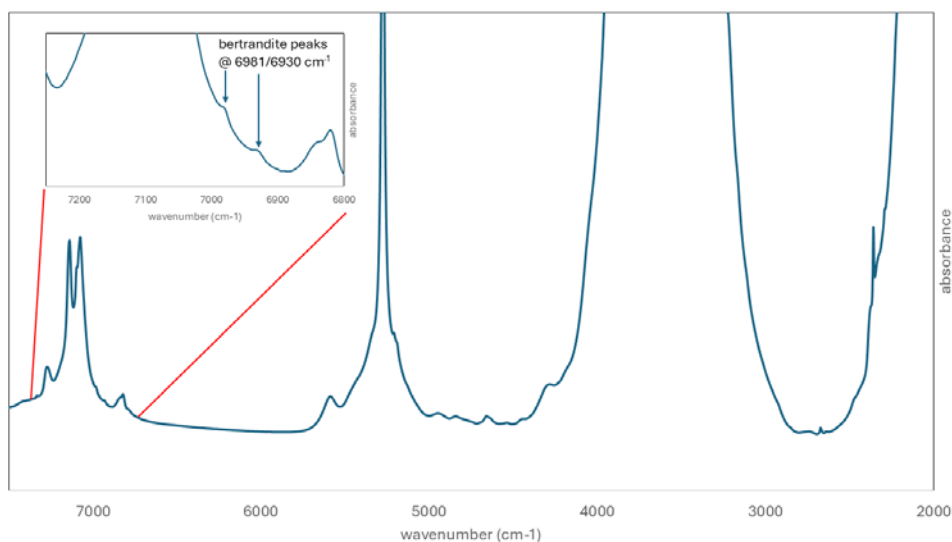


Figure 4: FTIR spectrum of an emerald showing two small bands at 6981 and 6930 cm^{-1} indicating the presence of bertrandite in this stone.

Bertrandite is a beryllium silicate - $\text{Be}_4(\text{Si}_2\text{O}_7)(\text{OH})_2$ - that Mindat.org describes as “a common hydrothermal alteration product of beryl”. It has an R.I. of approximately 1.59 – 1.61, which explains why fractures filled with it in high iron emeralds have such low relief. It is said to commonly replace beryl, often with a sheet silicate such as mica taking up the aluminum content of the beryl (Barton 1986). This could explain the detection of phlogopite in association with the bertrandite in fractures.

Most references discuss the formation of bertrandite in pegmatites, although there is some mention of it forming in non-pegmatitic environments (Barton, and Young, 2002). It is one of the last minerals to form in such pegmatites and requires water vapor to be present to form (Jacobson, 1988). Theoretically this could explain how bertrandite could form in the open fractures of a beryl crystal.

So, if all the fractures present in an emerald are filled with another mineral, then the stone has not been clarity enhanced

and we have seen quite a few stones where this was the case. However, we also have seen quite a few where some fractures were not filled with a mineral and these fractures were usually filled with a polymer. We have seen both oil and artificial resin that exhibited the properties we would expect from such fillers, such as flash effect with the artificial resin. These fractures fluoresce weakly while all the other fractures filled with bertrandite do not fluoresce at all. The polymers show up in the FTIR. The challenge then becomes trying to separate which fractures are filled with what and how much that is affecting the appearance of a stone.

At this point in time, the origin of these stones is still unclear. The people submitting the stones to the lab say they do not know the origin. In comparing the chemical plots against known samples from the GIA reference collection, some of these stones are similar to samples originating in Russia, while others are similar to Zambia. However, all of the stones containing bertrandite look very similar to one another, so we are convinced that they are coming from the same locality.

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Found, Lost, and Found Again: The Geological and Gemological Significance of the Salininha Emerald Deposit, Brazil

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Abstract

The Salininha emerald deposit (Figure 1), located in the municipality of Pilão Arcado, Bahia State, represents the first economically significant discovery of gem-quality emeralds in Brazil (Sauer 1982; 1992). Upon its recognition, controversy arose among specialists who questioned the classification of the Salininha crystals as true emeralds due to the predominance of vanadium, rather than chromium, as the chromophore element. The deposit was mined out after approximately one year of extraction (Sauer 1992) and subsequently abandoned. Currently, there are efforts underway to resume exploration activities in the region, with new material being extracted. Geologically, the occurrence is situated within the domain of deformed granitic rocks, which locally preserve remnants of metatexites, diatexites, and metabasalts that were not fully digested by granitization (Souza *et al.* 1979). These remnants can be interpreted as Archean granite-greenstone complex, later intruded by Paleoproterozoic granitoids, and deformed at ~2 Ga. in the events of the Transamazonian orogeny (Hurley *et al.* 1967). The emeralds are hosted within a talc-schist lens approximately 100 meters long and 50 meters thick, which is intruded by pegmatoid veins (Souza *et al.* 1979). In the samples analyzed, the emerald crystals grew over the meta-ultramafic host rock, showing no structural alignment with the main foliation of the talc-schist (Figure 2A). Micro-Raman spectroscopy was conducted on rough crystals and their inclusions, revealing the presence of talc and calcite. The inclusions in emeralds are also preferentially aligned with this foliation and are interpreted as relicts of the meta-ultramafic protolith, which likely served as the source of vanadium. The emerald crystals exhibit a flattened hexagonal prismatic habit and vary in color from very light to light yellowish green (Figure 2B

and Figure 2C), with pleochroism ranging from yellowish green to bluish green. Additionally, the analyzed crystals exhibit no fluorescence under UV light and display a stable green color when observed through the Chelsea filter, consistent with vanadium-based coloration. They exhibit specific gravity values ranging from 2.67 to 2.75 g/cm³ (measured in rough material) and refractive indices from $n_e = 1.588$ to $n_o = 1.595$ (measured in faceted samples from the same locality - Figure 2D). Given the absence of deformation in the emerald crystals, the mineralization is attributed to a metasomatic process similar to that observed in the Campo Formoso emerald district, located ~300 km to the southeast. In Campo Formoso emeralds are hosted in metavolcano-sedimentary sequences of Greenstone affinity, tectonically inserted into the granitic rocks of the Archean basement, sometimes intruded by younger granitic rocks. The emeralds crystallized within discordant pegmatitic veins and adjacent metasomatic zones under moderate temperature and pressure conditions, without direct genetic association with the nearby granitic intrusions (Capovilla 1995). The Salininha deposit not only marked the inception of Brazil's prominence in the global emerald market but also contributed significantly to the understanding of vanadium-induced coloration mechanisms in beryl. Its geological setting, characterized by the metasomatic transformation of meta-ultramafic rocks and the crystallization of emeralds within discordant pegmatitic veins, provides valuable insights into emerald genesis in greenstone-related environments. Renewed geological and mineralogical investigations can highlight the potential to identify new emerald occurrences in the Salininha area and its surroundings.

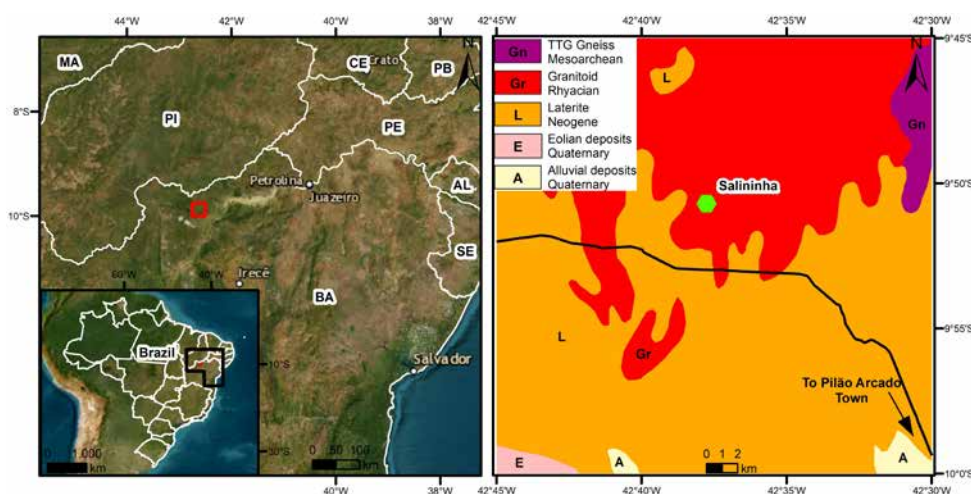


Figure 1: Location and geological map of the Salininha mine area. The geological mapping was modified from Souza *et al.* (2003)



Figure 2A: Emerald crystal from Salininha overgrowing the foliated talc-schist, with the main foliation of the sample lying horizontally in the image. Figure 2B: Artisanal miner (garimpeiro) holding rough emerald crystals. Figure 2C: Lot of rough emeralds including both small and large crystals. Figure 2D: Faceted emerald crystals displaying a light green color.

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We would like to thank the Laboratory of Fluid Characterization in Geological Systems (GeoFluid), of the Institute of Geosciences of the University of São Paulo for the micro-Raman analyses.

Automated Data Processing of Raman Spectra for Supporting Spinel Origin Determination

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Keywords: Spinel, Raman spectroscopy, provenance, automated data processing, origin determination, machine learning

Introduction

Spinel is a highly valued gemstone, with its color and quality influenced by trace-element chemistry and crystal structure. Accurate determination of its geographic origin is crucial for gemmological authentication, scientific research, and maintaining supply-chain integrity. Raman spectroscopy provides a non-destructive analytical approach, offering vibrational fingerprints that reflect both the crystal lattice and impurity-related features. However, manual interpretation is time-consuming and susceptible to operator bias, particularly when dealing with large datasets or subtle spectral variations. To address these challenges, we present an end-to-end automated pipeline that integrates spectral pre-processing, feature extraction, and chemometric analysis. Machine learning (ML) models are then applied to classify spinel samples according to their geographic origin. The performance of this approach is demonstrated using both red and blue spinels from three major deposits.

Materials and Methods

A total of 315 spinel samples (Figure 1) were sourced from local, trusted dealers and verified by gemmological means. Red spinel (MgAl_2O_4 ; $n = 154$) originated from Mogok (Myanmar), Luc Yen (Vietnam) and Lukande (Tanzania), while blue samples ($n = 161$) comprised spinel (MgAl_2O_4) from Luc Yen and Lukande and gahnite (ideally ZnAl_2O_4 ; $n = 52$) from Jemaa (Nigeria). Each sample was polished to present at least one flat surface for Raman analysis. Raman spectra were collected on a confocal Renishaw inVia Raman microscope equipped with a 50x objective (NA 0.50) using a 785 nm excitation laser. Three accumulations of 10s each were acquired over 200–1000 cm^{-1} with 1 cm^{-1} spectral resolution, conditions verified to avoid local heating (no peak drift between successive scans). Raw spectra were exported as CSV files and processed via the automated data processing pipeline.



Figure 1. Representative spinel samples used in this study, with weights ranging from 0.37 to 2.70 carats.
(Photo by M. Seneewong Na Ayutthaya)

Automated Data Processing Pipeline

Our data-processing pipeline was implemented in Python 3.12 and proceeds as follows. First, Raman spectra are smoothed using a Savitzky–Golay filter (window length 7, polynomial order 3) to suppress high-frequency noise while preserving peak shapes. The smoothed spectra are then baseline-corrected via Asymmetric Least Squares (ALS) algorithm with $\lambda = 10^7$ and $p = 0.01$, followed by max-normalization ($I/I_{\max}$) to remove intensity scale effects. Next, we isolate the four vibrational bands at 310, 405, 665, and 765 cm^{-1} and fit each with a Lorentzian profile (accepting fits only if $R^2 \geq 0.95$) to extract the full width at half maximum (FWHM). The resulting FWHM values, together with two dimensionless ratios (e.g., 405/665 and 405/765), are assembled into feature vectors and visualized in bi-scatter plots to assess natural clustering by geographic origin. Finally, we train both an artificial neural network (ANN) and a random forest (RF) models using 10-fold cross-validation and a 20% held-out test set (red spinel $n = 28$; blue spinel $n = 30$), optimize hyperparameters by grid search, and report overall accuracy, macro F1-score, and mean cross-validation accuracy (mean CV).

Results and Discussion

The full dataset of 315 spectra was processed on a MacBook Pro (M1 Pro, 16 GB RAM) in 149 seconds, averaging 0.47 seconds per spectrum. FWHM-based plots reveal well-defined clusters corresponding to geographic provenance. For red

spinel, plotting the FWHM of the 405 cm^{-1} band against that of the 665 cm^{-1} band distinguishes samples from Myanmar (MM), Vietnam (VN), and Tanzania (TZ), though some overlap occurs between MM and VN. In blue spinel (MgAl_2O_4), the 405 cm^{-1} and 665 cm^{-1} modes are decisive. Nigerian gahnite (ZnAl_2O_4) occupies a distinct field due to homologous peak shifts to approximately 415 cm^{-1} and 657 cm^{-1} . Applying adaptive fitting windows of $\pm 10 \text{ cm}^{-1}$ further isolates gahnite without disturbing the true-spinel clusters (Fig. 2). While gahnite is also distinguishable from blue Mg–Al spinels by RI, SG, and Zn-rich chemistry, Raman provides a complementary method, especially for small or mounted stones. These FWHM shifts likely reflect variations in A–B site inversion and mass-related frequency shifts from Mg^{2+} substitution with heavier Zn^{2+} (Malavasi *et al.*, 2002; Wang *et al.*, 2020). Additionally, the separation—and remaining overlap—between TZ and VN blue spinels may be influenced by Fe/Co variations affecting site distortion (Furuya, 2023). Classification models trained on FWHM features performed strongly, as summarized in Table I. Confusion matrix analysis indicates most misclassifications occurred between MM and VN red spinels, and between TZ and VN blue ones, reflecting partially overlapping geological and chemical characteristics (Chauviré *et al.*, 2015; Chankhantha *et al.*, 2020; Krzemnicki *et al.* 2023, Wu *et al.*, 2023). Overall, these findings demonstrate that deposit-specific FWHM features provide an effective foundation for automated, machine learning-based provenance determination of spinel.

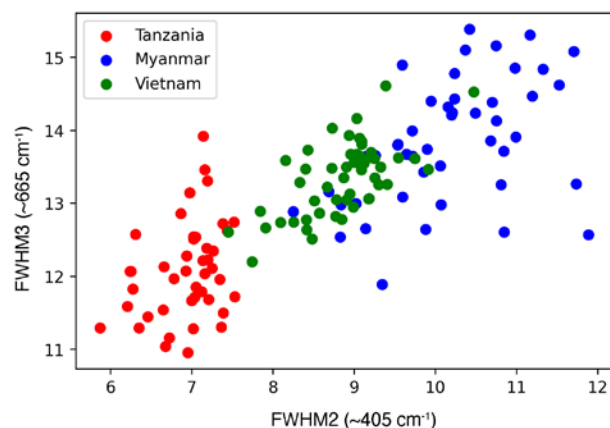
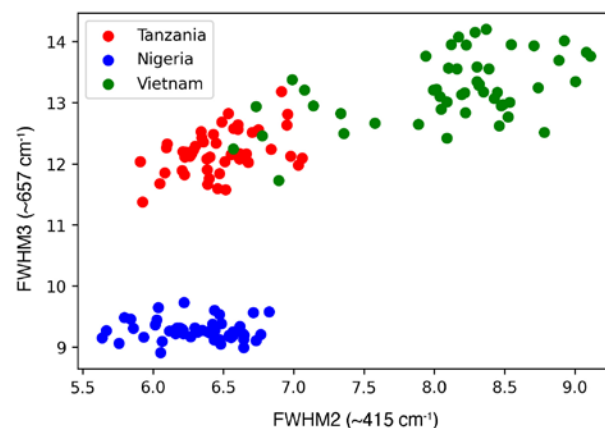


Figure 2. Scatter plots illustrating FWHM-based feature distributions for classification: (left) red spinel samples from



Myanmar, Vietnam, and Tanzania; (right) blue spinel (Vietnam, Tanzania) and gahnite (Nigeria).

Dataset	Model	Accuracy	Macro F1	Mean CV	Misclassified samples
Red spinel	ANN	93	93	89% $\pm$ 6.6%	2/28: (1MM $\rightarrow$ VZ, 1VN $\rightarrow$ TZ)
	RF	93	93	88% $\pm$ 7.1%	2/28: (1MM $\rightarrow$ VZ, 1VN $\rightarrow$ TZ)
Blue spinel & gahnite	ANN	97	97	95% $\pm$ 4.0%	1/30: (1VN $\rightarrow$ TZ)
	RF	93	94	96% $\pm$ 4.1%	2/30: (2VN $\rightarrow$ TZ)

Table I Performance summary of ML models applied to red and blue spinel classification

Conclusion

We present an automated, high-throughput workflow for Raman spectral processing and provenance classification of blue and red spinel. By using FWHM of key vibrational bands as features, the pipeline captures subtle lattice variations associated with geographic origin. This approach achieves over 90 % classification accuracy on the current dataset and significantly reduces analysis time, enabling scalable and reproducible gemmological assessments. The strong performance of FWHM-based features highlight their sensitivity to crystallographic disorder and trace-element chemistry—factors that are critical for provenance differentiation. Building on these results, future work will expand the geographic diversity of the spinel dataset and extend the pipeline to a broader range of gemstone species.

Limitations and Outlook

The current model is calibrated using a limited set of deposits; incorporating additional localities will likely introduce greater spectral variability, necessitating model retraining or, at minimum, site-specific recalibration. Broader datasets may also increase class overlap, which could complicate cluster separation and reduce classification accuracy. Furthermore, heating above ~ 800 °C has been shown to broaden the same Raman bands (Saeseaw *et al.*, 2009) used for provenance discrimination, potentially obscuring geographic signals. As a result, Raman-based FWHM classification should be integrated with complementary techniques—such as trace-element analysis—to disentangle provenance from heat treatment effects and establish a more robust, multi-modal framework for origin determination.

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Spinel from Sri Lanka - An update

Gamini Zoysa, Dietmar Schwarz, Arunas Kleismantas

Most of the spinels are found in shades of red to purple. Colourless, green and cobalt bearing varieties also been recorded in few locations. Same as other gemstones from Sri Lanka. Spinel also occur in alluvial deposits in south-western and north central parts of the island.

The following subjects will be discussed in the presentation.

1. Mineralogy and Crystallography
2. Geology and Genetic Aspects
3. Gemological and Mineralogical Properties
4. Internal Features
5. Comparison of Properties of Spinel from Different Locations from Sri Lanka (Ratnapura, Horana, Eheliyagoda, Elahera)
6. Chemical Fingerprinting
7. Spectral Fingerprinting

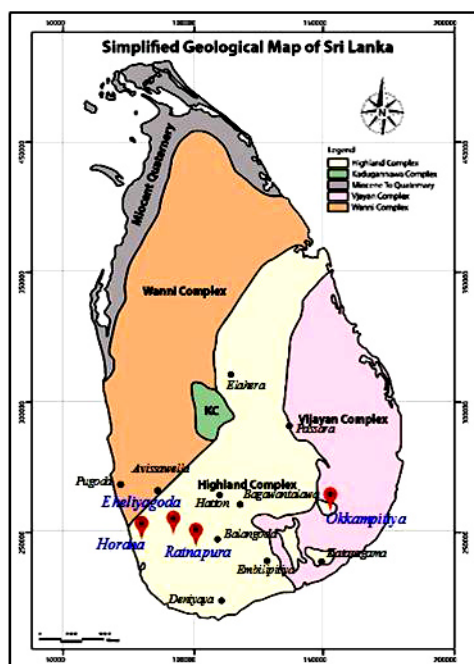


Fig. 1 - Geological map of Sri Lanka with litho-tectonic subdivisions. The locations of the spinel sampling areas used in this study are indicated.

The global distribution of spinel deposits is closely linked to collision, rift and subduction geodynamics. Spinel deposits located in marbles or in neighboring pelitic gneisses are widely known as sources of gem quality spinels. For example deposits in Myanmar, Vietnam, Afghanistan, Tajikistan, Tanzania. The formation of these deposits is explained by metamorphism of Limestones primarily contaminated by aluminous products of weathering, metasomatic transformations of terrigenous layers of marble by metamorphic solution and influx of Al into marbles by endogenic solutions related to alkalic magmatism.



Fig. 2 - Orangey-brown, transparent, irregularly rounded crystal of unknown natural origin hosted by a Horana spinel



Fig. 3 - Colorless-transparent, elongated-prismatic, rounded apatite crystal.

The Importance and Methodology of Undertaking a Literature Search for Gemmological Research and Article Publishing

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If I have seen further... it is by standing on the shoulders of giants. Sir Isaac Newton

Introduction

Performing a literature search is a very important—but often neglected—part of doing gemmological research. As indicated by Davis *et al.* (2012), “Any scientist needs to be familiar with the work that has been done in his or her area of research and with what others are doing concurrently. No scientist, however, can read all the books, articles and other printed matter related to a given area...Therefore, we need to be selective and to find the most pertinent literature on a subject as efficiently as possible and to use it effectively.” A literature review should be conducted *before* starting a research project, in order to avoid “reinventing the wheel” and to keep up with the most current methods for pursuing an experiment, as well as to gather data and information critical to a proper understanding of the subject matter. A literature search involves collecting, cataloguing and evaluating information in the literature in order to determine salient works and refine the subject of one’s research (Machi & McEvoy, 2022). Research findings and other scientific information can be published in books, monographs, proceedings, journals, dissertations, patents, standards, governmental bulletins or reports, and a variety of other sources.

In general, it is best to avoid using unpublished references (although in some cases a personal communication or report from a mining company etc. may be necessary to cite). As a general rule, it is also best to avoid referencing online chat groups/blogs or Wikipedia, due to variations on the quality of the information and lack of expert peer review.

Gemmological Literature Search

Gemmology poses special challenges for undertaking a literature search. Unlike other areas of earth sciences and related fields, there are no comprehensive reference tools for gemmological literature. Therefore, gem researchers will benefit from independently making their own literature collections and databases (typically composed of articles and other works in PDF and hardcopy formats). These can be organised and catalogued using reference management tools such as Endnote, Zotero, Mendeley and Papers, which also allow reference lists to be easily formatted according to numerous bibliographic style templates.

Scholarly journals (e.g. Figure 1) constitute a main source of primary literature (i.e. publications authored by the scientist(s) who personally carried out the research). An exhaustive list of 20 gemmological journals and 10 gem laboratory publications—including information on their time span, accessibility, website, etc.—is available by contacting the author. Only some of them offer indexes or table-of-contents listings that can be searched for keywords, authors, etc. These include *The Journal of Gemmology*, *Australian Gemmologist*, *Revue de Gemmologie A.F.G.* and *Gemmologie: Zeitschrift der Deutschen Gemmologischen Gesellschaft*. (Note that researchers should not omit foreign-language journals from their search, due to the availability of online translators such as Google Translate, DeepL, etc.)

It is highly recommended to download the entire available journal back-issue collections, so that the PDFs are readily accessible even without internet access, while travelling, etc. The PDFs for each journal should be saved into a separate folder. For those journals without indexes or contents

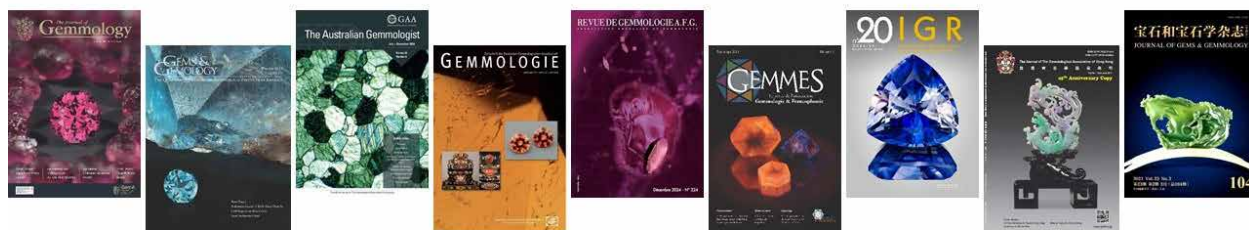


Figure 1: Examples of some scholarly gemmological journals in various languages.

listings, keyword searches can then be done for all files in a particular folder by using the “Advanced Search” function of Adobe Acrobat Pro. (Most PDFs are searchable, but those consisting of scanned images will first need to be processed using Adobe Acrobat Pro’s “Recognize Text” function.) Although more time-consuming than using indexes or contents listings, with this technique it is possible to search for specific content in any collection of PDFs within a single folder, whether from a particular journal or from miscellaneous sources that are grouped together according to gem material, locality, etc.

Other sources of gemmological information include the following:

- Gemmological reference books (e.g. Gübelin & Koivula, 1986, 2005, 2008; O’Donoghue, 2006; Dominy, 2024)
- GIA Library search (<https://g30010.eos-intl.net/G30010/OPAC/Index.aspx>)
- GIA’s historical reading lists (<https://www.gia.edu/historical-gemology-reading-lists>) and bibliographies (<https://www.gia.edu/search?q=bibliography>)
- CIBJO Blue Books and guidelines (<https://cibjo.org/industry-standards-resources>)
- Laboratory Manual Harmonisation Committee (LMHC) Gemstone Information Sheets (<https://www.lmhc-gemmology.org/gemstones>)
- Sheahan Diamond Literature Reference Compilation (<https://secure.kaiserresearch.com/a13/Education.asp?ReportID=364130>)
- For gem localities: John Sinkankas’ textbooks (*Gemstones of North America*, *Emerald and Other Beryls*, etc.); mineral

locality compilations (Bernard & Hyršl, 2015; Moore, 2016); and Mindat (<https://www.mindat.org>)

- For spectroscopic data: Mineral Spectroscopy Server (<http://minerals.caltech.edu>), RRUFF database (<https://rruff.info>) and Spec4Gem (www.spec4gem.info); the first two can be searched to provide reference listings for particular minerals.

Useful general search engines include Google Scholar (<https://scholar.google.com>) and Google Books (<https://books.google.com>); see search tips by Trauth and Sillmann (2018). A relatively new tool is R Discovery, which is an artificial intelligence tool for literature searches (<https://discovery.researcher.life>). In addition, keyword searches can be done using CrossRef (<https://search.crossref.org>), and this website can also be used to look up DOIs (digital object identifiers) for specific articles. DOIs should always be included in reference lists when available.

A tremendous amount of literature (mostly older works that are not under copyright) is freely accessible online via the Internet Archive (<https://archive.org>) and the HathiTrust Digital Library (<https://www.hathitrust.org/the-collection>). The Internet Archive contains resources of particular interest to gemmologists, such as a collection of documents from the GIA Library (<https://archive.org/details/gialibrary>) and gem-and-mineral bibliographies by John Sinkankas, Curtis P. Schuh and Joseph O. Gill. Valuable historical information on gem materials can also be found within archives contained in museums, monasteries, research institutions and government offices.

Local public university libraries are valuable assets for performing literature searches, encompassing numerous resources beyond the scope of this review (see, e.g., Mann, 2005). In addition to obtaining assistance from a reference librarian, it may be possible to use an interlibrary loan service to obtain materials held at other academic libraries. In addition, university libraries commonly have public computers that allow access to online academic databases including “online public access catalogues” (OPACs), WorldCat, GeoRef, Geobase, Web of Science, Scopus, ProQuest, EBSCO, Gale, etc. Importantly, these computers may also have full access to an extensive collection online journals, allowing members of the public to freely download selected articles

and books in PDF format to a flash drive. Local community libraries also have digital collections (although usually not academically oriented) and interlibrary loan services. Another way to obtain access to online journal collections is through membership with the Geological Society (London), which allows members to remotely access content from more than 100 earth science journals.

In conclusion, gemmologists have various options for performing literature searches and electronically accessing journals, books, etc., either using their own computer or by visiting local libraries, even without being a student or member of a university.

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Annex 1: Gemmological Journals^a

Title	Publisher	Language	Years	Access etc.	Website
<i>Australian Gemmologist</i>	Gemmological Association of Australia	English	1958–present	Subscription (hardcopies) or partially open access (html format 2021–2023)	https://www.gem.org.au/current-issue (contents listing available at https://www.gem.org.au/index)
<i>Bursztynisko: The Amber Magazine</i>	International Amber Association (Poland)	English and Polish	1996–present	Open access (online flipbook)	https://www.calameo.com/subscriptions/7391911
<i>Canadian Gemmologist</i>	Canadian Gemmological Assn.	English	1976–2010	Hardcopy back issues (some are still available)	https://canadiangemmological.com/shop/product/the-cga-official-journal-canadian-gemmologist-set-of-4
<i>GemGuide</i>	Gemworld International (USA)	English	1982–present	Partially open access (selected PDFs)	https://www.gemguide.com/news-archive
<i>GEMMAE: An International Journal on Glyptic Studies</i>	LIBRAweb (Italy)	English and Italian	2019–2023	Hardcopy back issues or partially open access (selected PDFs)	https://www.libraweb.net/sommari.php?chiave=142
<i>Gemmes</i>	Gemmologie & Francophonie (France)	French	2023–present	Open access (PDFs)	https://gemmologie-francophonie.com/la-revue-gemmes
<i>Gemmologie: Zeitschrift der Deutschen Gemmologischen Gesellschaft^b</i>	German Gemmological Association	German (and English for latest issue)	1969–present	Subscription (hardcopies or online flipbook since 2005)	https://www.dgmg.com/en/journal-gemmologie.html (author and subject indexes available)
<i>The Gemmologist</i>	N.A.G. Press (United Kingdom)	English	1931–1962	Hardcopies accessible at Gem-A and GIA libraries	None
<i>Gemmology Today</i>	World Gem Foundation (Spain)	English	2016–2024	Open access (non-printable PDFs)	https://www.amazonasgempublications.com/gemmology_today.html (contents listing available at https://www.amazonasgempublications.com/qt_articles.html)
<i>Gemologický Spravodajca / Gemmological Newsletter</i>	Gemmological Institute, Constantine the Philosopher University in Nitra (Slovakia)	English and Slovak	2011–present	Open access (PDFs)	https://www.gu.fpv.ukf.sk/index.php/2-uncategorised/24-gemologicky-spravodajca

^a This listing does not include trade journals or periodicals oriented towards mineralogy, mineral collecting or jewellery. An extensive listing of industry periodicals can be found at <https://www.gia.edu/library-industry-periodicals>. Gemmological publications titled *Diamond News*, *Diamond Review Africa*, *Gamma*, *Gemma News*, *Gemmological Digest*, *Journal of Gems and Precious Metals*, *Portugal Gemas*, *Rough Diamond Review* and *World Diamond Magazine* are not included here due to the relatively limited number of issues that were published while they were active.

^b Prior to 1995, the word *Gemmologie* was not included in the title; from 1952 until 1968 the journal's title was *Zeitschrift der Deutschen Gesellschaft für Edelsteinkunde*.

Annex 1: Gemmological Journals (continued)

Title	Publisher	Language	Years	Access etc.	Website
<i>Gems & Gemology</i>	Gemmological Institute of America (USA)	English	1934–present	Open access (PDFs) or subscription (hardcopies)	https://www.gia.edu/gems-gemology
<i>Gems&Jewellery</i> ^c	Gem-A (United Kingdom)	English	1991–present	Subscription (PDFs) or partially open access (PDFs >2 years old)	https://gem-a.com/publications/gems-and-jewellery
<i>InColor</i>	International Colored Gemstone Association (USA)	English	2006–present	Open access (online flipbook and html format)	https://www.gemstone.org/incolor-bookcase
<i>IGR (Rivista Italiana di Gemmologia / Italian Gemmological Review)</i>	Gem Tech Srls (Italy)	English and Italian	2017–present	Subscription (hardcopies and PDFs) or partially open access (html format)	https://www.rivistaitalianadigemmologia.com/en/home-english
<i>Journal of the Gemmological Association of Hong Kong</i>	The Gemmological Association of Hong Kong	English and Chinese	1979–present	Open access (PDFs from 2008)	http://www.gahk.org/en/journal.asp
<i>Journal of the Gemmological Society of Japan</i>	Gemmological Society of Japan	Japanese	1974–present	Subscription (hardcopies and PDFs) or partially open access (selected PDFs)	https://www.jstage.jst.go.jp/browse/gsjapan (contents listing available at https://doi.org/10.14915/gsjapan.38.4_58)
<i>The Journal of Gemmology</i>	Gem-A (United Kingdom)	English	1947–present	Subscription (hardcopies and PDFs) or partially open access (PDFs >2 years old)	https://gem-a.com/publications/the-journal-of-gemmology (index and bibliographies available online)
<i>Journal of Gems & Gemmology</i>	China University of Geosciences, Wuhan	Chinese	1999–present	Open access (PDFs) or subscription (hardcopies)	http://jogg.cug.edu.cn (search function available at https://jogg.cug.edu.cn/en/to_advance_search)
<i>Pearl Oyster Information Bulletin</i>	Fisheries Info. Unit, Secretariat of the Pacific Community (New Caledonia)	English	1990–2011	Open access (PDFs)	https://fame.spc.int/publications/bulletins/pearl-oyster
<i>Revue de Gemmologie A.F.G.</i>	Association Française de Gemmologie (France)	French	1965–present	Subscription (hardcopies; back issues available to 2013)	https://www.afgems-paris.com/revuesafg (index up to 2022 available at https://www.afgems-paris.com/index-revue-afg)

^c From 1991 until 2005 the journal's title was *Gem & Jewellery News*.

Annex 2: Gem Laboratory Publications

Title	Publisher	Language	Years	Notes	Website
Contributions to Gemology	GRS GemResearch Swisslab (Switzerland)	English	2002– 2014	Open access (PDFs; some issues unavailable)	https://www.gemresearch.ch/research/contributions-to-gemology
Facette Magazine	Swiss Gemmological Institute SSEF	English	1999– present	Open access (PDFs)	http://www.ssef.ch/ssef-facette
Gem Information	Japan Germany Gemmological Laboratory JGGL	Japanese (some English)	2013– present	Subscription (hardcopies) or partially open access (html format)	https://www.sapphire.co.jp/jggl/gi
GIT technical reports	Gem and Jewelry Institute of Thailand	English	2018– present	Open access (PDFs)	https://git.or.th/en/information-service/27/76
ICGL Newsletter	International Consortium of Gem- Testing Laboratories	English	2013– 2016	Open access (PDFs)	https://icglabs.org
Lab Information Circular	IIGJ – Research & Laboratories Centre, India	English	1994– present	Open access (PDFs)	https://qtliaipur.info/lab-information-circular1.aspx
Margaritologia Pearl Newsletter	Elisabeth Strack/ Gemmologisches Institut Hamburg (Germany)	English and German	2014– 2020	Subscription (hardcopies)	https://www.margaritologia.de
News & Events	Elisabeth Strack/ Gemmologisches Institut Hamburg (Germany)	German	2016– 2020	Open access (PDFs)	https://www.gemmologisches-institut-hamburg.de/news-events
Wooshin Gem Lab Magazine	Wooshin Gem Lab (Korea)	English and Korean	2014– current	Open access (PDFs)	http://www.wooshinlab.com/Search.wgk?keyword=%EC%9A%B0%EC%8B%A0%EB%A7%A4%EA%B1%B0%EC%A7%84

An approach to understanding the factors in the emerald enhancement clarity grading system used at CDTECGemlab

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Emerald is the green variety of beryl, with its distinctive color resulting from trace amounts of chromium, vanadium, and iron. These elements replace aluminum in the octahedral positions of its crystal structure. The clarity of an emerald depends on the presence of internal characteristics such as inclusions (solid and/or fluid) and fissures. Since ancient times, the clarity of emeralds has been improved through various treatments (Nassau K. 1984). Among these, fissure filling is the most common procedure, where substances with

optical properties like the host material are used to enhance clarity by improving light transmission within the gemstone. Emeralds may undergo treatments and/or enhancements before, during, and after the cutting process (Fig.1). Sometimes, the application of specific substances helps increase the material's stability, making it more resistant to damage during cutting, especially when large fissures or voids are present (Scarratt K. 2015). In most cases, fissures are filled after the cutting process to improve the gem's appearance.



Figure 1. Possible stages during which the enhancement of the stone can occur throughout its transformation.

Sometimes, an untreated gemstone with visible fissures can undergo treatments or enhancements that lead to a significant improvement in appearance changes that may not be apparent to the final consumer (Fig.2). For this reason, certification is essential, as it provides detailed information about any enhancement treatments, including the extent of the improvement and, in some cases, the type of substances used. Such certification serves as a reliable disclosure to the consumer, offering transparency regarding the gemstone's condition and the nature of any treatments at the time the report was issued.



Figure. 2 Emerald before enhancement (left) and after enhancement (right).

Gemlabs face the challenge of identifying both the type and quantity of substances used in clarity enhancement. This task involves considering various factors, such as identify-

ing the diversity of substances—including vegetable oils, hydrocarbon-derived oils, resins, balsams, and waxes—which can be of natural or artificial origin (Johnson M. *et al*,

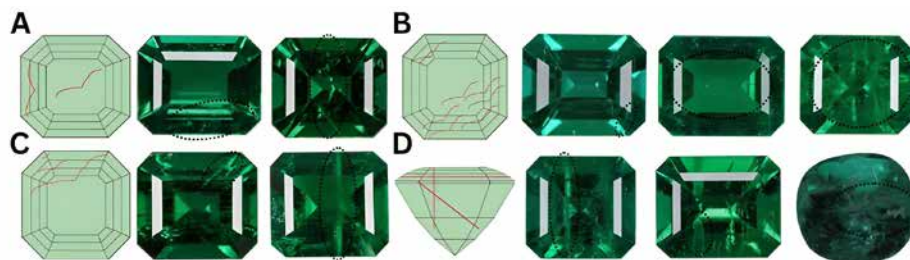


Figure 3. Primary factors used in grading emerald enhancement include, Location (A), quantity of fissures (B), length (C), and tilt (D)

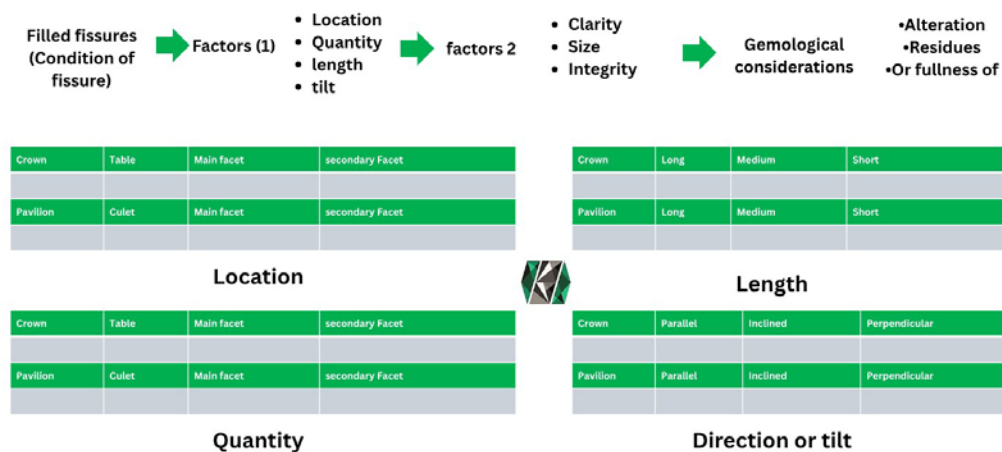
1990; Kammerling R. et al, 1991; Hanni H. 1992; Kiefert L. et al, 1999; Hainschwang & Notari. 2022), as well as the mixtures employed. Although cedar oil was historically a commonly used product, it is now less prevalent. Currently, substances such as natural hydrocarbon-derived oils—commonly known as mineral oil (e.g., baby oil)—and liquid or hardened artificial resins are predominantly used.

Additionally, it requires developing an assessment as objective as possible of the extent of clarity modification by fissure filling through qualitative analysis. The assessment of the extent or intensity of clarity improvement has been addressed by Shane McClure *et al.*, 1999, who classified the enhancement into three main categories: minor, moderate, and significant. This classification is based on factors such as the size, extent, or length of the treated areas; the number of fissures; the amount of treatment within fissures; and the location of filled fissures. The description of emerald enhancement is provided in Sheet # 5 (LMHC, 2014), which outlines a guide for determining the amount of filled fissures. This serves as the basis for quantifying the enhancement and assigning a corresponding quality grade.

However, one of the most important aspects of classification is the ability to identify fissures or voids containing substances. To achieve this, various methodologies have been developed that utilize different instruments and illumination techniques, along with careful inspection of various parts of the stone. These include transmitted light (bright field - dark field), reflected light, and UV light. Regarding UV illumination, caution should be exercised with certain UV ranges, as they may cause harmful effects on substances, such as polymerization. Additionally, it is important to remember that not all substances fluoresce, for example, some oils (Notari F. *et al.*, 2022). This comprehensive approach ensures the collection of complete information necessary for an accurate and reliable classification process.

The purpose of grading is to assess the extent of improvement in the clarity of the stone. While the number of fissures present may be important, it does not necessarily determine the degree of improvement. For example, a single fissure that is oriented parallel to the table and covers a large portion of the stone can result in a moderate grade because it significantly affects the stone's clarity. Conversely, several small fissures that are perpendicular to the table, located on the crown facets (not the table), and aligned with the edges of the crown facets, especially if they are small relative to the stone's dimensions—may warrant a minor grade.

Fissures typically behave as three-dimensional features, either resembling a plane or acting as pores or hollows within the stone, such as in emeralds. This means they possess volume. Factors such as their location, length, quantity, and inclination are highly important for determinate improve clarity. In certain areas of the stone, the presence and arrangement of fissures may have a greater impact on clarity than in other regions. Additionally, if a wide fissure is in a region that could affect the stone's integrity—such as on the pavilion, girdle, or crown—its significance increases. For example, a fissure that crosses the stone completely or is blunt can lead to chipping or even a fracture that splits the stone into two nearly equal parts (integrity factor). Another important consideration is size. It is rare to find stones over 5 carats—or even over 10 or 15 carats—that are free of fissures. However, a stone with several fissures that appear dense relative to its small volume does not necessarily indicate a similar density in a much larger stone (size factor). The clarity factor provides a general measure of the impact of fissures on the stone's overall appearance. If a stone has high clarity, filled fissures will be more visible. In such cases, the presence of fissures may be more significant in an untreated stone with high clarity than in one with lower clarity, where the abundance of inclusions makes fissures less noticeable (clarity factor). Other factors may also influence



Not indication – Insignificant - Minor (F1) - Minor to moderate (F1 - F2) - Moderate - Significant (F3)

Figure 4. Grading emerald enhancement system by CDTEC

the grading based on gemological considerations. These include residues, alterations of substances, or any other features within the stone that could impact its clarity. Such features should be identified, described, and evaluated to determine their effect on the stone's overall clarity.

To define the factors that enable the improvement of quality grade assignments related to enhancement and/or

treatment is an ongoing task for gemological laboratories, which helps foster consumer confidence. However, one of the objectives of this work is to contribute to the enhancement of methods that promote consistency in laboratory procedures. This includes supporting aspects such as record-keeping, traceability, and repeatability, all within the framework of international standards—not only for the gemstone industry but also for testing and analysis laboratories.

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Noble gas in corundum: interests in gemology

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Noble gas in geology

Noble gases—elements of the last column of the periodic table—are chemically defined by their full outer electron shells, making them highly inert and very volatile under ambient conditions. In geology, they are used to trace a wide range of processes, including mantle circulation and degassing, hydrocarbon reservoir formation, climate

change, and ocean circulation (Burnard, 2013). Among the noble gases the lightest (helium, neon and argon) are the most commonly used and possibility present in gems. Their lack of chemical interaction makes them useful in gemology for two main purposes: detecting heat treatment and determining geographic origin.

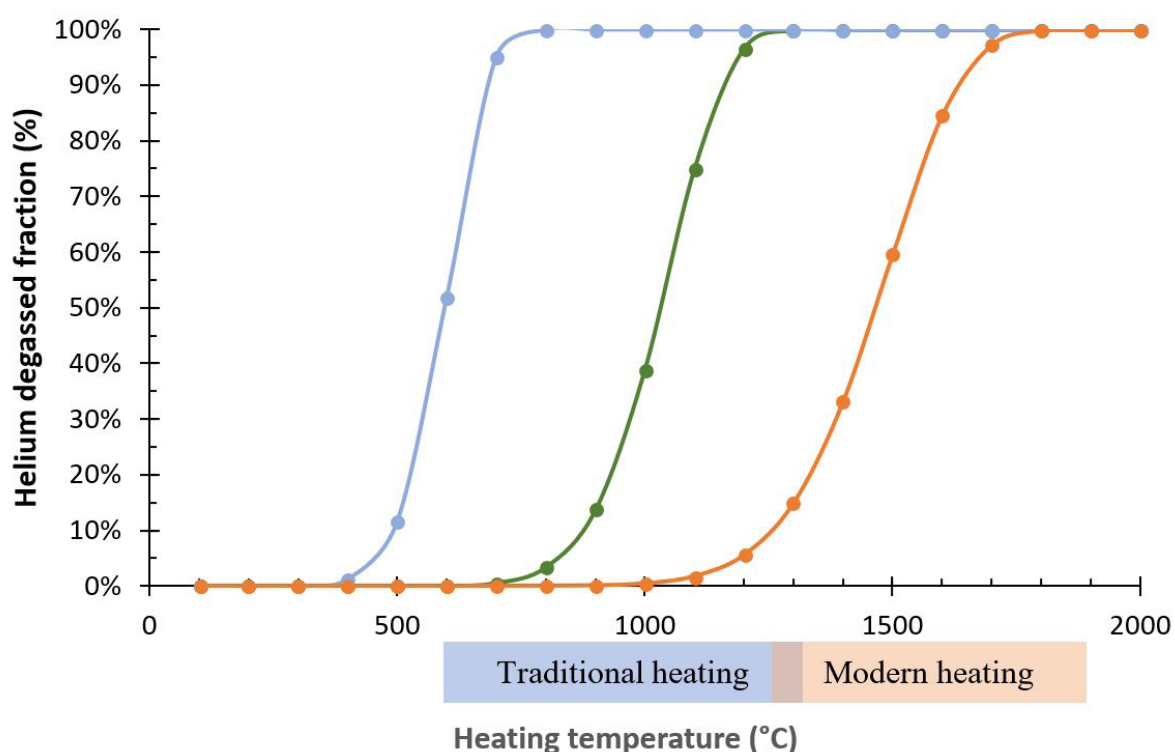


Figure 1 : Fraction of helium degassed during heating according to theoretical data from Zhang *et al.* 2016, with activation energy of 100 (blue), 150 (green) and 200 (orange) kJ/mol, calculated from the equation from Fechtig and Kalbitzer (1966).

Heat-treatment

Traditional heat treatment (~600°C–1200°C) is one of the oldest known gemstone enhancements, used over 4,000 years ago for improving corundum color (Notari *et al.*, 2019). Modern treatments reaching up to ~1850°C leave detectable traces, while low-heat methods remain to date an important challenge to identify and spur further research (Saeseaw *et al.*, 2021; Hughes & Vertriest, 2023; Karampelas *et al.*, 2023).

Due to their volatility, noble gases—especially helium—could serve as scientific indicators of heat treatment. Quantifying diffusion of noble gases in crystals is then necessary to use these volatiles as tracers. Theoretical models and experiments on synthetic materials suggest that helium diffuses out of the crystal between 500 °C and 1700 °C, depending on the nature of the defects in which it is trapped and the duration for which the temperature is maintained (*e.g.*: Campbell, 1960; Fechtig & Kalbitzer, 1966; Zhang *et al.*, 2016). The diffusion of noble gases is highly dependent on the local environment of diffusion, as demonstrated in high-density SiO₂ polymorphs (Figowy *et al.*, 2024). As noble gases easily may easily diffuse as low as 500°C, the absence of helium in natural corundum may indicate human-induced heating, potentially calling for a reassessment of current heat treatment detection methods. Ongoing experiments on six samples treated at vari-

ous temperatures aim at better constraining the conditions required for helium degassing. In parallel, new *ab initio* molecular dynamics calculations are performed and compared with experimental results to provide a quantification of diffusion processes and investigate the role of defects.

Origin

Noble gases are commonly used as tracers of geological processes, and their analysis may represent a novel approach for determining the origin of corundum. To date, no data on noble gases in natural corundum have been reported in the literature. To explore this potential, we measured the helium concentration from 116 crystals from 14 different sources. Our results show that sapphires from Kenya, Greenland, and Madagascar have significantly higher concentrations (>100 pmol/g) compared to those from other localities (<30 pmol/g), including Colombia, Vietnam, Mozambique, Kashmir, France, Sri Lanka, Cambodia, Australia, Myanmar, Tanzania, and Thailand. While most conventional provenance methods discriminate between deposit types (magmatic and metamorphic type), it seems that helium content does not directly correlate with the deposit type. While not immediately applicable to the gem trade yet, noble gas analysis could serve as a complementary scientific tool in the study of gem geology and origin determination.

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Gem zircon and sapphire age dating and application of origin determination: A study from New England sapphire fields, New South Wales, Australia

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Abstract

Chemical composition and U-Pb age determined by laser ablation (LA) ICP-MS are presented for zircon megacrysts found in alluvial gem corundum deposits associated with alkali basalts in the Inverell district in New England, New South Wales, eastern Australia. The three mine localities, Kings Plains, Swan Brook, and Mary Anne Gully, produce gem-quality brown to yellow zircon megacrysts characterized by relative enrichment in Hf and REE. We observed no differences in relative concentrations of transition metals in relation to color. Chemical homogeneity within the single zircon crystals indicates stable crystallization conditions. The 206Pb/238U age of zircon megacrysts from these three localities fall into two groups, an older group of about 174–216 Ma and a younger group of about 37.7–45 Ma, respectively. Based on our data, the zircons in the New England alluvial gem deposits may be related to two formation episodes, one in the Upper Triassic–Lower Jurassic and one in the Eocene (late Oligocene). Most originated from the lithospheric mantle, and later hosted basaltic magmas brought up. Gem quality blue sapphires are also directly collected from these mines, and zircon inclusions in Kings Plains and Mary Anne Gully sapphires gave U-Pb ages of about 35 to 37 Ma which fall within the range of basalt K-Ar ages of 19 to 38 Ma and close to the younger group of zircon megacrysts (37.7–45 Ma) thus indicating the timing of volcanism of the Inverell district–New England field.

Introduction

Zircon megacrysts are common within eluvial, paleo-alluvial, and alluvial sapphire/ruby deposits derived from basaltic sequences, with gem properties suitable for cutting as an accessory gemstone. Such zircons and sapphires/rubies are widely distributed along eastern Australia (Figure 1). The largest gem field in New England, New South Wales, has a limited systematic spread of U-Pb ages (Coenraads *et al.*, 1990; Sutherland & Fanning 2001; Sutherland *et al.*, 2002; Zaw *et al.*, 2006; Abduryim *et al.*, 2012). In 2010, the authors visited sapphire mines in Kings Plains, Mary Ann Gully, and Swan Brook and investigated the geological distribution and mining at each deposit. This study presents U-Pb age dating of zircon megacryst and zircon inclusion in host sapphire from the Inverell district in the New England gem field. These data are used to characterize and discuss the crystallization ages and source affinities (or genesis) of gem-quality zircon and sapphire megacrysts from a world-class basaltic gem field.

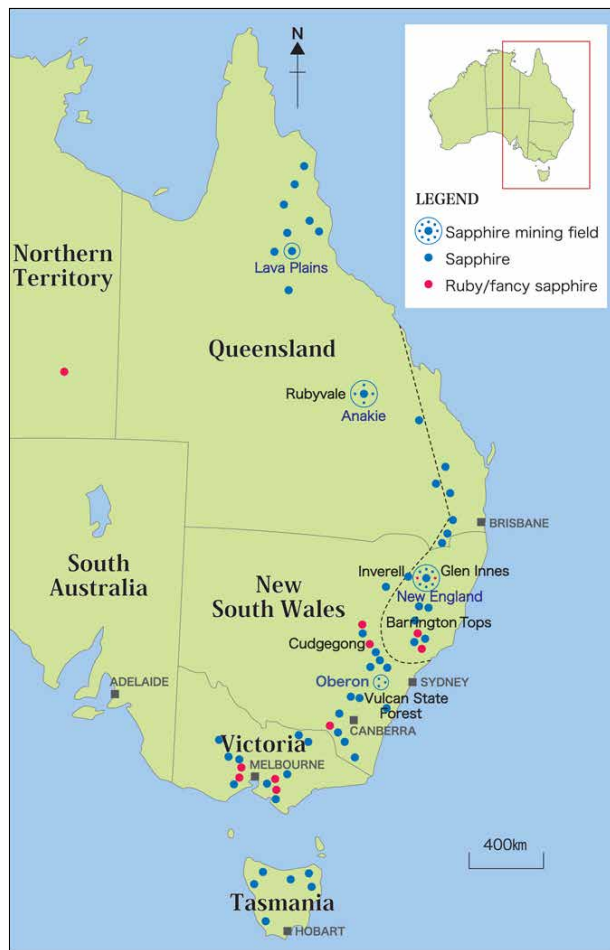


Figure 1. Distribution of sapphire/ruby and zircon in eastern Australia, showing mining locations of the New England, New South Wales, Anakie and the Lava Plains gem fields in Central and Northern Queensland, other sapphire/Ruby gem fields in Victoria and Tasmania.

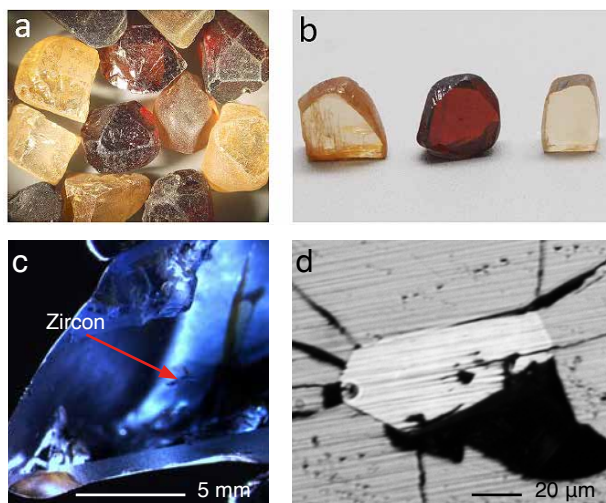


Figure 2. Zircon megacryst concentrate (a: left top) from Kings Plains mine. Zircon megacryst color range with polished surface for analysis (b: right top, sample weight, 2.46–4.11cts). (c) Blue sapphire from Mary Anne Gully (sample S-MA-0001) with a small zircon inclusion surrounded by tension cracks. (d) Zircon inclusion in sapphire (sample S-MA-0001) which was carefully polished to expose the zircon to the surface for LA-ICP-MS analysis.

Materials and Methods

Zircons from the Inverell district-New England gem deposits selected for analysis included 7 brown and yellow zircon megacrysts collected from the alluvial corundum mine in Kings Plains, 3 from the Mary Anne Gully mine and a color range of 5 faceted megacrysts from Swan Brook (Figure 2a and b).

125 samples of blue and greenish blue sapphire megacrysts were collected from the exact three mine locations. They show a glossy natural surface which is typical as a result of magmatic dissolution and etching. Two blue sapphire samples contained several large transparent euhedral zircon inclusions, and these sapphires were carefully orientated and polished to expose the zircon inclusion to the surface of the sapphire (Figure 2c and d). Their size (100 and

80 μm in length), was suitable for dating with conventional LA-ICP-MS (Agilent 7700 ICP-MS with 213nm YAG laser).

Results and Discussions

Chemical Composition: Zircon megacrysts from the Inverell district are significantly HREE-enriched over LREE (Figure 3). The chondrite-normalized REE patterns are moderately fractionated and show a positive Ce anomaly. Only a few samples show negligible Eu anomalies. The total content of LREE in the Inverell zircons is 0.01 to 0.3 (La) up to 20 to 300 (Dy) chondrite units. HREE content ranges from 30 to 500 (Ho) to 100 to 1250 (Lu) chondrite units. The REE patterns of the zircons from three localities in the Inverell district sapphire mine field have similar profiles, indicating related origins.

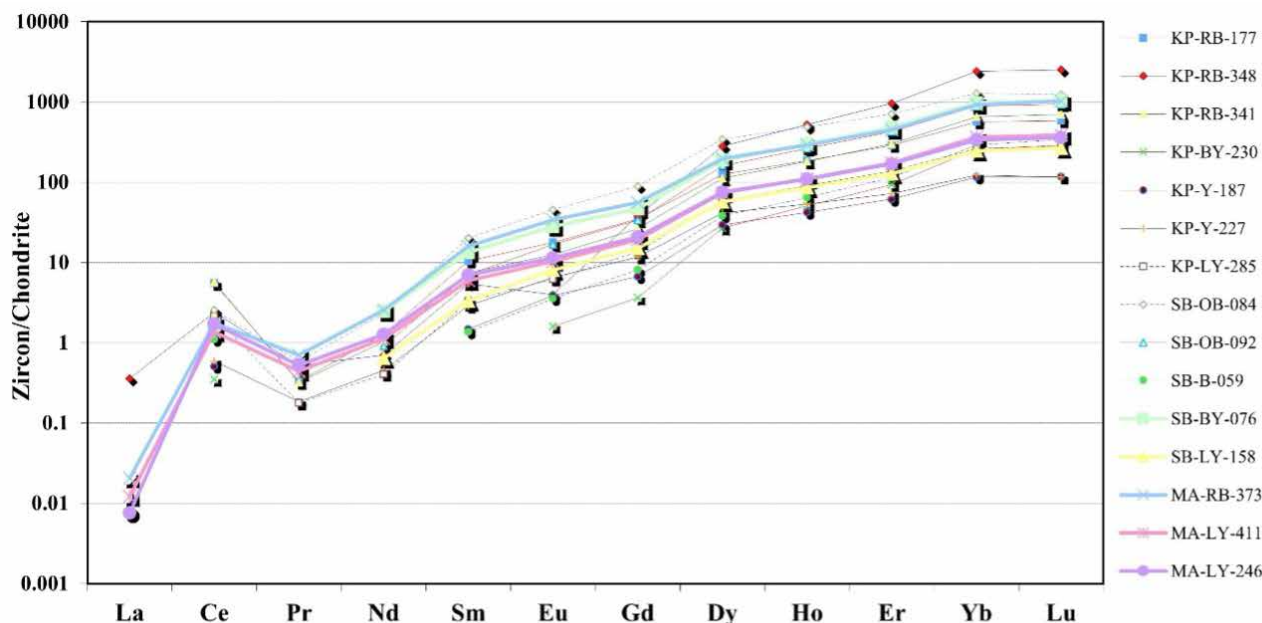


Figure 3. Chondrite-normalized REE patterns of zircon megacrysts from Kings Plains, Swan Brook, and Mary Anne Gully in the Inverell district, New England gem field, Australia.

The blue, yellow-blue (parti-color type) and greenish blue sapphires from three mine locations in Inverell district show relatively high Ga (> 120 ppm), low Mg (< 18 ppm) and low Cr (< 10 ppm), features that typify magmatic sapphires. The blue sapphire group, however, shows lower Fe, V, and Mg and higher Ti than the greenish blue group. Other minor trace elements such as Ni, Sn, and Ta values are mostly <1 ppm in blue and yellow-blue sapphires and below detection in greenish blue sapphires.

Geochronology: Six samples among the 7 zircon samples from Kings Plains reveal older ages, from the U-Pb concordance of $^{206}\text{Pb}/^{238}\text{U}$ age around 188.9 ± 4.5 Ma, 185.9 ± 3.3 Ma, 208 ± 4 Ma, 176 ± 6.1 Ma, 174 ± 7 Ma, and 207 ± 17 Ma at the 95% confidence level. One zircon gave a younger 45 ± 0.52 Ma age (Figure 4). Based on these results we assume that the deposits may represent two different volcanic activity periods, one in the upper Triassic and later in the late Oligocene.

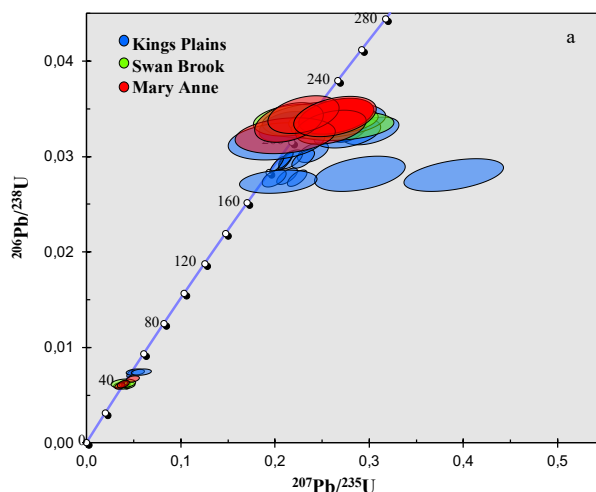


Figure 4. Concordia diagram showing U-Pb age data of zircons from Kings Plains, Swan Brook and Mary Anne Gully from the Inverell district, NSW, Australia.

Four samples among the 5 Swan Brook zircons show a younger concordant age around 37.94 ± 0.34 Ma, 37.22 ± 0.28 Ma, 36.2 ± 6.4 Ma, 37.66 ± 0.35 Ma at the 95% confidence level, except one zircon grain with an older U-Pb age of 215 ± 2 Ma.

This trend of two age groups is also found in zircons from the Mary Anne Gully alluvial deposit. Two samples resulted in a concordant age of 206 ± 11 Ma and 216 ± 5.7 Ma, while one zircon shows a concordant age of 37.7 ± 0.9 Ma.

Most of the zircons from the Kings Plains alluvial deposits belong to the older age group (the weighted mean $^{206}\text{Pb}/^{238}\text{U}$ age of 174 Ma to 216 Ma) and suggest its main volcanic eruption episode was in Triassic to Jurassic, but Swan Brook zircons are dominantly the younger age group (37.7 Ma to 45 Ma), relating them to the Eocene Maybole volcano eruptions.

In the blue sapphire sample from Kings Plains, two spot analyses were obtained on the zircon inclusion carefully polished to be exposed at the surface. The U-Pb concordant ages of this zircon inclusion were 37.2 ± 1.9 and 36.9 ± 1.6 Ma (Figure 5). Another zircon inclusion in the Mary Anne Gully sapphire gave similar concordant ages of 35.7 ± 2.1 and 34.9 ± 1.4 Ma, revealing that there is no significant difference between these two mine locations in terms of formation ages. These results show that the sapphires originated during the Eocene volcanism in the Inverell district. In addition, the age of the zircon inclusions in sapphires fully overlaps with the young age of the zircon megacrysts from the same locations, indicating that the zircon inclusions probably formed together with the sapphires or shortly before. Based on literature and our data, it is assumed that the sapphires formed from strongly evolved magmas in the deep crust or upper mantle and that they were carried up in large quantities within alkali basaltic magmas during volatile-rich and explosive volcanic eruptions.

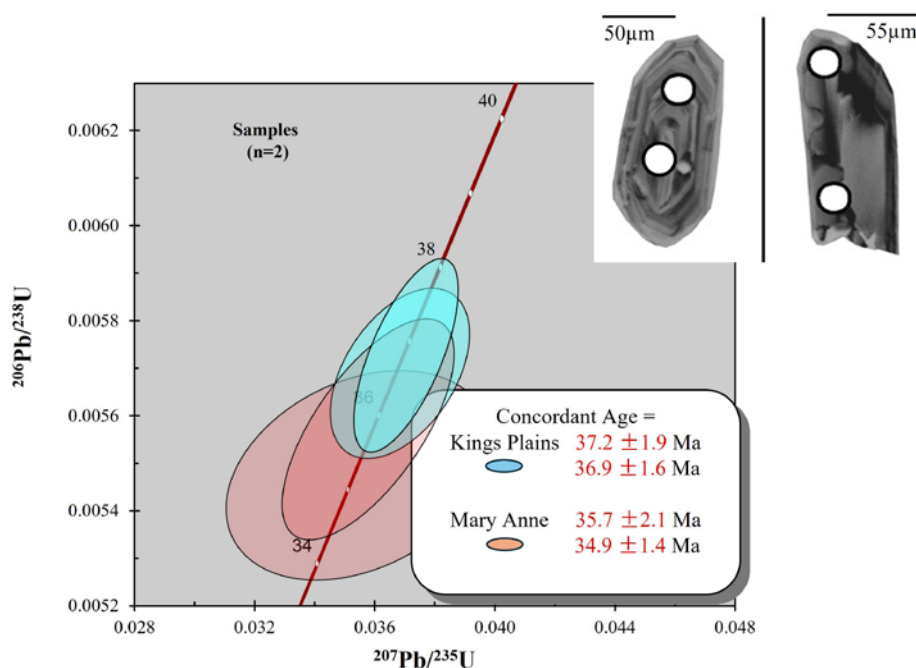


Figure 5. Concordia diagram for the zircon inclusion in sapphires from Kings Plains and Mary Anne Gully.

Geographic Origin Determination of Natural Yellow Sapphires: The Preliminary Study

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Keywords: Yellow sapphire, Geographic origin, Geochemical fingerprinting

Introduction

Determining the geographic origin of yellow sapphires plays a critical role in the gem trade, as origin often dictates both perceived quality and market pricing. Yellow sapphires, a popular variety of corundum, form in diverse geological environments, each imparting unique internal features and chemical profiles. In today's market, where ethical sourcing and traceability are increasingly prioritized, reliable methods for origin determination are essential. Traditional gemmological approaches, such as analyzing inclusions, can offer helpful insights but are often insufficient—particularly when stones have undergone heat treatment or when inclusions appear similar across different mining regions. This research seeks to overcome these challenges by analyzing yellow sapphires from importantly global sources, including Sri Lanka, Madagascar, Tanzania, and Thailand. Through detailed geochemical profiling, the study aims to uncover distinguishing chemical traits that can serve as dependable indicators of geographic origin. The goal is to improve precision in origin determination, thereby reinforcing

consumer trust, enhancing valuation, and promoting transparency across the gemstone supply chain.

Materials and Methods

In this investigation, a total of 151 yellow sapphire samples were analyzed. These originated from Sri Lanka (36 samples), Ilakaka, Madagascar (22), Tanga, Tanzania (45), and Bang Kacha, Chanthaburi, Thailand (50), with weights ranging from 0.23 to 4.30 ct and displaying various yellow colour tones (Figure 1). All samples had undergone traditional heat treatment, except for the Thai samples, which were treated using beryllium heat treatment. The gemmological properties of these samples were examined using both standard gemmological tools and advanced analytical instruments, including Raman (Renishaw inVia Qontor Raman microscope), UV-Vis-NIR (PerkinElmer Lambda 1050), and LA-ICP-MS (LA: ESI Industrial NWR-213, ICP-MS: Thermo-Scientific iCAP™ RQ).



Figure 1: Representative samples of yellow sapphire from Sri Lanka, Madagascar, Tanzania, and Thailand ranging in weight from 0.23 to 4.30 ct (Photograph by M. Seneewong-Na-Ayutthaya)

Results and Discussions

Internal features.

Sri Lankan yellow sapphires are frequently characterized by rectilinear, zigzag-shaped fingerprint inclusions (Figure 2a) and CO₂-filled negative crystals (Figure 2b) (Palke *et al.*, 2019). Additional commonly observed internal features include silk, various fingerprint patterns, needle-like inclusions, minute particles, milky clouds, twinning, as well as crystalline inclusions such as apatite and graphite. In contrast, yellow sapphires from Ilakaka, Madagascar typically contain abundant rounded zircon inclusions (Figure 2c) (Saeseaw *et al.*, 2020). Other diagnostic features in these samples include rutile needles (Figure 2d), silk, complex fingerprint patterns, iron staining, and mica inclusions. Tanzanian yellow sapphires are often distinguished by the presence of hematite silk (Figure 2e) and rose channels (Figure 2f).

(Figure 2f). Additional inclusions observed in these stones include iron staining, hematite, zircon, amphibole, apatite, rutile, black inclusions, various crystalline phases, fingerprint patterns, silk, needles, clouds, and lamellar twinning. Notably, the zircon inclusions in Tanzanian sapphires may closely resemble those from Madagascar, complicating origin determination based solely on inclusion characteristics and necessitating further analytical investigation. Thai yellow sapphires frequently exhibit dissociated inclusions, particularly following high-temperature heat treatment (Figure 2g), along with angular bands of brownish particles (Figure 2h) (Saeseaw *et al.*, 2017). Other internal features include feldspar, crystals, negative crystals, minute particles, clouds, blue colour zones, fingerprint patterns, silk, needles, and iron staining.



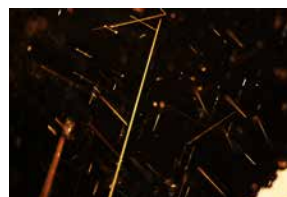
a) Rectilinear zigzag fingerprint in Sri Lankan stone



b) CO₂-filled negative crystals in Sri Lankan stone



c) Cluster of rounded zircon inclusions in Madagascan stone



d) Rutile needle-like inclusions in Madagascar stone



e) Hematite silk in Tanzanian stone



f) Rose channels in Tanzanian stone



g) Dissociated crystal (heat treatment) in Thai stone



h) Angular bands of brownish particles in Thai stone

Figure 2: Inclusions observed in yellow sapphire samples from various deposits examined in this study; field of view 3.8 mm (a, d, e, g), 2.8 mm (b), 1.8 mm (c), 4.5 mm (f) and 9 mm (h). (Photomicrographs by M. Seneewong-Na-Ayutthaya)

UV-Vis spectra.

Yellow sapphires exhibit characteristic UV/Vis/NIR spectra, with their colouration primarily attributed to two distinct mechanisms: colour centres ($h\nu$ - Fe^{3+}) and the presence of the iron (Fe^{3+}) trace element (Dubinsky *et al.*, 2020). Figure 3 illustrates the absorption spectrum of a yellow sapphire from Sri Lanka, which is representative of the colour produced predominantly by the colour centre chromophore. In contrast, for yellow sapphires with higher iron concentra-

tions, such as those from Madagascar, Tanzania, and Thailand, their spectra are characterized by absorption features at approximately 378 nm and 450 nm attributed to $\text{Fe}^{3+}/\text{Fe}^{3+}$ pairs, and an absorption band at approximately 387 nm indicative of isolated Fe^{3+} (Emmett *et al.*, 2023; Dubinsky *et al.*, 2020). These spectral features confirm the dominant role of iron (Fe^{3+}) as the primary chromophore for yellow colouration in these latter groups.

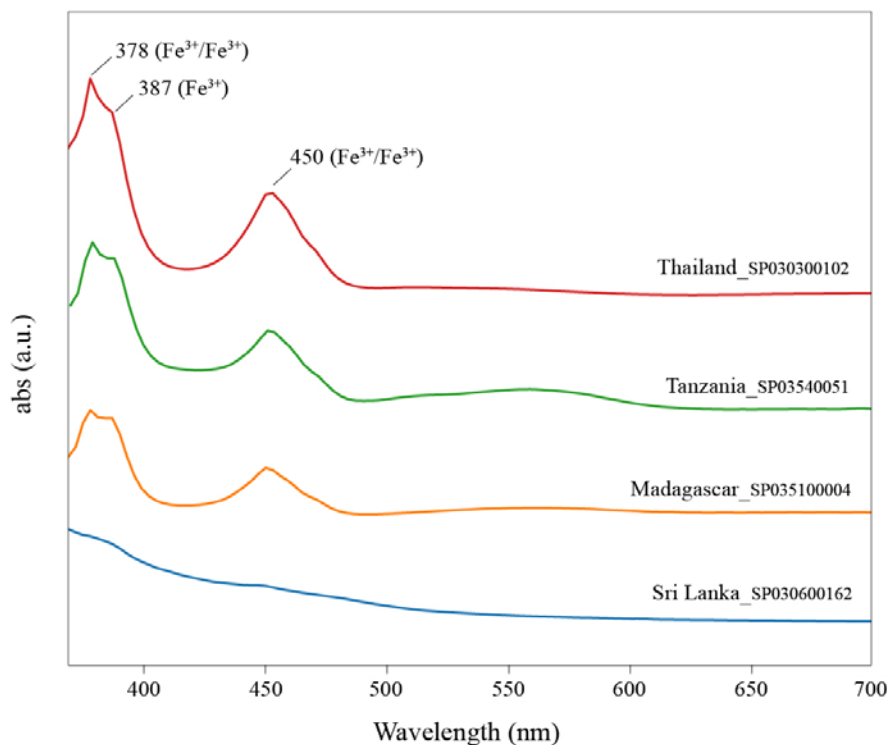


Figure 3: UV-Vis-NIR spectrum of yellow sapphire samples from various deposits examined in this study

Chemical analysis.

Chemical composition analysis of yellow sapphire samples using LA-ICP-MS revealed the presence of important trace elements, including Fe, Mg, Ga, Ti, Cr, and V. Yellow sapphires from Thailand exhibited the highest average Fe content (~15,000 ppm), followed by samples from Tanzania (~7,800 ppm), Madagascar (~4,500 ppm), and Sri Lanka (~440 ppm). The relatively high Fe content in Thai is attributed to their basalt-related geological origins, in contrast to Sri Lankan sapphires, which are typically associated

with metamorphic rocks. As shown in Figure 4, 2D (Fe–Cr) and 3D (Fe–Mg–Cr) scatter plots clearly illustrate the differentiation of Sri Lankan yellow sapphires from those of other origins. Distinct clustering patterns were observed for each locality, enabling effective discrimination among samples from Sri Lanka, Madagascar, Tanzania, and Thailand. Among the trace elements, Fe, Mg, and Cr are identified as a key indicator for determining the geographical origin of yellow sapphires.

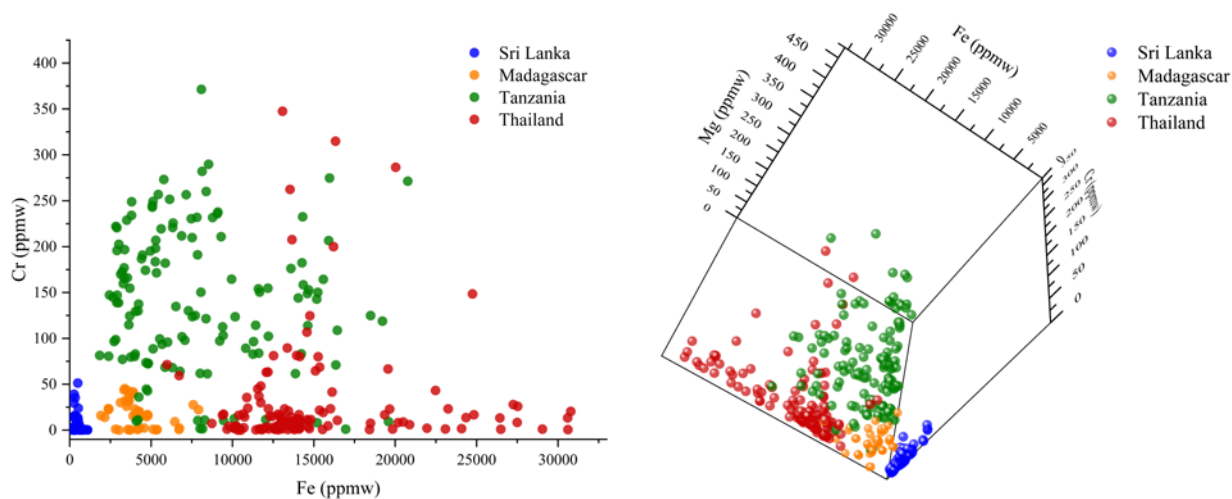


Figure 4: 2D and 3D scatter plots of trace element contents in yellow sapphires in this study.

Conclusion

This preliminary study demonstrates the potential for determining the geographic origin of yellow sapphires using a combination of various analytical techniques. The results highlight the ability to distinguish samples from Sri Lanka, Madagascar, Tanzania, and Thailand. Scatter plots of Fe-Cr and Fe-Mg-Cr concentrations show good clustering patterns, with Fe, Mg and Cr emerging as a key geochemical marker that reflects differences in geological formation. The

integration of multiple methods, including inclusion observation, chemical analysis, and spectroscopic techniques, further enhances the accuracy of origin identification. For future research, increasing the sample size and incorporating a broader range of sample types and origins, colour variations, and treated stones from each locality will help strengthen and validate these findings.

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Rubies from Greenland

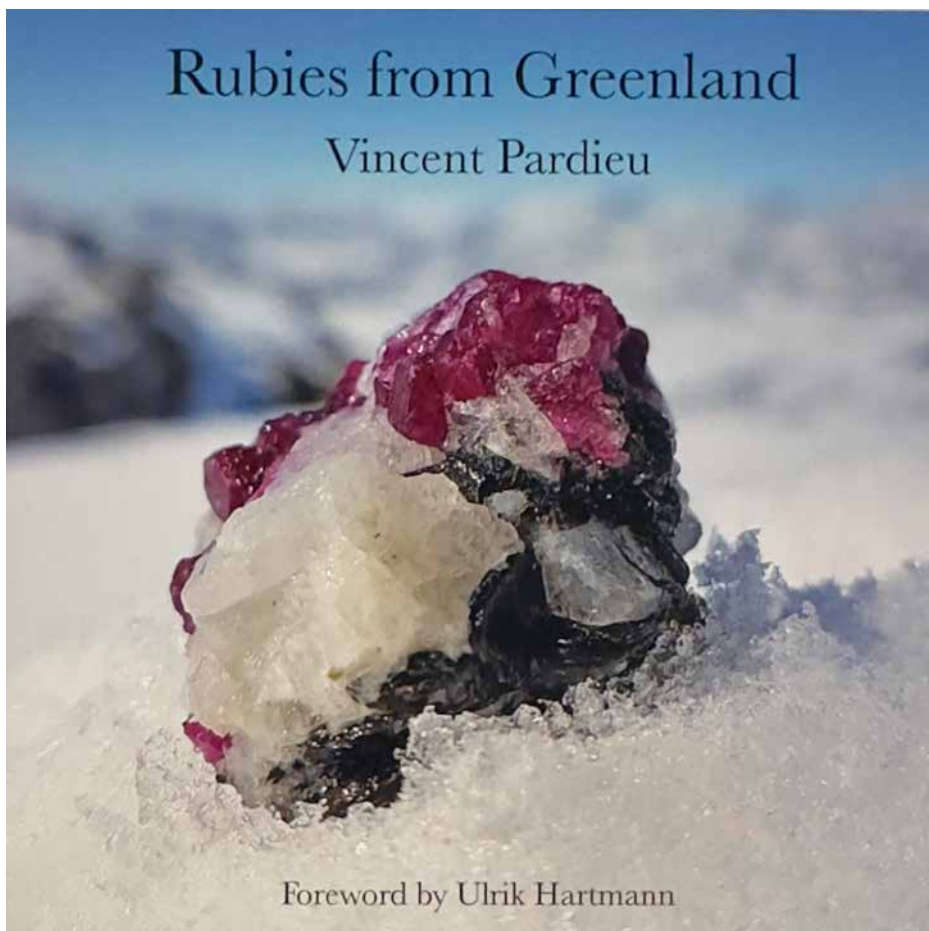
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“Rubies from Greenland – A Field Gemology Travelogue” a book by Vincent Pardieu

Gemology is traditionally practiced in laboratories, focused on identifying and analyzing gemstones. Vincent Pardieu’s journey, however, took a different path. In 2005, he coined the term *field gemology* to describe the practice of studying

gems at their source—in mines, markets, and remote locations. Since 2004, he traveled the world collecting research samples, carefully documenting each using what he calls the “Four Ws”: When, Where, From Whom, and How it was collected. As Sir David Attenborough aptly noted, without context, a specimen loses much of its scientific value (Pardieu, 2020).



Beyond collecting reliable reference samples for gemological research, the author aims to capture the full story—through photographs of landscapes, miners, and the often-overlooked realities of extraction. While most in the gem trade focus on the polished final product, the origin of a gem profoundly influences its scientific, commercial, and cultural significance. This book, *Rubies from Greenland – A field gemology travelogue*, offers a visual and narrative account of the eleven visits Vincent Pardieu had to the Aappaluttoq ruby mine in Greenland. It follows the journey of these rubies from the mine to Bangkok—where they are heat-treated, cut, and polished—and finally to Denmark, where Hartmann's sets them into fine jewelry, some of which is now worn by Queen Margrethe II.

Rubies were first discovered in Greenland in 1965 (Herd & al, 1969). In the early 2000s, commercial interest grew, eventually leading to the development of a mining project. VP was invited in 2017 as a consultant when the mine officially began production. Remarkably, the rubies from Greenland belong to the same geological type as those from East Africa (Madagascar, Tanzania, Mozambique, and Malawi), associated with amphiboles—deposits that have now overtaken marble- and basalt-related sources as the

primary origin of rubies in the global market (Giuliani *et al*, 2020). Yet, aside from Greenland, these amphibole-related ruby deposits remain under-researched. Beyond the scientific value, VP was struck by the paradox of a modern, high-tech mining operation in such a remote, untouched landscape. He proposed conducting heat-treatment experiments in Thailand to enhance the rubies' suitability for the jewelry market. The successful results led to the establishment of a Bangkok office and the development of traceability software that tracked each gem from mine to market. From 2017 to 2022, VP worked closely with the Aappaluttoq team—offering training, technical advice, and documenting the process through photography and video. Despite significant achievements, the mine struggled with challenges, especially during the COVID-19 pandemic (Pardieu, 2021). It ceased operations in 2022, and in 2024, Greenland Ruby declared bankruptcy.

This book is a tribute to those five unforgettable years. With hundreds of photos over 280 pages, it is visual reference on an extraordinary project. It offers a unique window into the complex and often secretive world of gemstone sourcing; sheds light on Greenland's mining potential and logistical challenges and honors the people who brought to life what are now recognized as the oldest known rubies on Earth.

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Blue sapphire from Malacacheta of Minas Gerais, Brazil

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Malacacheta, located in Minas Gerais, Brazil, is renowned for its alexandrite deposits and complex geological setting. Mining of alexandrite began in 1975, peaked around 1987, and continued into the 2000s. Brazilian cat's eye alexandrite from this locality remains available in the market today. In addition to alexandrite, sapphires of various colors have also been reported from Malacacheta (Liccardo *et al.*, 2006). However, sapphire has not been the primary focus of mining activity. These sapphires are recovered from the same alluvial deposits as alexandrite and are estimated to have formed in a metasomatic environment.

In this study, 13 faceted blue sapphire samples (Figure 1) from Malacacheta, commercially available on the market, were analyzed. According to the supplier, the sapphires were recovered together with alexandrite as a byproduct. The samples ranged in size from 0.35 to 0.69 ct and displayed colors ranging from light blue to vivid violetish blue, with some exhibiting visible color zoning. All samples were transparent, although some contained visible inclusions.

The sapphires exhibited gemological characteristics consistent with typical blue sapphire. Re-fractive indices ranged from 1.76 to 1.77, and specific gravity ranged from 3.98 to 4.00. Pleochroism ranged from pale to vivid blue. The samples were inert to both long-wave and short-wave UV radiation.

About microscopic features, most samples showed strong strain and color zoning (Figure 2). Comet tail-like inclusions were observed in some samples (Figure 3). Numerous samples contained crystal inclusions identified as plagioclase (Figure 4). Other mineral inclusions were rare; only one sample exhibited carbon and calcite inclusions within a surface-reaching cavity. Raman spectroscopy of a negative crystal inclusion (Figure 5) revealed CO₂ peaks at 1279.8 and 1384.6 cm⁻¹ (Frezzotti *et al.*, 2012). Additionally, no two phase inclusions were found in the samples.



Figure 1. Tested samples of blue sapphire from Malacacheta



Figure 2. Strong strain with colour zoning



Figure.3 Comet like inclusions

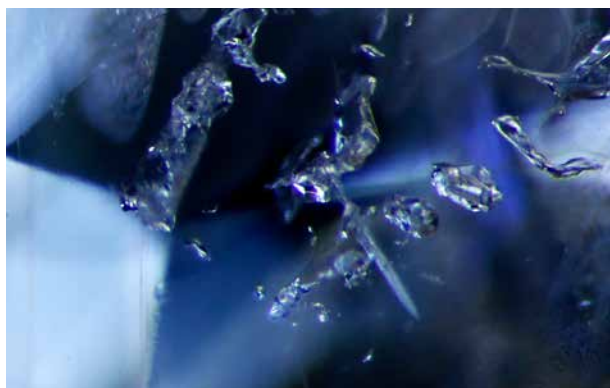


Figure 4. Plagioclase inclusion

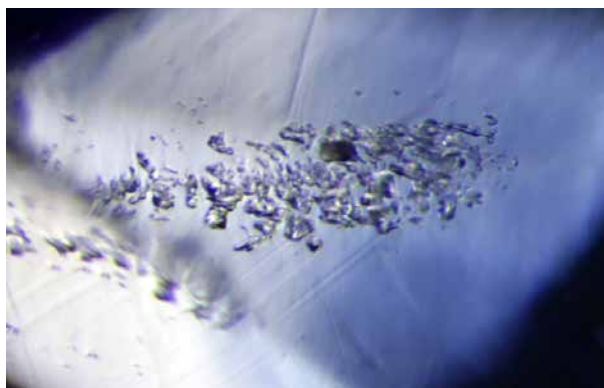


Figure.5 Negative inclusions tested with Ra-man

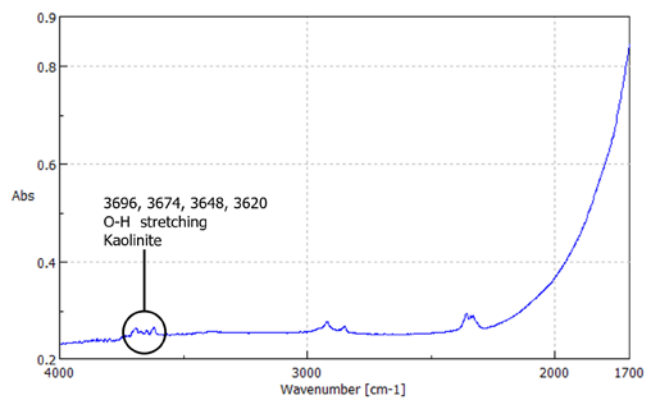


Figure 6. FT-IR spectrum with kaolinite absorption

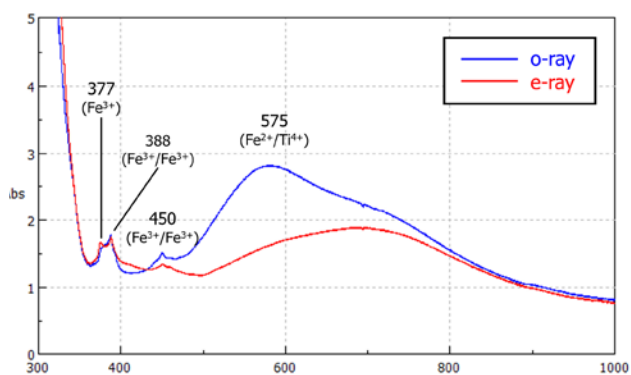


Figure.7 Polarized UV-Vis spectra

Origin	Malacacheta	Myanmar	Madagascar	Sri Lanka	Montana (Blue)
Elements	range	mean			
Be	0.2-3.1*	1.2*	-	-	-
Mg	41.9-138.4	81.5	25	63	37
Ti	127.2-608.5	320.2	59	164	50
V	8.7-30.5	20.3	1.5	23	5.2
Fe	1,845-6,393	3,722	2,040	924	4,598
Ga	53.7-124.0	83.0	89	63	54
Source	This study	Kreb, 2020			Zwaan, 2016

(*except high points)

Table 1. Trace elements of sapphire from Malacacheta and other origins (in ppmw)

FT-IR spectroscopy revealed almost no significant absorptions, except for weak kaolinite peaks in one sample. No OH-related absorptions around 3309 cm⁻¹ were detected, not even as an almost insignificant feature (Figure 6). UV-Visible spectroscopy indicated characteristics typical of non-magmatic sapphires, with no broad-band absorption near 880 nm and a UV cut-off between 320 and 330 nm (Figure 7). These features suggest a metamorphic origin and the absence of heat treatment under reducing conditions.

LA-ICP-MS analysis detected Be in all samples, with concentrations ranging from 0.2 to 3.1 ppmw across most areas, and up to 6.8 ppm in specific regions containing cloud-like inclusions. High field strength elements such as Sn, Nb, and Ta were also detected in proportion to Be. Compared to the average trace element concentrations reported by Kreb *et al.* (2020) and Zwaan *et al.* (2016), the Malacacheta sapphires exhibited higher levels of Fe, Ti, and Mg than

meta-morphic sapphires from Myanmar, Madagascar, and Sri Lanka. In particular, the Fe concentrations were similar to those found in Montana sapphires.

The Malacacheta sapphires are mined from alluvial deposit together with alexandrite. As the host rock for alexandrite formation in Malacacheta remains unidentified (Basílio *et al.*, 2013), the exact origin of the sapphire is also uncertain. However, Basílio (2013) proposed that the alexandrite in this region formed via a Cambrian-aged metasomatic system, in which beryllium-rich fluids from nearby granite intrusions interacted with meta-ultramafic rocks and peraluminous schists. The features observed in this study—including strong strain patterns, CO₂ rich fluid inclusions and plagioclase inclusions, the absence of OH absorption in FT-IR spectra, and elevated Be, Fe, Mg, Ti concentrations—support the hypothesis that these sapphires might have crystallized in peraluminous schists and undergone metasomatic alteration by Be-rich fluids derived from granite intrusions.

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Gemological Characteristics of New Transparent Brown Sapphire

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A newly observed variety of faceted transparent brown sapphire has recently entered the gem market, prompting gemological attention due to its highly unusual appearance (Figure 1). Unlike traditional brown sapphires—commonly referred to as geuda in Sri Lanka or brown sapphires from basaltic-related origins—which are generally opaque or only weakly translucent due to dense microscopic inclusions and iron-related chromophores (Li and Shaokui, 2022; Wongkokua *et al.*, 2023; Zheng *et al.*, 2013; Zwaan, 1998). These new samples exhibit high transparency and attractive brown coloration. Such features suggest a treated origin rather than a naturally occurring chromophore system. This study investigates the gemological, spectroscopic, and chemical properties of these stones, with a focus on identifying the origin of color, the presence of any treatments, and the probable geological source.



Figure 1 Rough and faceted transparent brown sapphires. The pear-shaped stone in the middle weighed 2.29 ct. Photo: T. Sripoonjan

Gemological testing confirmed that the stones are corundum, with refractive indices between 1.760 and 1.770 and specific gravities close to 3.98–4.00. Spectroscopic analysis revealed a gradual absorption rise beginning near 550 nm and extending into the UV region (Figure 2), attributed to charge transfer between trapped holes (h^+) and Fe^{3+} , as well as a small distinct absorption at 388 nm caused by isolated Fe^{3+} ions (Dubinsky *et al.*, 2020). Notably, absorptions typically associated with Fe^{3+} – Fe^{3+} pairs, particularly at 377 and 450 nm, were absent, indicating that iron does not serve as the dominant chromophore in these samples. Microscopic observation revealed minute particulate inclusions and hazy zones causing Tyndall scattering. In some specimens, discoid fissures, glassy internal features, and dotted whitish minute particles consistent with rutile breakdown were observed—

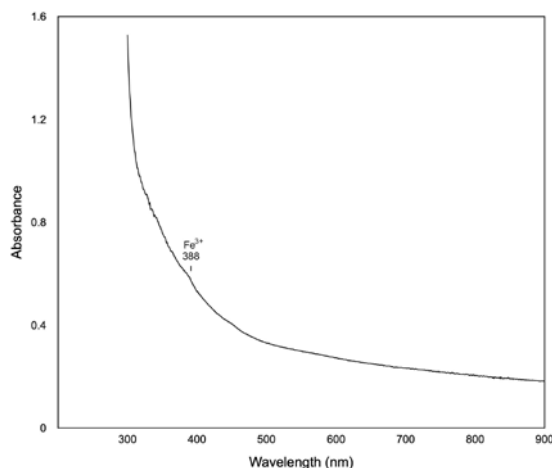


Figure 2 Representative UV-Vis-NIR spectrum of brown sapphire from this study

features indicative of exposure to extreme temperatures and supporting evidence of high-temperature heat treatment.

Laser ablation ICP-MS analysis detected beryllium concentrations ranging from 5 to 62 ppma, strongly suggesting that these sapphires have undergone heat treatment involving the diffusion of beryllium. Iron concentrations were low and closely aligned with those typical of metamorphic sapphires such as those from Sri Lanka, as opposed to those derived from basaltic sources. FTIR spectroscopy showed no hydrous or hydroxyl-related absorption bands, consistent with oxidative heating at elevated temperatures. DiamondView imaging revealed chalky blue fluorescence localized in the brown zones, attributed to titanate group-related structural features (Vigier *et al.*, 2023). This

fluorescence behavior, combined with the low iron content, reinforces the hypothesis that the brown coloration is primarily due to submicroscopic titanate-related particles and associated defect centers, rather than conventional Fe-based chromophores.

These findings reveal a previously undocumented treatment-induced appearance in sapphires and highlight the importance of advanced analytical methods in establishing origin and treatment history. This study underscores the need for continued research; particularly experimental work aimed at reproducing this brown color through controlled heating processes—a technically complex challenge that may yield deeper insights into color development mechanisms in corundum.

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Acknowledgements

The authors gratefully acknowledge the Gem and Jewelry Institute of Thailand (GIT) for their valuable assistance in providing advanced analytical facilities for this study. Special thanks are extended to Mr. Pukkapon Piantumdee of Sunset Gems Co., Ltd. for generously supplying the sapphire samples that were critical to the completion of this research.

HFSE-enriched sapphires of gem quality: A combined FTIR and trace element study and implications for heat treatment detection

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Gem-quality sapphires containing ‘exotic’ elements such as Sn, Nb, Zr, Ta, W, are of special interest in gemmological research, as the concentration of these so-called high-field-strength-elements (HFSE) often correlate with detectable traces of naturally incorporated beryllium (Whatanakul *et al.* 2004, Pardieu 2007, Shen 2007, Zhou *et al.* 2022 & 2024). It is this correlation, which enables gemmologists in most cases to safely separate them from Be-diffusion treated sapphires which show no such correlation when analysed with LAICPMS. In addition, some of these sapphires may also contain thorium and lead traces, which enable us to carry out direct radiometric dating on such sapphires (Wälle & Wang 2023).

Interestingly, such ‘exotic’ trace elements are found in both, basaltic sapphires (e.g. Nigeria, Tasmania, N-Madagascar) and metamorphic sapphires (e.g. Madagascar, Sri Lanka, Afghanistan) and are commonly related to metasomatic

processes during crystal growth. How these incompatible elements and beryllium were incorporated into sapphires has been investigated in several studies, with models explaining the presence of these elements either by primary integration of nano-inclusions during growth of the sapphire or by direct incorporation of these elements into the corundum structure with a possible epigenetic segregation into nanoclusters or precipitates as nanoparticles of separate phases (Shen & Wirth 2012, Emori *et al.* 2019; Oto *et al.* 2023, Jin *et al.* 2024 and references therein).

This study focuses mainly on FTIR spectroscopy of HFSE-enriched sapphires of metamorphic origin from Madagascar and Sri Lanka. We analysed 18 untreated but gem-quality sapphires ranging in weight from 0.45 ct to 50 ct, partly from the SSEF research collection and from reliable trade members (Figure 1). All samples contained either goethite (e.g. in deep hollow tubes) or diaspore in fluid inclusions,



Figure 1: Selection of investigated sapphires (5.7 ct to 50 ct) from Madagascar and Sri Lanka.

thus confirming them to be unheated samples (Sripoojan *et al.* 2016; Krzemnicki *et al.* 2023). Interestingly, most of the investigated samples were of very fine quality with a velvety blue colour generally related to a certain amount of turbidity and occasionally dense growth structures and thin laces of very dark ink-blue colour.

Most of these HFSE-sapphires showed a broad absorption band at about 3300 cm⁻¹, partly accompanied by peaks at 3182, 3232, and 3308 cm⁻¹ related to OH- groups associated with Al vacancies (Moon & Phillips 1991; Beran & Rossman 2006; Jollands *et al.* 2023 and references therein). By comparing their trace element content, we found best correlation with the tantalum concentration in these samples, although contribution by other HFSE-elements including Sn cannot be fully excluded. In general, HFSE-bearing sapphires with a low Ta concentration exhibited none to a small band, whereas stones with higher Ta were characterised by an increasingly strong band at 3300 cm⁻¹. As a special case, we found in sample 032 a similar but slightly shifted broad band centred at about 3385 cm⁻¹ with two small side peaks at 3254 and 3310 cm⁻¹. Although high in tantalum (Ta_{mean} 67 ppm), the reason for this 'shift' and the side peak at 3254 cm⁻¹ is not yet fully understood and needs further research.

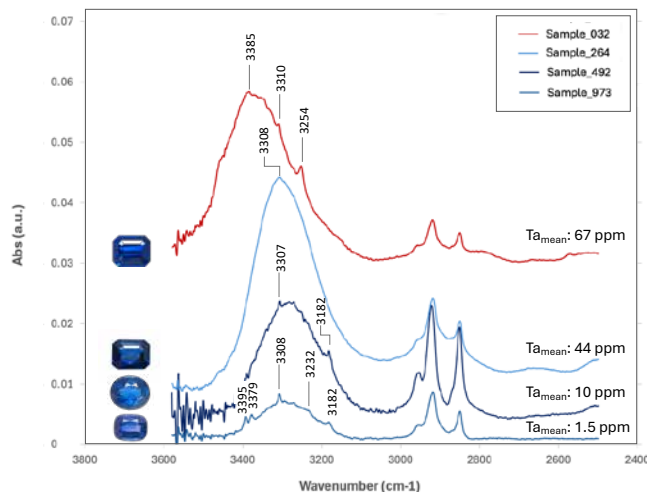


Figure 2: FTIR spectra of selected HFSE-sapphires characterised by a broad band centred at 3300 or 3385 cm⁻¹. Sample 973 generally has a mean Ta concentration of 1.5 ppm, but contains a micro-zonation enriched up to 345 ppm Ta, which may have a certain (small) influence on the observed FTIR spectrum.



Figure 3: Thin Ta-rich (and HFSE-rich) zone in HFSE-sapphire sample 973 seen under microscope as colourless band and as chalky white zone in deep UV (DiamondViewTM).

Conclusions:

This study on unheated sapphires enriched in HFSE-elements reveals the presence of a broad absorption band at about 3300 cm⁻¹, seemingly best correlated with the Ta-concentration in these sapphires. The 'shift' of this band to 3385 cm⁻¹ observed in one sample (Sample 032) is yet not fully understood, although assumed to be also related to some extent to tantalum.

Notably, some of these unheated sapphires show small OH- related peaks (3182, 3232, and 3308 cm⁻¹). In addition, these unheated sapphires also exhibit varying intensities of chalky white reactions. Although both features

(OH- peaks in FTIR and chalky reaction in SWUV) are well known and described in heat treated metamorphic sapphires, this study reveals that such features may also be encountered in unheated stones. As such this study further underlines, that a simplistic approach for no-heat/heated distinction can be erroneous specifically for metamorphic sapphires with velvety appearance and turbidity. Only a full characterisation of these sapphires including Raman spectroscopy on inclusions (diaspore, goethite, zircon) and detailed trace element analyses (LA-ICPMS) may provide the answer whether such a sapphire had been heated or not.

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Low Temperature Heat Treatment of Madagascar Sapphire

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One of the challenges facing gemologists today is the detection of low temperature (below 1200°C) heat treatment in corundum (Hughes *et al.*, 2017). At these temperatures, the color of ruby and sapphire can be altered, but some common features such as rutile silk may remain intact.

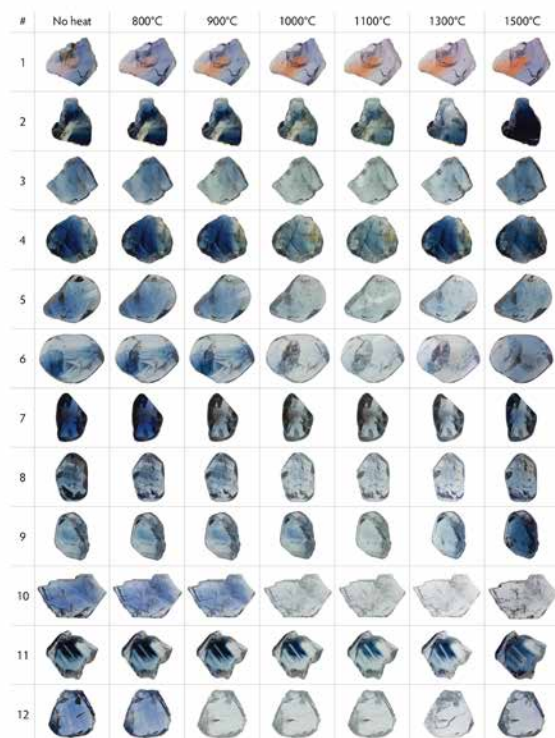


Figure 1. Madagascar sapphire before and after heating the 12 samples, shown before and after heating to each temperature. Most of the samples began to lighten significantly after heating to about 900–1000°C and started to deepen in color again around 1300°C. After heating to 1500°C, many pieces became significantly deeper in color. Photos not to scale. Photos by Rosey Perkins and Sora-at Manorotkul.

To aid in detection, the authors heated Madagascar sapphires at a range of temperatures from 800°C to 1500°C. The samples were documented using macrophotography in the untreated state and after heat treatment at elevated temperatures (figure 1), photomicrography of inclusions (figure 2), ultraviolet fluorescence imaging (figure 3), and spectroscopic analysis (ultraviolet/visible/near infrared and infrared) to record any changes, with a focus on features that could help detect heat treatment.

Results showed that a combination of inclusion studies, infrared spectroscopy, and observation of fluorescence could help detect treatment. We also noticed a fascinating change in UV-Vis-NIR spectra that can have implications for origin determination of blue sapphire (figure 4) (Schmetzer and Bank, 1980 and Emmett and Douthit, 1993). Many of the samples developed a peak at 880 nm after heat treatment. This can make it challenging to separate heated metamorphic sapphires from Madagascar from the basalt related material.

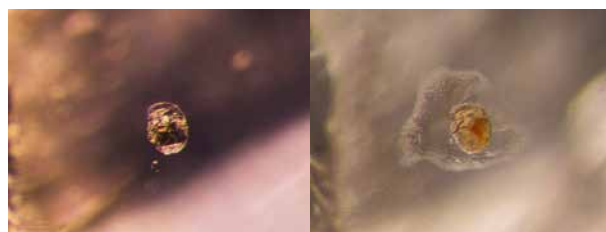


Figure 2.

Left: A small, flat crystal is pictured in sample 10 before heat treatment.

Right: After the first heating to 800°C, a reflective fissure begins to appear around the crystal. By 1100°C (pictured), the fissure has begun to heal at the edges.

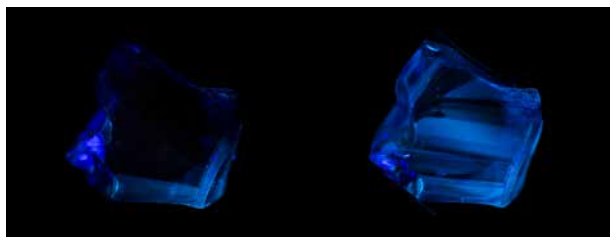


Figure 3. Short-wave UV fluorescence reaction of a Madagascar sapphire during heating

Left: Sample 11 began to show a short-wave fluorescence reaction after heating to 1000°C. This was mainly limited to one side of the stone, where a chalky blue band can be seen.

Right: After heating to 1100°C, the chalky blue reaction became stronger and more widespread, with chalky blue bands across the stone.

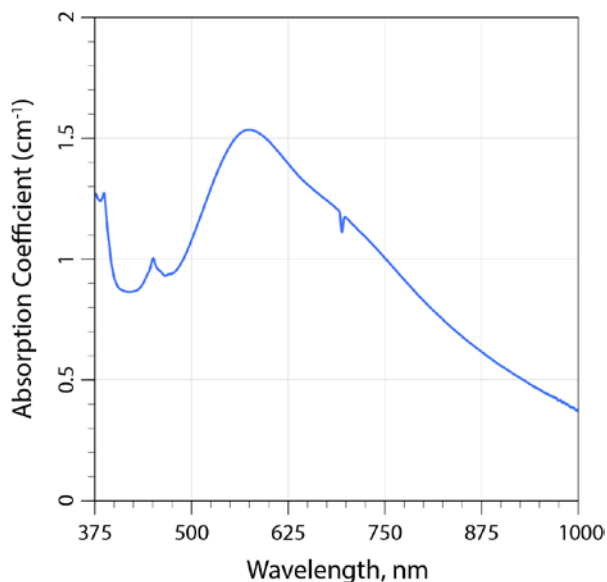
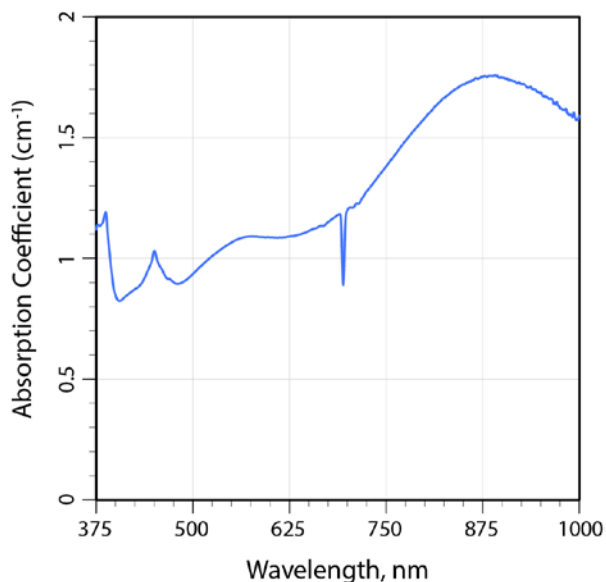


Figure 4. UV-Vis-NIR spectrum of a Madagascar sapphire before and after heating

Left: The UV-Vis-NIR spectrum of sample 7 before heat treatment. Note that the most prominent feature is the absorbance between 500 and 600 nm, as well as a small Fe-related peak around 450 nm.



Right: The UV-Vis-NIR spectrum of sample 7 after heating to 900°C. Note the development of a large peak at ~880 nm.

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Effects of Gamma Irradiation on Corundum: Towards a Potential Detection Method

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Introduction

Corundum, encompassing the highly prized ruby and sapphire varieties, stands as a cornerstone of the global gemstone market. The allure and value of these gems have historically driven the development of various treatment techniques aimed at enhancing their colour and clarity. While many treatments are stable and widely accepted when properly disclosed, the emergence of new or less understood methods presents an ongoing challenge for the gemological community. Accurate detection and disclosure of all treatments are paramount for maintaining market confidence and ensuring fair trade. Gamma irradiation has been explored as a potential method to alter the colour of various gemstones, including corundum (Ashbaugh, 1988; Pardieu *et al.*, 2022). However, the effects of such irradiation on corundum, the stability of any induced colour changes, and robust methods for their detection remain areas requiring comprehensive investigation. This contribution builds upon our preliminary work, presented at the IGC Tokyo 2023 (Wang *et al.*, 2023), to provide a more in-depth understanding of gamma irradiation effects on a diverse suite of corundum samples and to outline a potential detection method.

Materials and methods

• **Sample selection:** For this study, we carefully selected 29 natural and synthetic corundum samples. This selection was designed to represent a variety of colours, including colourless, pink to purple fancy sapphires, rubies and sapphires. The rationale was to assess how differing trace element compositions, which are fundamental to corundum's colour, influence the response to gamma irradiation. Four selected samples representing different initial colours and elemental concentrations (Mg, Ti, V, Cr, Fe) are presented in Table 1. Elemental concentrations were determined by LA-ICP-TOF-MS.

		Init. Colour	Mg	Ti	V	Cr	Fe
Sample 1	Nat. Crd.	blue	100	175	11	10	2800
Sample 2	Nat. Crd.	purple	25	50	10	400	540
Sample 3	Syn. Crd.	purple	-	60	0.6	360	95
Sample 4	Syn. Crd.	purplish red	-	5	1	1700	-

Table 1. Information and initial colours of the selected four corundums as well as their trace element composition. All numbers are in unit of ppm. “-” indicates that the concentrations are below detection limit ~1ppm.

- **Gamma irradiation:** The samples were subjected to gamma irradiation at FRM II Facility for a continuous period of 70 hours, achieving a total dose of approximately 33 kGy (Li, *et al.* 2022). This represents a substantially higher dose than in our preliminary investigations, intended to maximize any potential colour alterations.
- **Post-irradiation handling:** Immediately following irradiation, samples were carefully transferred into an aluminum container. This precaution was taken to shield them from ambient light exposure, thereby preserving any unstable colour centers that might have been restored until systematic laboratory analysis could start. Upon return to the laboratory, the initial post-irradiation colour of each sample was documented.
- **Colour stability testing:** To test the colour stability we followed an altered procedure based on the standard testing procedure as used also for client stones in the SSEF laboratory (Krzemnicki, 2022). The colour shift of 4 selected samples after each step are shown in Figure 1.

o **Fading test:** To assess the light stability of any induced colour, samples were exposed to a 20 W high-power LED light source, for a duration of 3 hours. Importantly, this LED source was characterized by the absence of long-wave UV (LWUV) component at 365 nm and only a negligible contribution of short-wave UV (SWUV) at 254 nm (Palke *et al.*, 2023).

o **Activation test:** Following the fading test, samples were exposed to standard gemological UV lamps to probe for the presence of colour centers that could be reactivated. This primarily involved exposure to LWUV (365 nm) for 10 minutes. For specific samples that showed no discernible colour shift after LWUV activation (e.g., Sample 3), more energetic activation methods were employed, including LWUV exposure for 2 hours, SWUV exposure for 1 hour

and deep UV exposure using a DiamondView instrument for 10 minutes, to explore the potential activation of other types of colour centers.

o **UV-VIS-NIR spectroscopic analysis:** UV-VIS-NIR absorption spectroscopy was the primary analytical technique used to characterize colour and to identify specific chromophores or colour centers. Spectra were recorded over a range of 290 nm to 1600 nm. While the majority of analyses focused on polarized light through C-axis (O-ray), E-ray measurements were also conducted on select samples to further investigate anisotropic absorption features.

o **Mild heat treatment:** After the colour stability test, a mild heat treatment at 350 °C for 20 minutes was applied to all samples, in order to see the colour stability under heat condition.

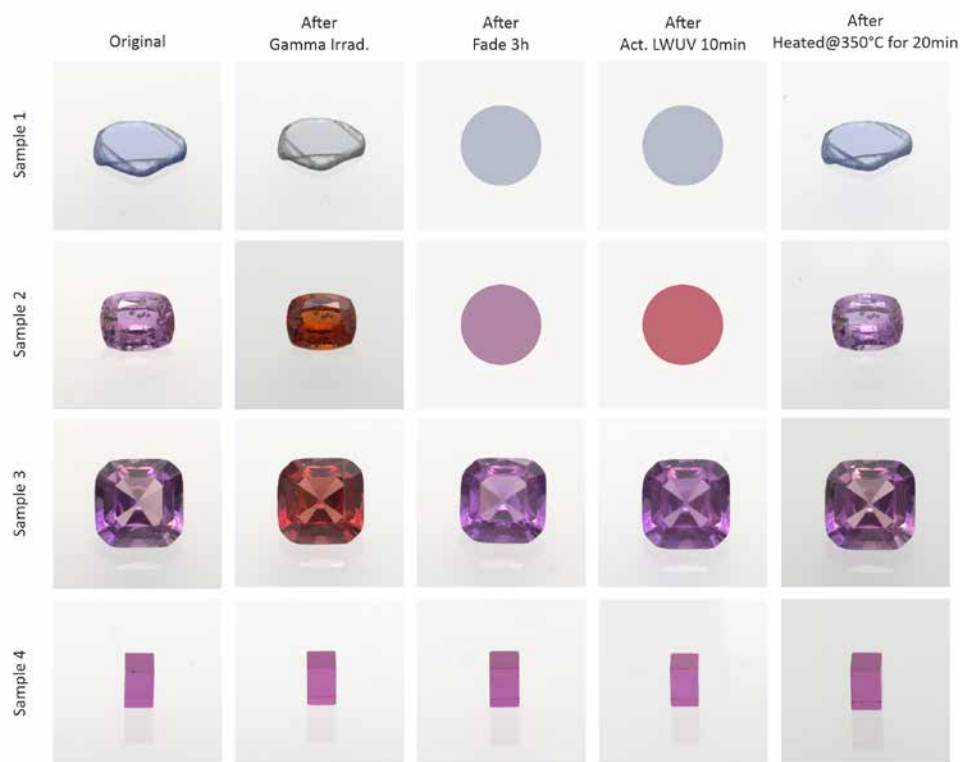


Figure 1. Photos of selected four corundums undergoing colour shift (except for sample 4 no obvious colour shift) after Gamma irradiation, colour stability test in the lab and a mild heat treatment. After fade and after activation photos of Sample 1 & 2 were unfortunately not properly taken, hence the colours are represented by the colour disks.

Results and conclusion

Gamma irradiation induces diverse, chemistry-dependent colour shifts in corundum, which we have categorized into four distinct groups based on their behaviours (Figure 1). These responses include irreversible (during fading or UV activation) blue colour reduction in Fe-Ti rich sapphires (Group 1), which might be one of the reasons that blue/purple hue can be reduced in some rubies. We also observed the development of unstable, light-sensitive, colour reversible orange hues in some natural pink/purple corundum (Group 2) and the appearance of semi-stable orange hues in certain varieties (Group 3) that are bleachable by intense light but not readily reactivated by standard UV sources, nor higher energy UV lights. Notably, high concentration Cr-only synthetic corundum (Group 4) showed an inert response to the irradiation. Those samples with colour shift after irradiation can be restored to their original colours after a mild heat treatment.

The variability and potential transience of these induced colour shifts require a detection approach beyond simple observation. We will propose a strategy based on differential UV-VIS-NIR spectroscopy, meticulously tracking spectral

changes during colour stability tests (controlled fading and UV activation). This method involves comparing spectra taken before, during, and after these tests to identify characteristic patterns of colour centres which are unstable, or which can be activated again. We will also discuss spectral features serving as illustrative examples of ongoing investigations into diagnostic markers. This emphasis on the dynamic response of the stone offers greater diagnostic potential than static measurements.

In conclusion, while gamma irradiation effects in corundum are complex and present detection challenges, our methodology may hold promising potential. Future research will focus on expanding the sample database, refining stability testing protocols to enhance diagnostic spectral features, correlating these features with specific colour centres and defects, and validating the method through broader studies. This work aims to provide the gemmological community with more robust tools for identifying gamma-irradiated corundum, thereby supporting market transparency and consumer confidence.

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Challenges in the Detection of Ruby and Sapphire Treatment in the Thai Market

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Keywords: ruby, sapphire, corundum, treatment detection, heat treatment, FTIR, Raman spectroscopy, diffusion, glass filling, gemology

Abstract

The detection of treatments in ruby and sapphire remains a critical aspect and has become an increasingly complex issue in the gem trade and laboratories alike. Treatments such as traditional heating, flux-assisted heating, various types of glass filling, lattice diffusion heating using elements from an external source, irradiation, and pressure-assisted heating have been widely applied to enhance colour, clarity, and durability of corundum, directly affecting the stone market values and ethical disclosure. The advancement of these techniques, particularly those designed to evade detection, have outpaced many traditional gem detection methods, necessitating the integration of advanced analytical technologies.

Introduction

The commercial value of ruby and sapphire is closely linked to their natural, untreated or treated state. The treatments, usually common, must be accurately detected and disclosed to ensure transparency and maintain consumer trust. However, as treatment technologies evolve, so do the challenges in their detection. This presentation is aimed at reviewing and updating both traditional and advanced methods for identifying treatments in the ruby and sapphire and discusses the identification challenges, ongoing research and future directions in this field.

Overview of Treatment Methods

Corundum treatments are commonly employed to enhance a gem's appearance, and they can be broadly categorized into heat-involved and non-heat-involved methods (Pisutha-Arnond, 2017). Heat-involved treatments include conventional heating (typically around 1200 – 1700 °C), flux-assisted heating using substances like borax and/or sil-

ica to (partially) assist healing of fissures. In addition, heat treatment combined with diffusion is used, such as shallow surface-diffusion with added colouring agents such as Ti alone or Ti with Be (as well as Ti and Fe in case of synthetic colourless sapphire) to create blue colouration (e.g., Kane *et al.*, 1990, Leelawatanasuk *et al.*, 2014, Pisutha-Arnond *et al.*, 2019), Cr to produce red hue (McClure *et al.*, 1993), and high-temperature beryllium diffusion (above 1700°C) often resulting in yellow colour (e.g., Emmett *et al.*, 2003, Pisutha-Arnond *et al.*, 2004). Other heat treatments of corundum include fracture filling with high-refractive-index glasses (e.g., lead or bismuth-based, colourless or coloured) at rather moderate temperatures and heating under pressure (e.g., Krzemnicki *et al.*, 2019), as well as other low temperature heat treatments. In contrast, non-heat-involved treatments of corundum consist of oil or resin filling to improve clarity, dyeing to enhance or alter colour, and irradiation to induce a change of stone colour (e.g., introducing an unstable yellow hue). Each treatment may produce specific diagnostic features. However, certain corundum treatments - particularly low-temperature heating and recent glass filling – often are challenging for gemologists to accurately identify and properly disclose the treatment of a stone.

Basic and Advanced Techniques for the Detection of Ruby & Sapphire Treatments

- Microscopy: Identification of specific diagnostic features, e.g., altered inclusions, newly healed fissures, presence of flux residue, colour concentration along stone surface and fractures.
- FTIR Spectroscopy: May assist heat treatment detection of corundum, e.g. based on characteristic absorption bands

(e.g., 3232, 3309, 3185 cm⁻¹), though not always conclusive for all samples (Soares *et al.*, 2025, Phlayrahan and Homkhajorn, 2021)

- Raman Spectroscopy: Detects phase transformation of some inclusions (e.g., goethite to hematite). The presence of goethite (or diasporite) in corundum is a strong indication that a corundum sample has not been heated (Krzemnicki *et al.*, 2023)
- EDXRF and LA-ICP-MS: Identify trace elements and diffusion of foreign elements (e.g., Be, Pb, Ba, Bi) associated with specific treatments (Krzemnicki *et al.*, 2004)
- LIBS: Elemental analysis for diffusion treatments, especially Be-diffusion (Krzemnicki *et al.*, 2004)
- Luminescence Analysis: Observes changes in silk inclusions and UV-induced luminescence as indicators of heating, especially in Geuda sapphires (Pluthametwisute *et al.*, 2025)

Key Challenges in Identification

- Low-temperature heat treatment of corundum is often difficult to detect, as it may not altered inclusions significantly or produce clear spectroscopic markers (e.g., Atichat *et al.*, 2011, Hughes *et al.*, 2022)
- Heat treatment detection in corundum from basaltic origins is particularly difficult, as natural heating during ascent in alkali basalt and high iron content—which suppresses diagnostic spectral features—complicate differentiation from heated stones. Accurate trace element

analysis and close inclusion observation are essential. (e.g., Soonthorntantikul *et al.*, 2019).

- Some glass-filled stones may appear highly transparent, masking fractures quite effectively; detection may be supported by chemical analysis (e.g., lead content) and X-ray imaging (see figure 1).
- Surface diffusion treatments create color only on the outer rim, requiring immersion techniques and careful chemical profiling for identification (see figure 2).
- Irradiation treatments can create an unstable (in some cases also stable) yellow colouration, but it could also induce a subtle change of colour that is difficult to distinguish from natural colouration. Currently, only the colour stability of such stones can be checked by a fading test, whereas the irradiation treatment itself cannot be detected up to this day. Recently, advanced spectroscopic and fluorescence techniques are under investigation allowing possibly a reliable detection in the near future (e.g., Wang *et al.*, 2023).
- Detection the presence of oil in ruby and sapphire: especially colourless or low-viscosity oil could be a challenge without the backing of other techniques, such as Raman microprobe, FTIR spectroscopy, and hot point tester.
- Overlapping spectroscopic features in treated and untreated stones complicate detection, but the most reliable results come from combining multiple analytical techniques (Soares *et al.*, 2025, Phlayrahan and Homkhajorn, 2021).

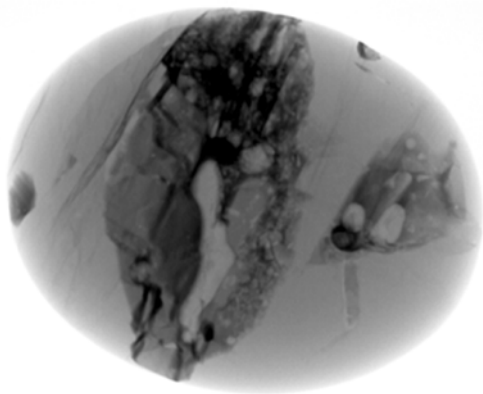


Figure 1. X-ray image of a glass-filled (and heated) ruby showing high density glass along the fractures within the stone. Photo by W. Chongraktrakul; image widths 7.0 mm.

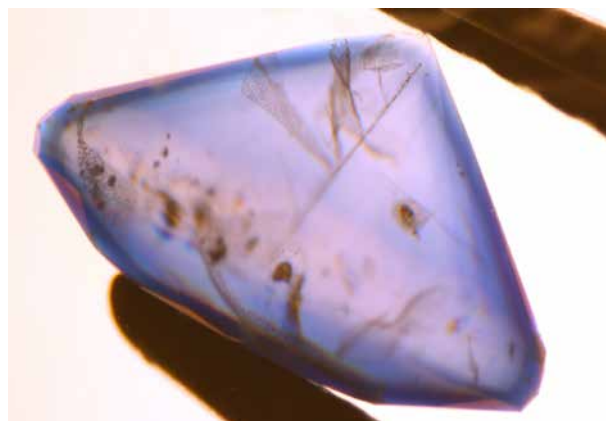


Figure 2. A diffusion-treated sapphire exhibits blue colour concentration along the stone surface, while the interior is noticeably lighter or even colorless. Photomicrograph by S. Promwongnan; image widths 7.0 mm.

Conclusion

The detection of treatments in ruby and sapphire is an ongoing challenge due to the sophistication of modern enhancement techniques. Accurate identification requires a combination of traditional gemological skills and advanced analytical technologies. Continued research, method devel-

opment, and collaboration among laboratories are essential to keep pace with evolving treatment methods. Above all, transparency and full disclosure of treatments are essential to uphold ethical standards and maintain consumer confidence in the gemstone industry.

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Padparadscha Sapphire & heat treatment

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Keywords: Padparadscha sapphire, heat treatment, Laboratory Manual Harmonisation Committee

In 2018 the Padparadscha engagement ring of Princess Eugenie (Figure 1), the niece of the king, gave a boost to the popularity of the gemstone in the UK. Its subtle colour makes it ideal for the independently minded looking for something different (Figure 2).



Figure 1 –Princess Eugenie's engagement ring.
Photo: Hello Magazine.



Figure 2 – Padparadscha Sapphire. Photo: SSEF.

In gemmological circles the Padparadscha sapphire moved centre-stage with the discovery of beryllium diffusion in 2002 - treatment may have been occurring the year before (Emmett, J.L., et al 2003). There is much on the accepted colour of padparadscha and treatments (Notari, Franck, 1996). The present Laboratory Harmonisation Committee Information Sheet 4 (Version 10, February 2023) standardises the nomenclature they use to describe a 'padparadscha sapphire' (www.lmhc-gemmology.org) I wish to

thank the individuals involved, who have provided these Information Sheets on many gemstones over the years. They are an excellent tool to explain the issues of report wording to both trade and public alike. In general I inform customers that I follow the World Jewellery Confederation (CIBJO) and LMHC guidelines. However I do not issue reports on heated 'padparadscha' sapphires', which is still acceptable according to LMHC wording. It is this I would like us to consider.

Many gem dealers in the UK have informed me that they do not trade in heated padparadscha sapphires. It may be due to the lack of availability of heated (not diffused) material, or maybe those seeking Padparadschas for commissioned jewellery are only interested in unheated gemstones. It may be a pragmatic decision in that the gem dealer or trader, in the UK or from abroad, may be asked to provide a laboratory gem report to prove it is only heated and not diffusion treated. The LA-ICP-MS spectroscopy to check for diffusion is not widely available and testing them might be expensive. So it is likely they will be traded as untested heated padparadschas. This temptation is removed if any heated pink/orange sapphire, along with diffused pink/orange sapphire, is not allowed to have the padparadscha label.

I understand the quandary of the LMHC. If padparadscha is a variety name then it has to follow the allowances for

all variety names in that they can be heated or unheated. I would prefer the variety name to be pink/orange sapphire and the commercial name of padparadscha only applied to unheated gemstones. However the variety name of padparadscha is so well established in the trade it probably cannot be changed. Even if we keep the variety name as padparadscha a precedent has already been set that padparadscha cannot be applied to diffusion-treated pink/orange sapphire and it is therefore only a small step further to say it cannot be applied to heated pink/orange sapphires. I know padparadscha is arguably only a colour description in Sinhalese but the term has taken on a meaning in its own right of a premium gemstone with a premium price. We have 'out-lawed' padparadscha being applied to diffused orange/pink sapphires so let's take it a small step further and remove the premium padparadscha label being applied to heated orange/pink sapphires.

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Advancing Fei Cui Origin Authentication: A comparison of green Fei Cui from Myanmar, Guatemala, Italy and Russia

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Keywords Origin determination, Fei Cui, Jadeite jade, Omphacite jade

In recent years, an increasing number of sources of fei cui, such as Guatemala, Italy and Russia, have entered the Chinese market, leading to a growing demand for precise origin determination of fei cui. Building upon our previous study (Liu *et al.*, 2024), this study progresses towards a more comprehensive understanding. In November 2024, author Liu conducted a geological field trip to the fei cui mines in Guatemala (Figure 1), collecting a new set of Guatemalan samples directly from the source. To expand the study's scope, green samples from Russia were also included (Figure 2).

Following the standard tests outlined in the fei cui standards (GAHK, 2016; GIT, 2022), which encompass RI, SG, and FTIR analyses, all samples (BU, n = 21; GU, n = 29; RU, n = 13; IT, n = 7) have been confirmed as fei cui, devoid of any resin or dye. Moreover, the FTIR results, consistent with

the literature (Abduriyim *et al.*, 2017; Miura *et al.*, 2019), support the classification of the Russian samples as jadeite jade. In alignment with our previous study, the recent batch of Guatemalan samples collected between 2023 and 2025 also exhibits jadeite dominance, while the Italian samples are characterised by chrome-omphacite dominance (Liu *et al.*, 2024). Compared with other localities, Russian fei cui relatively coarse granular and loose texture, and contains phlogopite and molybdenite (Figure 3).



Figure 2. The samples of green fei cui from Russia (left, RU-01 1.03 ct), Guatemala (centre, GU-G31 0.83 ct), and Myanmar (right, BU-41 0.78 ct) analysed in this study



Figure 1. Outcrop of jadeitite and omphacitite (white area on the left) with intrusive dykes of albitite and amphibolite in serpentinite mélangé in a new mining site on the north - eastern side of the Motagua fault zone in Guatemala. Photo © S.I Liu

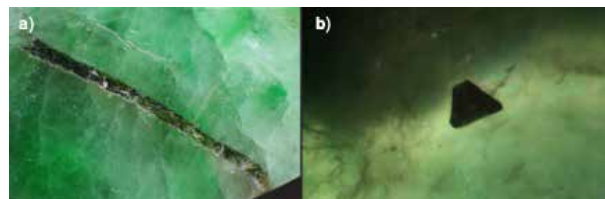


Figure 3. (a) Phlogopite and (b) molybdenite inclusions found in Russian fei cui

LA-ICP-MS were used to analyse the characteristic trace element profile of fei cui from different origins. The 3D-PCA score plot of LA-ICP-MS (Figure 4a) indicates a distinct separation of Italy from other regions, with slight overlap

observed among Myanmar, Guatemala, and Russia. Among the four localities, Russian fei cui has the highest Ga, Zn, Li, and Fe content, while Italy has the highest Cr content (Figure 4b) due to its chrome-omphacite-dominated nature.

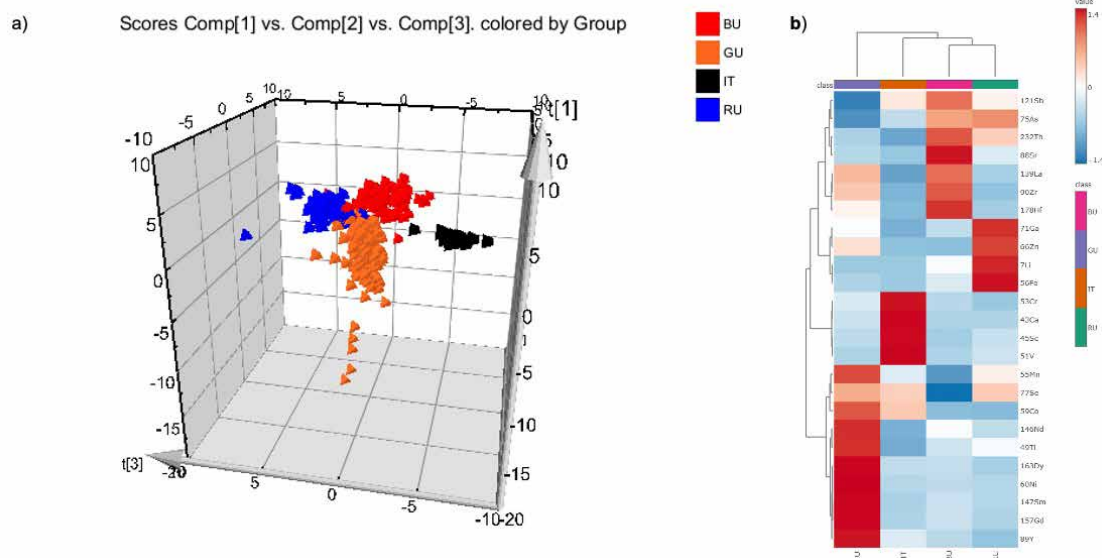


Figure 4. (a) 3D-PCA score plots of LA-ICP-MS data of green fei cui from Myanmar (BU), Guatemala (GU), Italy (IT) and Russia (RU); (b) Heatmap of LA-ICP-MS data.

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Color Genesis Analysis and Characteristics for Origin Identification of Purple Jadeite

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Introduction

Purple jadeite, renowned for its distinctive purple hues and rich cultural connotations, is highly admired and sought after. This study investigates purple jadeite samples from Burma and Guatemala, each exhibiting different color variations, as shown in Figure 1. The research commences with a conventional gemological analysis, followed by a comprehensive examination of the color genesis through various spectroscopic techniques, including X-ray fluorescence (XRF) spectra analysis, ultraviolet-visible (UV-Vis) absorption spectra analysis, infrared spectroscopy (IR), Raman spectra analysis and Photoluminescence Spectroscopy (PL) analysis.



Figure 1: Purple Jadeite Samples from Burma and Guatemala.

Results

The results reveal that the coloration of purple jadeite is primarily attributed to the presence of manganese ions (Mn), iron ions (Fe), and titanium ions (Ti). The intensity of the purple coloration is directly correlated with the concentration of manganese ions, while iron ions and titanium ions producing a more violet color. Notably, there are significant differences in trace element concentrations between the two sources: Burmese purple jadeite exhibits higher manganese and lower titanium content (MnO: 0.038-0.065%, TiO₂: 0.015-0.200%), whereas Guatemalan purple jadeite shows lower manganese and higher titanium content (MnO: 0.011-0.032%, TiO₂: 0.506-0.697%). Furthermore, the jadeite samples examined in this research all exhibit purple hues, primarily attributed to the chromophoric contributions of manganese (Mn), iron (Fe), and titanium (Ti) ions, with no significant relationship to chromium (Cr) ions. The chromium content is relatively low, as seen in Table 1.

Sample	Fe ₂ O ₃ %	MnO%	Cr ₂ O ₃ %	TiO ₂ %	NiO%
B-001	0.098	0.044	0.012	0.024	0.052
B-002	0.073	0.040	0.004	0.015	0.043
B-003	0.115	0.038	0.001	0.029	0.045
B-004	0.246	0.041	0.027	0.079	0.043
B-005	0.087	0.034	0.017	0.054	0.044
B-006	1.291	0.055	0.013	0.200	0.056
B-007	0.093	0.044	0.005	0.025	0.040
G-001	0.210	0.013	0.001	0.643	0.032
G-002	0.193	0.011	0.009	0.557	0.027
G-003	1.063	0.030	0.014	0.506	0.021
G-004	0.133	0.032	0.011	0.697	0.025
G-005	0.188	0.027	0.009	0.559	0.037

Table 1: Trace element content in Burmese and Guatemalan Purple Jadeite.

The ultraviolet-visible absorption spectra of purple jadeite from the two regions exhibit distinct differences. Burmese purple jadeite is primarily characterized by absorptions at 314, 437, and 581 nm, as illustrated in Figure 2, whereas Guatemalan purple jadeite displays characteristic absorptions at 309, 380, 437, 551, 626, 768, and 947 nm, as shown in Figure 3. The absorptions at 381 and 437 nm are attributed to d-d electronic transitions of Fe^{3+} , while the absorption at 581 nm is associated with d-d electronic transitions of Mn^{3+} . The absorption at 626 nm is due to electronic transitions of Ti^{3+} , the absorption at 768 nm results from charge transfer between Fe^{2+} and Fe^{3+} , and the absorption at 947 nm is attributed to d-d electron transitions of Fe^{2+} [which reference?].

Therefore, 581 nm represents a characteristic peak for Burmese purple jadeite (Minghui *et al.*, 2023), while 551 nm is a characteristic peak for Guatemalan purple jadeite. The 581 nm feature appears as a peak in the UV-Vis spectra of Burmese purple jadeite, whereas it manifests as a trough between 551 nm to 626 nm in the spectra of Guatemalan purple jadeite. The UV-Vis spectra of Burmese purple jadeite in the 580–950 nm range exhibit a single peak, whereas the spectra of Guatemalan purple jadeite within the same range display multiple peaks.

In terms of infrared spectroscopy, Guatemalan purple jadeite lacks the absorption lines at 1167 cm^{-1} and 745 cm^{-1} . However, no significant differences were observed in the Raman spectroscopy and photoluminescence spectra. The Raman spectral results indicate that purple jadeite from both sources is relatively pure jadeite.

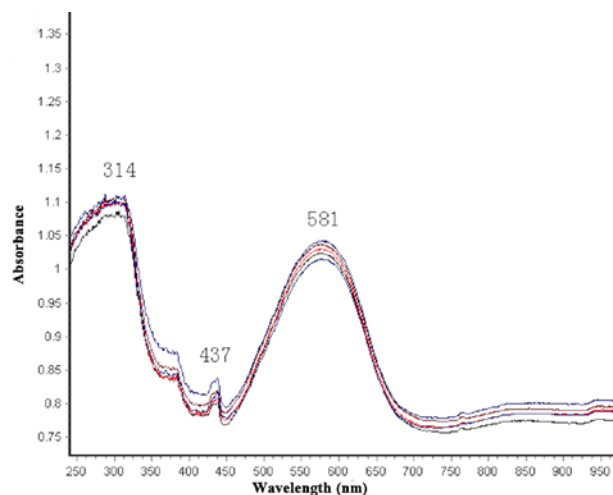


Figure 2: UV-Vis absorption spectra of the purple jadeite from Burma.

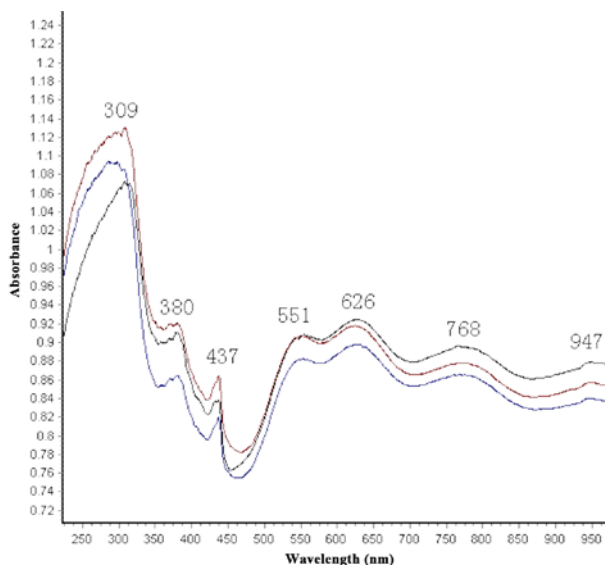


Figure 3: UV-Vis absorption spectra of the purple jadeite from Guatemala.

Conclusion

The findings of this study indicate that both Burmese and Guatemalan purple jadeite are primarily composed of relatively pure jadeite. However, significant differences in trace elements are observed, with Burmese purple jadeite exhibiting elevated manganese and lower titanium concentrations, whereas Guatemalan purple jadeite displays reduced manganese and higher titanium levels. These compositional variations contribute to the distinct differences in their absorption spectra. Through detailed compositional analysis and ultraviolet-visible absorption spectroscopy, a reliable distinction between Burmese and Guatemalan purple jadeite can be made.

The samples examined in this abstract predominantly consist of purplish-red jadeites from Burma and bluish-purple jadeites from Guatemala. However, it should be noted that bluish-purple jadeites are also present in Burmese jadeite on the market, and premium-quality Guatemalan jadeite may exhibit purple hues similar to that of the Burma origin. For instance, B-004 (purple jadeite sample from Burma) and G-002 (purple jadeite sample Guatemala) display remarkably similar purple coloration in both hue and saturation, rendering them difficult to differentiate by visual observation alone. Therefore, I conducted a comprehensive analysis using several types of advanced instrumentation. Among these, the ultraviolet-visible (UV-Vis) absorption spectroscopy proved to be diagnostically significant, offering an effective means of distinguishing the origin of the jadeite samples.

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Dyed purple Jade (Lavender jade or purple “Fei Cui”) from Mandalay's jade market, Myanmar

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Keywords: Fei Cui, Jadeite, Lavender jade, Dyed jade

Introduction

Jade or “Fei Cui” are trade names for a stone that typically contains only one or more main mineral components, including jadeite, omphacite, and kosmochlor. These mineral components exist in the form of small crystallites tightly packed together (aggregate), with many grain boundaries between them. As a result, jade also known as “fei cui” commonly exhibits varying degrees of translucency. Not only green jade, but also purple jade is increasingly popular in the gem market. This is due to its beautiful, rare and unique colouration. Moreover, there is a belief that purple jade (also known as lavender jade or purple “fei cui”) is an auspicious gem, bringing good luck to the person who owns or wears it. Due to the rarity of such shades in nature, high-quality purple jade holds significant value and is in demand in the current market. Since 2023, The Gem and Jewelry Institute of Thailand (Public Organization) or GIT, has received reports from entrepreneurs indicating an

increase in the quantity of various shades of purple jade entering the market compared to previous years. Furthermore, there is a significant possibility that some of the purple jade available in the market may have been artificially enhanced colour through dyeing processes. This situation has raised significant concerns within the trade, impacting both entrepreneurs' and consumers' confidence in the authenticity of purple jade available in the trade and market.

Materials and Methods

GIT received a collection of purple jade samples, acquired from a dealer at the Mandalay Jade and Gemstone Market in Myanmar. The collection comprises 15 samples with a weight range of 3.30 to 9.16 ct. Among these, three samples were identified as natural purple jade while the remaining twelve were suspected to have been dyed, as illustrated



Figure 1. 12 Dyed purple jade samples from the Mandalay jade market in Myanmar, ranging from 0.11 to 1.04 ct.

Photo by M. Seneewong-Na-Ayutthaya.

in Figure 1. To determine evidence of dye treatment, the samples were analyzed using both microscopic examination and advanced instruments, including a Raman microscope spectrometer (Renishaw inVia), FTIR spectrometer (Thermo Scientific, Nicolet iN10) and UV-Vis-NIR spectrophotometer (PerkinElmer, Lambda 1050).

Results and Discussions

All samples have been confirmed to be natural jadeite-jade through basic instrument testing. Fluorescence testing revealed that all samples were inert when exposed to short-wave and long-wave UV radiation. Visual observation of the samples with the naked eye and a 10x loupe revealed that they can be grouped into two categories: dark purple and light purple. Most of the samples exhibited irregular colour and texture characteristics, typical for jade which is dyed. Poor-quality jade, characterized by pale colour and larger crystal grain size, are often treated by a dyeing process. This textural property allows the dye to penetrate and spread more effectively along the crystal grain boundaries. Additionally, it's worth noting that jade with purple colour is rare in the famed jade deposits in Myanmar. High-magnification

observation under a gem microscope, as illustrated in Figure 2, revealed distinct internal characteristics. The natural purple jade sample exhibited a light purple colour that is smoothly distributed among the fine, small crystal grains. The more intense purple jade samples revealed evident features indicating that they were dyed, such as distinct colour patches and concentrations along grain boundaries as a result of the dyeing treatment.

Subsequently, the purple jade beads were cross-sectioned to study the extent of dye penetration, as represented in Figure 3. The results revealed significant colour modifications in all the purple jade samples through the dyeing treatment. Strong evidence of dye infiltration was observed within the central area of the samples, with the intensity of the dye gradually decreasing to a very light colour toward the edge of the samples (in the thinnest area of the cross-section). Based on the observed characteristics of the dye penetration we assume that a vacuum-assisted dyeing technique was used. For this, the jade bead is first placed in a vacuum chamber to remove air, water, or small particles from the grain boundaries. This pre-treatment enables the jade to absorb the dye more effectively, as the dye can quickly

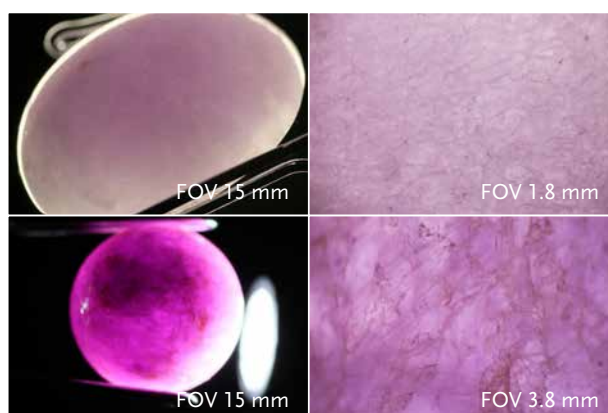


Figure 2. (top) Natural purple jade exhibiting a light purple colour with smooth distribution in fine crystal grains, and (bottom) dyed purple jade displaying areas of colour patches caused by dyeing, with dye concentration observed along the crystal grain boundaries. Photomicrographs by M. Seneewong-Na-Ayutthaya.

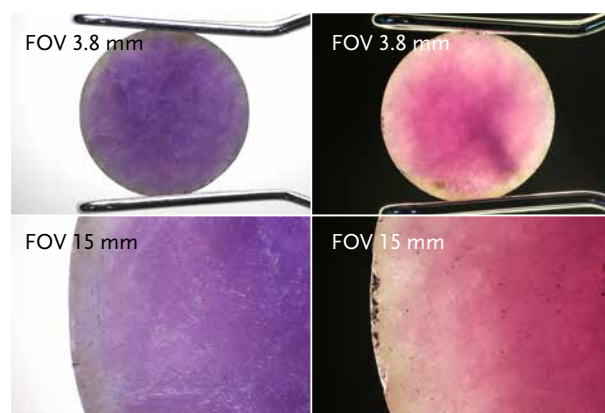


Figure 3: A cross-section of dyed purple jade sample was studied in bright-field and dark-field illumination to observe the extent of dye penetration. Dye concentration was observed in the centre of the bead, while a pale colour was evident near the surface of the bead. Photomicrographs by M. Seneewong-Na-Ayutthaya

penetrate between the crystal grain boundaries (Ou Yang & Humphrey, 2015) and thus allows the dye to penetrate deeply into the crystalline texture of the jade bead. Furthermore, the lighter colouration observed at the edge of the cross-section sample may result from cleaning procedures commonly employed during the process of making jade jewelry. Specifically, washing the bead with a mild acidic solution, such as plum juice, is a traditional practice used to clean off various chemical residues from the grinding and polishing process (Yan, 2019).

An additional experiment involved soaking purple dyed samples in ethyl alcohol for 48 hours. After this test it was observed that the jade samples displayed their original colour (pale yellowish green) near the edge of the samples (see Figure 4) as they had assumingly before the dyeing treatment.

Figure 5 displays the absorption spectra of a purple jade of natural colour (spectrum at the bottom), a reference sample of dyed purple jade, and two spectra of dyed jade

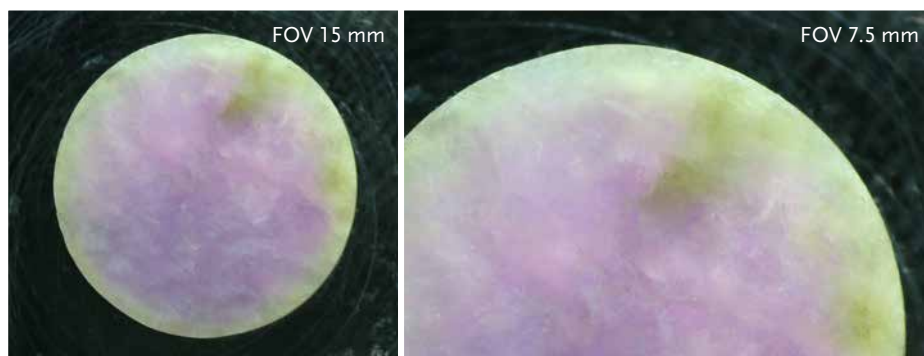


Figure 4: A purple dyed jade cross-section after soaking in ethyl alcohol for 48 hours. It clearly displays its original colour before the dyeing treatment along the edge of the sample. Photomicrographs by M. Seneewong-Na-Ayutthaya

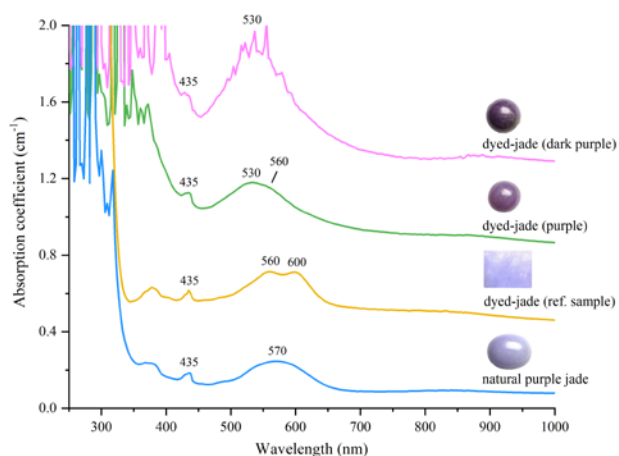


Figure 5: UV-visible spectra of natural purple jade and dyed purple jade samples.

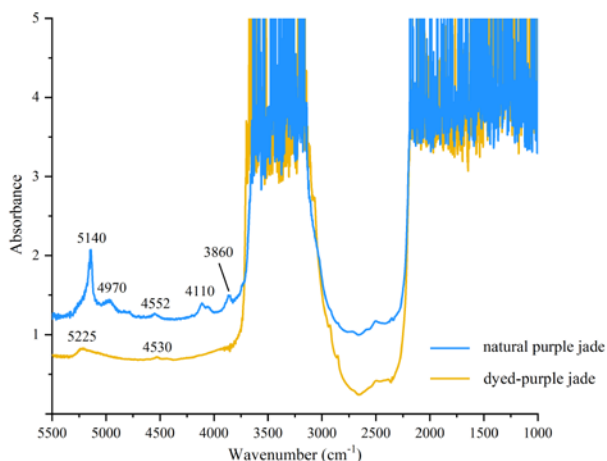


Figure 6: FTIR spectra of natural purple jade and dyed purple jade samples.

of purple and dark purple colour. In dyed samples investigated for this study, a main absorption band was observed at 530 (with a small shoulder at 560 nm) related to the purple dye. The dyed reference sample of light purple colour exhibits two broad absorption bands at 560 and 600 nm, respectively. In contrast to this, the purple jade of natural colour exhibits an absorption peak at 573 nm due to the presence of manganese (Mn^{3+}), causing the purple colour in natural jade, and in addition a peak at 453 nm, which represents iron (Fe^{3+}) (Lu, 2012; Abduriyim *et al.*, 2017). The comparison of the absorption spectra clearly indicates that the purple jade beads investigated had been dyed. Figure 6 shows the FTIR spectra of a dyed purple sample compared to the untreated natural purple jade. A small wax-related appears around 2900 cm^{-1} , with no absorption peaks indicating polymer treatment (Zhang *et al.*, 2013; Senee-wong-Na-Ayutthaya, 2022; Promwongnan *et al.*, 2023). This suggests that the dyed purple jade sample can be classified as C-Jade, a type of jade that has only been enhanced in quality through dyeing.

Concluding remarks

From the analysis of purple jade (also known as lavender jade or purple “fei cui”) samples from the Mandalay Jade Market in Myanmar, it was suspected that the colour had been modified by dyeing. From a total of 15 purple jade beads investigated for this study, 12 samples were found to be impregnated with purple dye. The experimental results showed that the dye penetrated deeply into the jade, likely facilitated by using a vacuum technique to help spread the dye along crystal grain boundaries. Consequently, this type of treatment can be identified through careful microscopic examination and confirmed using standard spectroscopic techniques by experienced gemmologists.

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Pink to Red and Violet Diamonds from the Argyle Mine, Australia: Properties, Color Origin and Latest Research Results.

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Fig. 1. A range of pink to red to violetish gray to violet diamonds from the Argyle mine in Australia.

The Argyle Mine was by very far the largest producer of pink to red diamonds worldwide and the only producer of violet diamonds (Fig. 1). The mine that started its operation in 1983 was finally closed in November 2020, after 37 years of successful operation. While the mine was famous for these rare colored diamonds, these represented only a very small percentage of its production, far more what was produced by the Argyle deposit were brown diamonds.

Significant research has been conducted by our laboratory concerning the still puzzling color origins of these diamonds and their defect characterization. While being produced in the same Lamproite body which is characterized by its high percentage of diamonds that have undergone plastic deformation – namely brown, pink and red diamonds – the extremely rare and unusual violet diamonds have no link to plastic deformation whatsoever. These stones are much higher in nitrogen and hydrogen than the pinks and reds and we found that their color is linked to a combination of absence of N3 center absorption and distinct content of nickel nitrogen defects, possibly in conjunction with hydrogen. Whether the hydrogen plays an actual role in the coloring defects is not clear, as contrary to the classical attribution of the violet color to hydrogen, we could not find any major link between the color and

(IR active) hydrogen content. During this research we have defined the spectral structures of the major defects that are responsible for the violet color, and curiously one of the main defects involved has its ZPL quite far in the NIR, namely at 1330 nm (Hainschwang et al, 2021). We were able to define the spectral structure of the 1330 nm center at 77K and define its link to part of the absorptions seen in the visible spectral range (Fig. 2).

Work performed by our lab on “olive”, brown to pink and red diamonds with the aim to understand why plastic deformation can produce these colors has pointed towards the temperature during plastic deformation events being responsible for the different hues. The different temperature conditions result in somewhat different vacancy-cluster-type defects and in consequence the different colors. Annealing experiments of type I diamonds exhibiting two colors of “graining” revealed that the “olive” hue is least stable, the brown hue is more stable and the pink hue is most stable to annealing. A link between the amber center (AC) absorption in the infrared spectrum at 4063 cm^{-1} with “olive” color and the AC at 4167 cm^{-1} with brown color could be clearly established: diamonds of “olive” color with distinct 4063 cm^{-1} AC absorption turned brown upon annealing at 1500°C and the AC at 4063 cm^{-1}

annealed out while the AC at 4167 cm^{-1} increased distinctly (Fig. 3). The brown colors start to anneal out at temperatures from 1900°C while pink seems to remain unaffected to over 2000°C . Since at that temperature other defects emerge in type Ia diamonds, it is difficult to define a temperature when the pink color is effectively annealed out. From experiments with type IIa diamonds of mixed brown-pink color, we know that pink color in type IIa diamonds is more stable than brown color, as such stones can be HPHT treated into a purely pink color, hence the brown color anneals out earlier than the pink color. All indications show that the same is true for the pink color of type Ia diamonds.

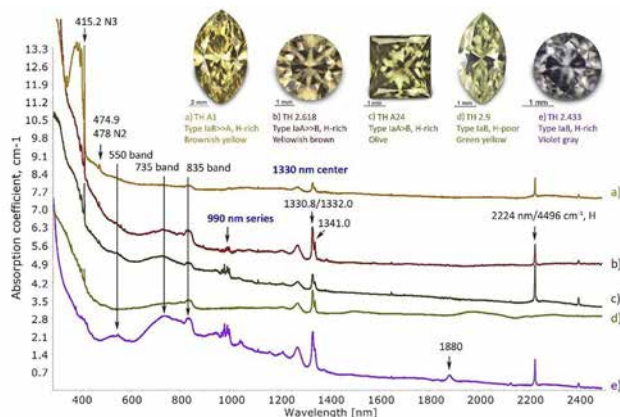


Fig. 2. The normalized and extended low temperature UV-Vis-NIR spectra of some differently colored "1330 nm diamonds" included in this study. It is obvious that sample TH 2.9 is different to all others, since while it exhibits a strong 1330 nm center, it is very low in hydrogen and lacks the broad bands at 735 and 550 nm. The spectra have been shifted vertically for clarity.

It is suggested that the three different colors of plastically deformed diamonds are the result of plastic deformation at different temperatures: olive is the result when plastic deformation occurs at the lowest temperature, brown is the resulting color when plastic deformation occurs at an intermediate temperature and pink results from plastic deformation at the highest temperature. This model is the only way one can logically explain pink and brown graining in one and the same diamond: two separate events of plastic deformation, the first one at higher temperature resulting in pink graining, and the second one at lower temperature resulting in brown color (Hainschwang *et al.*, 2020b).

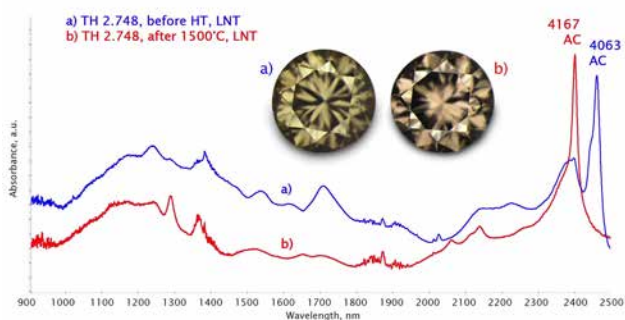


Fig. 3. The normalized and low temperature NIR spectra of a type Ia "olive" diamond before and after annealing at 1500°C . The sample changed color from "olive" to brown by the annealing while the amber center at 4063 cm^{-1} anneals out and the amber center at 4167 cm^{-1} increases distinctly.

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Shining light on “480 nm band” diamonds: Gemmological characteristics of an unusual group of diamonds with yellow and red luminescence

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Fancy colour diamonds that derive their colour from nitrogen-related defects (such as N3/N2 centers or C-centers) are relatively common and well-studied. An underexplored group derives its colour from a broad absorption band centered around 480 nm, which is hypothesized to be related to oxygen impurities (Hainschwang *et al.* 2008). These diamonds are referred to as “480 nm band diamonds” and their properties have been described in a number of studies (e.g. Breeding *et al.* 2020, Lai *et al.* 2024). A rare subgroup of “480 nm band” diamonds are “chameleon diamonds”, which temporarily change colour upon heating or prolonged storage in the dark. Common features of these diamonds are (De Weerd & Van Royen, 2001, Fritsch & Delaunay, 2018):

- 1) A band at 426 nm in UV-Vis-NIR absorption spectroscopy, related to the 480 nm absorption band.
- 2) A peak at 1241 cm⁻¹ in FTIR, which is thought to be linked to the 480 nm band.
- 3) Yellowish-green fluorescence in DiamondView, often accompanied by irregular zones of blue and bright green fluorescence.
- 4) Nickel-related defects in Photoluminescence Laser spectroscopy (PL), such as the 883/885 nm doublet, S2 and S3 centers.
- 5) Hexagonal or rounded plate-like inclusions, identified as graphite platelets by Shiryaev *et al.* (2023)
- 6) Hydrogen-related absorptions in FTIR spectroscopy.

An often-overlooked feature of “480 nm band” diamonds, however, is their luminescence reaction. When illuminated with long-wave ultraviolet (LWUV) light (365 nm) these diamonds typically show a yellow luminescence, whereas under blue light (450 nm) they display red luminescence (Collins & Mohammed, 1982). The red luminescence is produced by the 480 nm absorption band (2.6 eV), while yellow luminescence is linked to a vibronic system with a zero-phonon line (ZPL) at 456 nm (2.721 eV).

The aim of this study is to provide a detailed gemmological characterization of the relatively unknown “480 nm band” diamonds. Diamonds were screened with a custom-made luminescence tester, containing a LWUV and blue light source (Figure 1). In this way, we could quickly identify whether diamonds have yellow and red luminescence, which implicates that they contain the “480 nm band”.

More than 50 diamonds were selected for detailed spectroscopic and gemmological analysis. All diamonds show yellow and red luminescence under the luminescence tester. The diamonds were studied by FTIR and UV-Vis-NIR spectroscopy, DiamondView imaging, microscopy, long-wave and short-wave UV excitation, and some of them by other techniques such as PL and ED-XRF spectroscopy. Special attention was paid to the presence of Y centers and CO₂ content in FTIR, since a genetic relation with “480 nm band” diamonds has been suggested before (Kupriyanov *et al.*, 2020, Hainschwang *et al.* 2008).



Figure 1: Two “480 nm band” diamond under normal light (left), LWUV light (middle) and blue light (right). Both display yellow and red luminescence under LWUV and blue light, respectively.

For the selected diamonds, brown is the dominant colour grade (usually ‘reddish brown’), followed by yellow, orange and rarely olive and pink. Weights are in the range of 0.14 ct to 2.27 ct. FTIR results show that two diamonds contain distinct CO₂ bands around 655 and 2370 cm⁻¹, known as the u2 and u3 bands (Schrauder & Navon, 1993). The main Y center band at 1145-1150 cm⁻¹ is found in 7 diamonds. This is lower than expected, as Kupriyanov *et al.* (2020) concluded that the typical “480 nm” red luminescence is genetically related to Y centers. Therefore, we would expect to find Y centers in all studied diamonds. It is, however, possible that the Y centers are overshadowed by other defects (A, B, C centers), and further analysis using spectral deconvolution is necessary.

Although the “480 nm band” is responsible for red luminescence, only 15 diamonds show a distinguishable 480 nm band in their UV-Vis spectrum (Figure 2, red spectrum).

Most of the studied diamonds owe their colour to a broad absorption continuum (Figure 2, black spectrum). For these stones, other properties such as inclusions, DiamondView and PL spectroscopy correspond to those of typical “480 nm band” diamonds. We can conclude that red luminescence is a more sensitive indicator than UV-Vis spectroscopy for the presence of the “480 nm band” defect.

In conclusion, our study shows that screening for luminescence reaction is a useful gemmological tool to quickly identify diamonds with “480 nm band” properties, since the combination of yellow and red luminescence is unique to these diamonds. We identified “480 nm band” diamonds over a range of colours, extending beyond the yellow and orange diamonds that have been typically studied. This allows us to investigate possible genetic links between the “480 nm band”, CO₂ diamonds and Y centers.

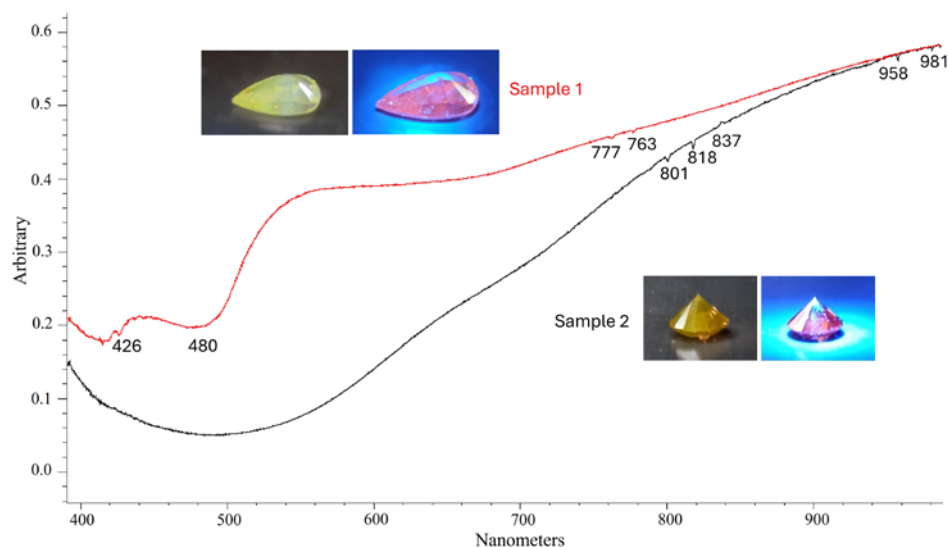


Figure 2: UV-Vis-NIR transmission spectra of two “480 nm band” diamonds. Red spectrum: Diamond with a typical 480 nm absorption band (Sample 1). Black spectrum: Diamond with yellow and red luminescence but no 480 nm band in UV-Vis-NIR (Sample 2). The samples are the same as those shown in Figure 1.

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Identification of melee size synthetic coloured diamond for jewellery

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Introduction

Synthesis methods to produce large and high-quality diamond for jewellery have been improved over the years, and large, colourless faceted stones weighing more than 50 ct have been realized (Eaton-Magaña *et al.*, 2024). In addition, melee size fancy-coloured diamonds are also useful as components of jewellery, and the melee size synthetic coloured diamonds have been seen at jewellery shows in Japan and at the Tucson Show in USA. Jewellery assembled with the melee size diamonds is also sold in shops. Two methods are used to produce synthetic diamonds for jewellery: the HPHT method and the CVD method. It is important for gemmological identification to know clearly the different characteristics of the synthetic diamonds made with the two methods.

Material and Methods

Samples we bought at the 2024 IJT (International Jewelry Tokyo) for the present study are 35 pieces of melee size synthetic fancy-coloured diamonds with size of 0.10 ct or less which were labeled as CVD synthetic diamonds. After testing with standard gemmological instruments, the 35 samples were classified into seven categories according to colour, with five pieces for each category: (1) Green, (2) Greenish Blue, (3) Yellow, (4) Pink, (5) Orangy Pink, (6) Reddish Orange and (7) Orange (Figure 1).

These stones were observed under a gemmological microscope, and were measured with infrared and UV-VIS absorption spectroscopy. Deep UV luminescence images were obtained using DiamondView™. Photoluminescence spectra were taken for all samples being immersed in liquid nitrogen, using a Raman spectrometer with microscope with five different lasers for excitation at 457, 488, 514, 633 and 830 nm.



Figure 1: Fancy colour melee size diamond studied in this study (Upper row, from left: Orange, Reddish Orange. Middle row, from left: Yellow, Pink, Orangy Pink. Lower row, from left: Green, Greenish Blue)

Results and Discussion

(1) Green and (2) Greenish Blue: In all samples in these categories, the concentrations of nitrogen were below the detection limit according to the IR absorption spectroscopy, and deep UV fluorescence images showed characteristic patterns of HPHT synthetic diamond. From both green and greenish blue samples, a distinct 741 nm (GR1) peak was observed in the UV-VIS absorption spectra (Figure 2) and PL spectra. For green samples, peaks at 575 nm (NV⁰), 741 nm (GR1), 488.9 nm and 470.2 nm (TR12) were detected in the PL spectrum. The 575 nm peak was stronger than the 741 nm peak. These results suggest that the green and greenish blue samples have been produced by irradiation of type IIa HPHT-synthetic diamonds. The greenish blue samples have been annealed at a lower temperature (around 500 °C) after the irradiation.

(3)Yellow: All samples are type Ia with high concentrations of A-centre nitrogen according to the IR absorption measurements. Weak C-centre and 3107 cm⁻¹ (N₃VH) peaks were also detected. In addition, deep UV fluorescence images confirmed that all of them are HPHT synthetic diamond. 544.5 nm and 523.8 nm peaks associated with cobalt were detected in PL spectra (Figure 3). These suggest that they are so called “HIH” diamond which means a HPHT diamond were irradiated and treated in a HPHT condition (Hainschwang and Notari, 2011.).

(4)Pink, (5)Orangy Pink, (6)Reddish Orange and (7)Orange: In all samples, faint nitrogen-related absorption was observed in the IR absorption spectra. A very strong 575 nm (NV⁰) peak and a weak 637 nm (NV⁻) peak were observed in

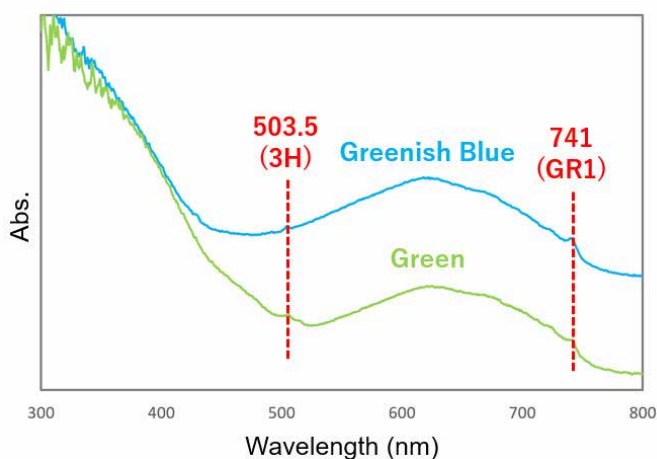


Figure 2. UV-VIS absorption spectra of the blue and greenish blue samples

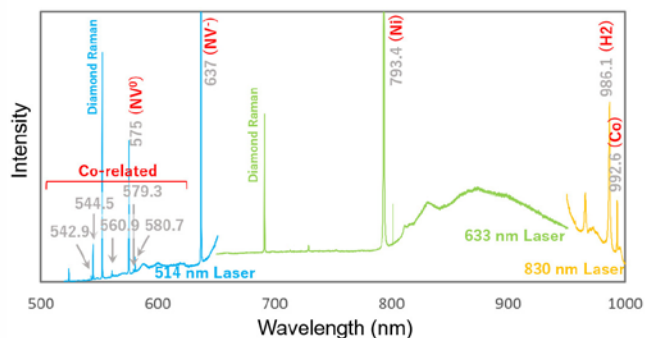


Figure 3. PL spectra of yellow samples (514, 633 and 830 nm laser excitation)

a PL spectrum, while in reddish orange samples a 741 nm (GR1) peak was observed in both UV-VIS absorption s and PL spectra. The 575 nm (NV⁰), 637 nm (NV⁻) and 741 nm (GR1) centers were formed by irradiation and subsequent annealing. The intensity ratio of the 575 nm (NV⁰) peak to the 637 nm (NV⁻) peak was in the order of pink < orangy pink < reddish orange < orange. This is the same order as the concentration of C-centres, and the concentrations may influence the colour. For two of the orange samples? There are five right?, the deep UV fluorescence image and the presence of 737 nm in the photoluminescence spectra indicate that they are CVD synthetic diamond, while all the others are HPHT synthetics due to their specific fluorescence images (Fig. 4).

Conclusion

In this study, gemmological examinations were carried out on melee-sized fancy coloured synthetic diamonds in loose form. We bought all 35 samples which were labelled as CVD synthetic diamonds, but 33 pieces of them were HPHT synthetic diamond and two of them were CVD synthetic diamonds. They had undergone multiple treatments with irradiation, annealing and HPHT treatments. If these were submitted in the form of gems set in jewellery for identification, the analysis methods would be limited and poor report may have been published. Therefore, we need to have full knowledge of characteristics of all type synthetic diamond made with the different methods and treated in the different ways so that error-free reports on the identification are published.

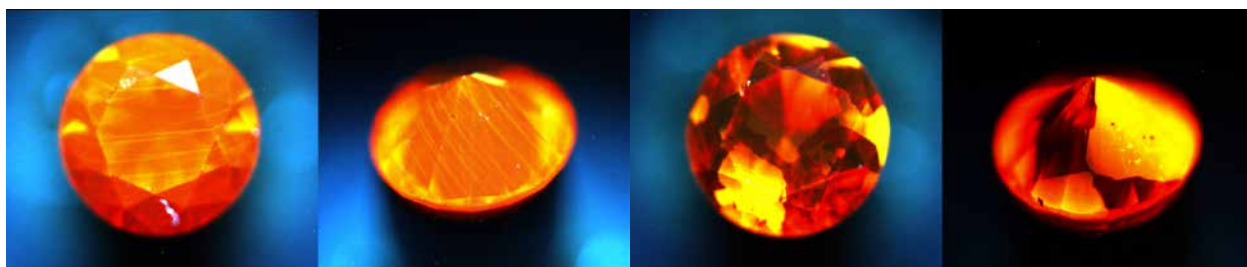


Figure 4. Step-flow structure specific to CVD synthetics (left) and sector zoning structure specific to HPHT synthetics (right)

References:

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Nitrogen aggregates in yellow HPHT and CVD synthetic diamonds.

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The identification of natural and synthetic diamonds is an important gemmological task. Structural impurities in diamonds play a critical role in this task because they characterize the diamond growth and post-growth conditions, which differ between natural and synthetic diamonds. Optical centers in yellow diamonds partially overlap with those in colorless diamonds but also exhibit distinct differences. In recent years, there has been a growing number of synthetic yellow diamonds on the market, many of which are irradiated and annealed to alter their color. As a result, gemmological laboratories face challenges in the identification of yellow diamonds.

We studied synthetic yellow diamonds of both HPHT and CVD origin using a combination of spectral methods. Since 2023, the N3 center (415 nm) - previously considered an indicator of natural diamonds - has been observed in synthetic diamonds. In natural diamonds, the N3 center can be detected in both absorption and photoluminescence (PL) spectra, whereas in synthetic diamonds, its peak is weaker and observable only in PL spectra. Beginning in 2024, the N3 center has become common in synthetic yellow diamonds submitted by clients, with a more distinct peak in HPHT samples (Fig. 1a) and a weaker peak in CVD

samples (Fig. 2). In natural diamonds, the N3 center arises from nitrogen aggregation during prolonged geological processes, while in synthetic diamonds, it can form during post-growth treatments (Vins *et al.*, 2021).

In addition to the N3 center, a prominent PL series with zero-phonon lines (ZPLs) at 523.6 nm and 626.3 nm was detected in all HPHT yellow diamonds (Fig. 1a). These optical centers form in type Ib diamonds after irradiation and annealing (Dobrinets *et al.*, 2013). The 580 nm center is attributed to Ni-related defects (Yelisseyev and Kanda, 2007).

FTIR spectroscopy revealed that the HPHT yellow diamonds are type Ia, with nitrogen detected in the 1100–1300 cm⁻¹ range. One sample exhibited a platelet-related peak at 1364 cm⁻¹, a feature typically associated with natural post-growth annealing. Two synthetic yellow CVD diamonds (1.03 ct, cushion-cut) submitted by clients showed weak nitrogen-related peaks at 1250–1380 cm⁻¹ and hydrogen-related peaks at 3107 cm⁻¹ in FTIR spectra. Their PL spectra displayed weak N3 and H3 centers, along with a [Si-V]⁻¹ doublet at 736,6/736,9 nm (Fig. 2).

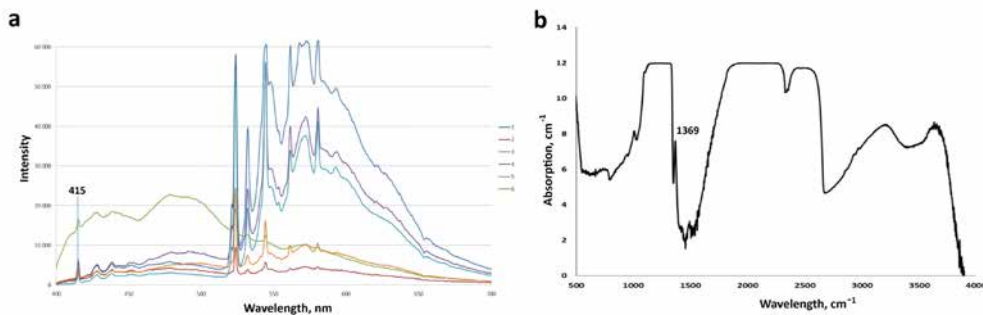


Fig. 1. a – PL spectra of 6 yellow synthetic HPHT diamonds, recorded at liquid nitrogen temperature. Excitation 365 nm. b – FTIR spectrum of yellow HPHT synthetic diamond with B2 (platelets) peak.

Experimental Stage

To investigate nitrogen aggregation mechanisms, we conducted post-growth experiments on three CVD diamond plates (0.15 ct, 0.12 ct, and 0.65 ct) and one HPHT diamond plate (6×6×1 mm, 0.86 ct). The samples were electron-irradiated (3 MeV) and subsequently annealed at 1750–1800°C.

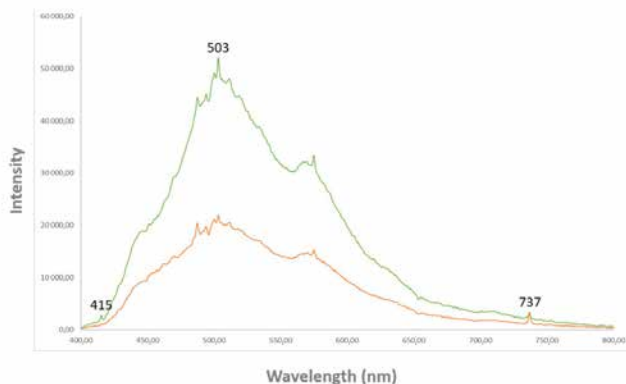


Fig. 2. PL spectra of two yellow CVD diamonds recorded at liquid nitrogen temperature with excitation 365 nm. There are peaks at 415 nm, 503 nm, 575 nm, and 737 nm.

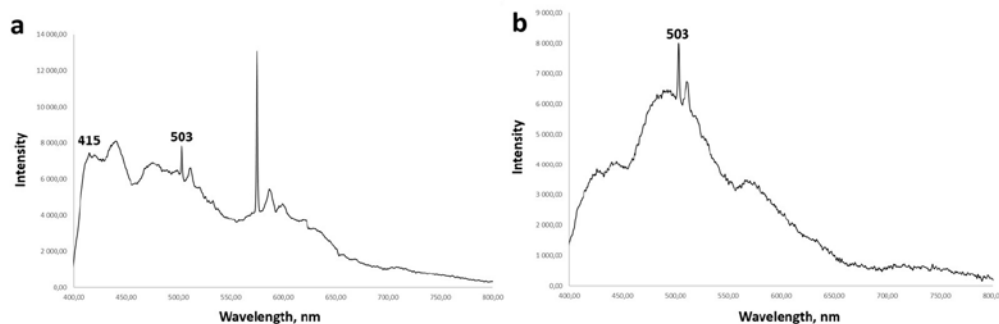


Fig. 3. PL spectra of synthetic CVD (a) and HPHT diamond (b) plates after irradiation and annealing.

Spectral characterization was performed before and after each stage. After irradiation, vacancies or GR1 centers (741 nm) appeared in the spectra. Following annealing, the N3 center emerged in CVD plates in PL spectra recorded at liquid nitrogen temperature. In annealed CVD plates, apart from 415 nm, additional 503 nm, 575 nm, and 737 nm peaks appear in PL spectra recorded at liquid nitrogen temperature (Fig. 3a). Same peaks are recorded at polished CVD diamonds (see Fig. 2). In the annealed HPHT plate we did not observe the N3 center in PL spectra (Fig. 3 b) but see the H3 center (503 nm) that appeared after annealing. In the FTIR spectrum of the annealed HPHT plate there are A and C centers recorded as well as weak B2 (platelets) peaks. Probably further annealing is required for this sample in order to induce N3 center.

Our experiments confirm that the N3 center forms in synthetic diamonds after irradiation and annealing. The proposed mechanism involves:

1. As grown stage: Single nitrogen atoms are present (more abundant in HPHT, less in CVD diamonds).
2. Irradiation: Generates vacancies in the diamond lattice.
3. Annealing: Facilitates the formation of H3 aggregates, a fraction of which evolve into N3 aggregates.

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Diamonds from Than Lwin River near Mong Hsu, Myanmar

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Introduction

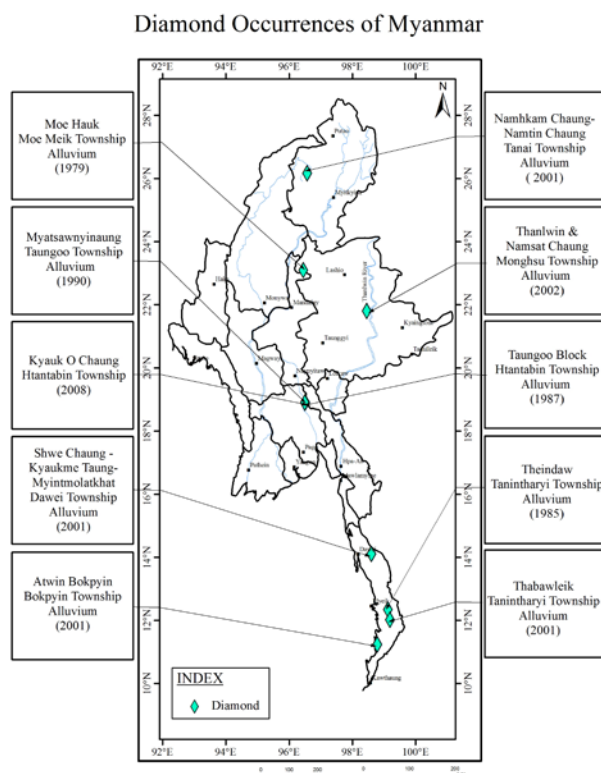
Sixteen rough diamonds were acquired in 2023 by co-author TTS from a gem dealer linked to Mong Hsu, Shan State, Myanmar. While diamonds have been reported in Myanmar, most are from limited alluvial deposits. Discoveries include Kyeindaw (1959), Mohaung near Momeik (1989), and tin mining areas in Kyaukmedaung (1981) and Myat-

sawnyinaung near Taungoo, both within the Mergui Group. (Griffin *et al.*, 2001; Win *et al.*, 2001; Kyaw Thu & Khin Zaw, 2017). In 1985, 3,000 alluvial diamonds, including a 10.3-carat stone, were recovered from Theindaw mine near Myeik. Between 1995–1998, small quantities of diamonds were found during gold panning along the Than Lwin River near towns including Tangyan, Pangsang, and Mong Hsu (21°51'22"N, 98°32'36"E) (Personal communication: Nyunt Htay & Ma Gjam). The sixteen diamonds used in this study originate from this locality. As noted by Griffin *et al.* (1998), Win *et al.* (2001), and Nyunt Htay (2012), Myanmar's diamonds are largely alluvial, with no confirmed primary sources to date.

Materials and Methods

Sixteen rough diamonds ranging from 0.18 to 1.02 carats (Fig. 2) exhibited various crystal habits, including octahedral, trisoctahedral, tetrahedral, macle, and rhombic dodecahedral forms. Surface and internal features were examined using a GIA binocular microscope and long-wave ultraviolet (LWUV) fluorescence. Raman spectra were collected with a Renishaw inVia™ confocal Raman microscope (532 nm, spectral range from 100 to 2000 cm⁻¹), while FTIR analysis was performed using a JASCO FT/IR-4X spectrometer (absorbance range 400–4000 cm⁻¹). These methods assessed structural integrity, surface alteration, and defect-related features.

Fig. 1. Various location of diamond deposits discovered in Myanmar over the years including the recent finding along Than Lwin river near Mong Hsu Township (map prepared by Thet Tin Nyunt, 2025)



Samples & Results

Microscopic Observations

Microscopic analysis revealed a variety of surface and internal features across the specimens. Notable observations included dark crystal inclusions, green and brown surface spots, percussion marks indicative of mechanical stress, hexagonal etch pits, trigons, and lamellar growth zoning. Green spots were especially prominent on some samples and suggested exposure to radioactive environments. Per-

cussion marks implied histories of plastic deformation and mechanical impact.

Raman Spectroscopy

Raman spectroscopy revealed a dominant diamond peak at 1331 cm^{-1} across all specimens, confirming preservation of the crystalline diamond structure. Secondary peaks between $1442\text{--}1449\text{ cm}^{-1}$ were detected in several specimens, attributed to disordered carbon associated with surface-reaching inclusions or stress-altered regions. Additional features at 381 cm^{-1} and 1216 cm^{-1} (Fig. 4) observed near green surface spots suggest radiation-induced defects, while a 1419 cm^{-1} peak, found in diamonds with brown spots and percussion marks, reflects plastic deformation. Low-frequency peaks around 154 cm^{-1} (Fig. 5) were also recorded in some samples, indicating localized lattice stress or vibrational modes activated near inclusion zones (Pasteris & Wopenka, 1991; Nasdala *et al.*, 2004; Zaitsev, 2001; Fischer *et al.*, 2009; Smith & Dent, 2010).



Figure 2. Sixteen pieces of rough diamonds from Mong Hsu, Myanmar with size range from 0.18 ct to 1.02 cts, with total weight 7.86 carats. (Photo by Tay T.S.)

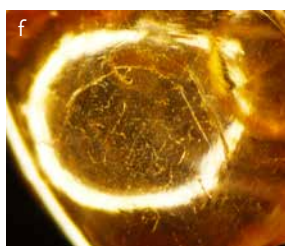
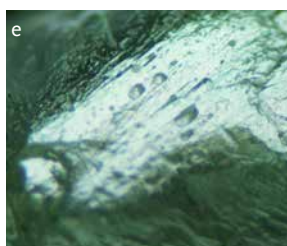
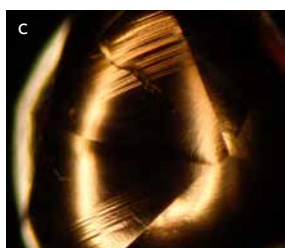
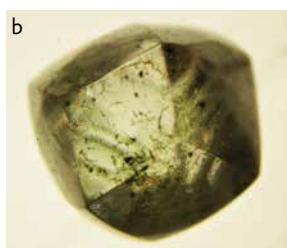


Fig. 3. Surface features of the 16 samples: (a) brown radiation spot (20x); (b) green radiation spots (20x); (c) lamellae growth marks (10x); (d) hexagonal growth marks (10x); (e) Etch marks (10x); (f) percussion marks (10x) (Microphotography by TTS).

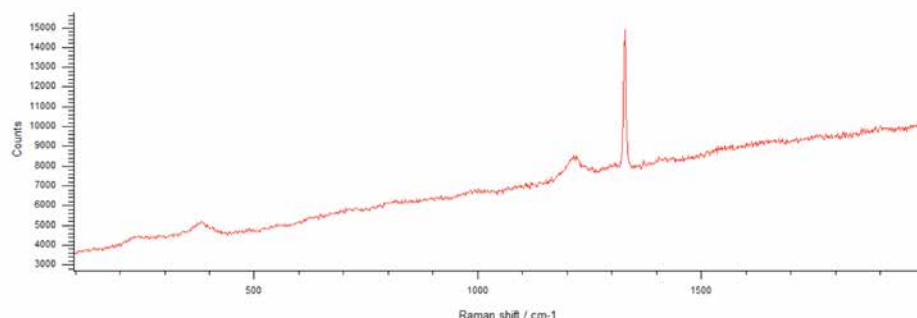


Fig. 4. Raman shift spectrum for specimen no. 11 showing peaks at 381 cm^{-1} , 1216 cm^{-1} , 1331 cm^{-1}

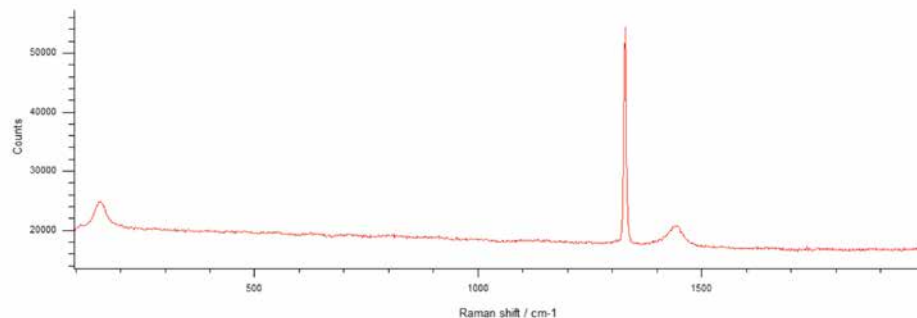


Fig. 5. Raman shift spectrum for specimen no. 6 showing peaks at 154 cm^{-1} , 1331 cm^{-1} , 1442 cm^{-1}

FTIR

FTIR spectroscopy confirmed that all diamond specimens are Type IaAB, characterized by absorption features at $\sim 1004\text{--}1013\text{ cm}^{-1}$ and $1170\text{--}1370\text{ cm}^{-1}$, corresponding to A-aggregated and B-aggregated nitrogen defects. Additional absorptions near $3106\text{--}3215\text{ cm}^{-1}$ indicate the presence of hydrogen-related defects. These results are consistent with prolonged high-temperature mantle residence, allowing nitrogen aggregation into stable forms, and suggest possible interaction with mantle fluids during or after formation (Boyd *et al.*, 1994; Woods, 1986).

Ultraviolet Light

Under long-wave UV light, most specimens exhibited weak to moderate fluorescence ranging from bluish white to chalky green, while almost all were inert under short-wave UV. Stronger fluorescence, as seen in samples 6, 7, and 9, may relate to nitrogen-related defects or localized structural distortion (Zaitsev, 2001). Specimens with green or brown radiation spots frequently displayed moderate blue fluorescence. Inert samples, including specimen 16, may

reflect low nitrogen concentrations or absence of optical defects (Boyd *et al.*, 1994). These observations support the natural origin and defect diversity among the Mong Hsu diamonds.

Discussion

Microscopy revealed features such as percussion marks, radiation spots, and resorption patterns. Under long-wave UV light, most diamonds fluoresced faint to moderate blue. Raman spectroscopy confirmed a stable diamond lattice (1331 cm^{-1}) with additional peaks ($1442\text{--}1449$, 1419 , 381 , 1216 cm^{-1}) indicating disordered carbon, plastic deformation, and radiation-related defects (Pasteris & Wopenka, 1991; Zaitsev, 2001). FTIR analysis identified all specimens as Type IaAB, showing A- and B-aggregated nitrogen and hydrogen-related defects (Boyd *et al.*, 1994; Woods, 1986), supporting a natural mantle origin with complex post-growth histories. A summary of results is given in Table 1 below.

Sample no.	Weight (Carats)	Colour	Shape	Surface Features/ Inclusions	UV	Raman cm ⁻¹	FTIR Wave-length cm ⁻¹	Diamond Type
01	0.295	Very light greenish yellow	Octahedral	Semi-transparent; Curved percussion marks on surface	LW – Faint blue SW – inert (very faint orangy yellow)	1331	482, 1010, 1175, 1286, 1364, 3106, 3200	IaAB
02	0.294	Light yellow	Octahedral	Semi-transparent; Resorbed surface with curved per-cussion marks	LW – faint yellow SW – inert	1331	482, 1007, 1278, 1372, 2850, 2918, 3106, 3200	IaAB
03	0.349	Light yellow	Tris-octahedron	Transparent; Re-sorbed surface	LW – very faint yellow SW – inert	1331	481, 1007, 1172, 1284, 1367, 3106, 3195	IaAB
04	0.396	Light yellow	Tetra-hexahedron	Semi-transparent; Frosty surface; Brown radiation spots	LW – moderate bluish white SW – inert	1331	488, 1010, 1176, 1224, 1363, 3106, 3201	IaAB
05	0.613	Yellow-ish brown	Rhombo-dodecahedral	Transparent; Re-sorbed surface with curved per-cussion marks; Brown stains along growth zones	LW – faint yellowish green SW – inert	1331, 1419	485, 1013, 1177, 1287, 1363, 3192	IaAB
06	0.792	Yellow	Irregular/ Twinning	Transparent; Graining; Black included crystal	LW – strong yellowish-blue SW – inert	154, 1331, 1442	477, 1009, 1175, 1280, 1362, 3106, 3200	IaAB
07	1.018	Greenish yellow	Irregular	Transparent; Brown & green radiation spot inclusion; Re-sorbed surface with curved and rhombic percus-sion marks	LW – strong blue SW – inert	1331	480, 1009, 1171, 1281, 1362, 3107, 3215	IaAB
08	0.931	Brownish Yellow	Irregular	Semi-transparent; Black included crystals; Resorbed surface; Cube-shaped growth marks	LW – moderate yellowish blue SW – inert	156, 1331, 1449	488, 1010, 1174, 1280, 1362, 3106, 3203	IaAB
09	0.299	Yellow-ish Brown	Irregular	Transparent; Brown radiation spots; Resorbed surface;	LW – strong yellowish brown SW – inert	156, 1331, 1445	478, 1008, 1177, 1276, 1362, 3106, 3200	IaAB
10	0.636	Green	Irregular	Semi-transparent; Green radiation spots; Curved percussion marks; Shield-shaped growth marks	LW – moderate blue SW – inert	156, 1331, 1447	498, 1010, 1174, 1281, 1327, 1363, 3107, 3208	IaAB
11	0.525	Green	Dodecahedral	Transparent; Green radiation spots; Brown staining along fractures; Curved and rhom-bic percussion marks	LW – moderate blue SW – inert	381, 1216, 1331	1008, 1173, 1278, 1367, 3107, 3207	IaAB
12	0.473	Brown	Irregular	Semi-transparent; Curved percussion Marks; Resorbed surface	LW – moderate yellowish blue SW – inert	154, 1331, 1447	481, 1009, 1177, 1288, 1331, 1359, 3204	IaAB
13	0.234	Brown	Macle	Transparent; Re-sorbed surface; Rhombic percus-sion marks	LW – very faint blue SW – inert	1331, 1419	475, 1004, 1174, 1280, 1362, 3107, 3182	IaAB
14	0.287	Brown	Tetra-hexahedron	Transparent; Shield-shape; Resorbed surface and lamellae	LW – inert SW – inert	1331	474, 1006, 1185, 1280, 1362, 3186	IaAB
15	0.552	Green	Irregular	Semi-transparent; Hexagonal surface etch marks; Green radiation spots (also in fissures)	LW – moderate blue SW – inert	1331	480, 1010, 1183, 1288, 1362, 3106, 3208	IaAB
16	0.179	Colour-less	Octahedral	Transparent; Octa-hedron growth marks	LW – inert SW - inert	1331	482, 1010, 1180, 1282, 1371, 3107, 3192	IaAB

Conclusions

The rough diamond specimens show evidence of natural radiation exposure, plastic deformation, and surface stress features, as revealed by microscopy, UV fluorescence, Raman, and FTIR spectroscopy. It is likely to be due to exposure to alpha radiation and regional metamorphism in the alluvial deposits. Radiation defects may be further confirmed with UV-Vis spectroscopy. Despite localized defects, the primary diamond lattice remains intact. These findings reflect a complex post-growth history involving deep

mantle conditions, transport-related stress, and surface alteration. Further advanced spectroscopic studies such as photoluminescence and cathodoluminescence could refine the understanding of these processes. The study of inclusions after polishing, may also potentially be useful in studying the origins of diamonds in the future, however, it may only remain as a useful information rather than giving definitive conclusions of a diamond's origin (Smith et al, 2022).

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Morphologies and Surface Features of Diamond Samples from Southern Thailand

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Introduction

In 1955, the first report on Thai diamond was made by Dr. Payome Aranyakanon, a renowned tin exploration geologist of Thailand Department of Mineral Resources. He visited a tin mine on Phuket Island and picked a shiny grain, later identified as diamond, from a tin-tailing pile. However, stories told by villagers revealing diamonds in southern Thailand provinces have been found since about a century. These so-called headless diamonds were surprisingly found in very much younger geological terranes (cf. most of the world diamond deposits) of Southeast Asia, such as in Myanmar, Indonesia (Kalimantan) and Thailand, with no old cratons,

and no report of kimberlite/lamproite pipe occurrences in the vicinities. Studies of diamond samples from Myanmar and Thailand were also published (Griffin *et al.*, 2001; Win *et al.*, 2001; Wathanakul *et al.*, 2005). As for southern Thailand, diamonds have been collected by tin-mine workers and sold to the wealthy locals for several decades, actually since the tin mining and dredging started. The diamond samples collected for this study were mainly bought and collected time to time from the miners and villagers around Phuket, Phang Nga areas (Figure 1).

Diamond Samples

More than a hundred samples of diamond from southern Thailand were collected. Thirty-five samples were selected for being representative for this study, including fifteen new diamond samples collected from the locals off the Phuket coast. The samples were primarily studied for their colours, types, morphologies, and surface features. Examples of diamond colours as well as their morphologies are shown in Figure 2.

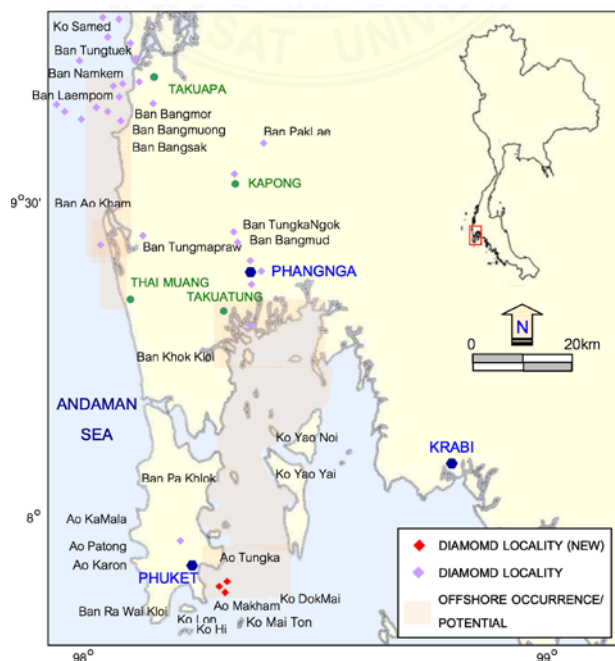


Figure 1. Diamond localities from southern Thailand, samples collected from both around Phuket and Phang Nga areas



Figure 2. Representative samples, showing Thai diamond colours

Thai Diamond Types

FTIR and UV-Vis spectroscopies reveal Thai diamond types are mainly IaAB, moderate amount of IaA, and lesser IaB, and very few of type Ib. One out of fifty samples is type IIb.

Morphologies & Surface Features

Diamond samples were inspected for their morphologies and surface features using a stereo microscope, some selected samples were carefully studied using SEM and a cathodoluminescence (CL) image. The results show that morphologies mainly include resorbed tetrahedroids (THH), a

few rounded dodecahedrons, multiple twins, octahedrons and macles. Thai diamonds show about six characteristic patterns of CL images (Figure 3), which show entirely yellow or blue, yellow/blue oscillation, a yellow centre and a blue rim, a blue core with a yellow rim, and blue with yellow patches/sectors. Radiation spots are often present in Thai diamond samples.

Their surface features show straight lamination lines, shield-shaped lamination, cross-hatched lamination, hillocks, trigons, pits, radiation brown spots, percussion scars and ruts (Figure 4). Those features reveal the diamond crystals experienced high resorption of corrosive magma, which also commonly causes a glossy surface.

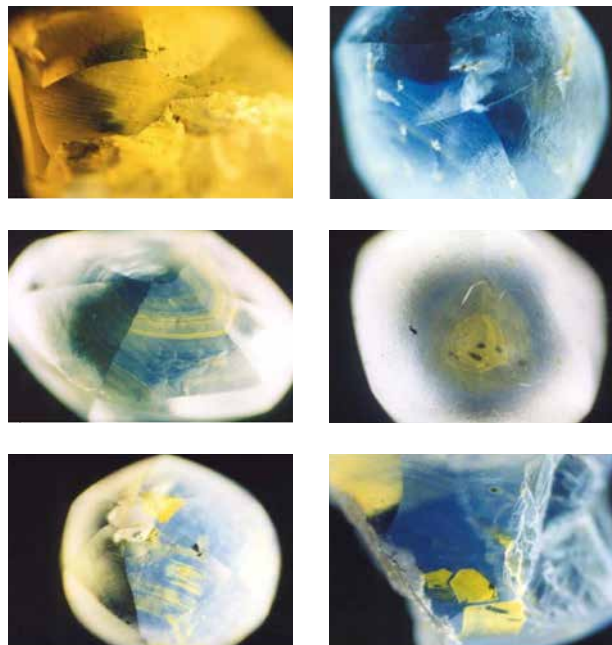


Figure 3. Examples of CL images of diamond samples, southern Thailand

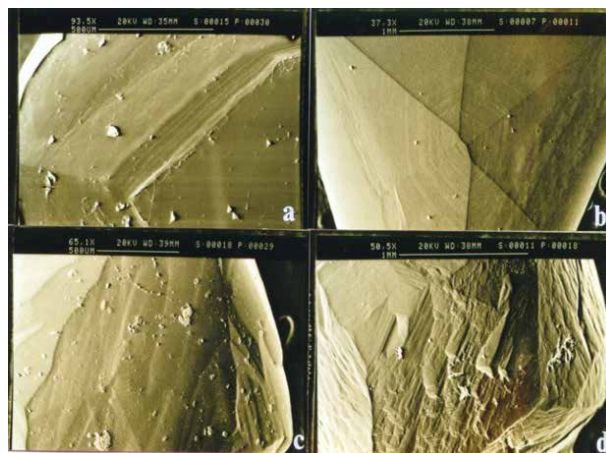


Figure 4. Examples of Thai diamonds with surface features under SEM showing (a) lamination lines, (b) shield-shaped laminae, (c) resorption etching and (d) hillock surfaces

Concluding Remarks

Thai diamonds have been found for about a century in southern Thailand. Their morphologies include mainly resorbed tetrahedroids (THH), but also rounded dodecahedrons, multiple twins, octahedrons and macles. Their surface features show straight laminae, shield-shaped lamination, cross-hatched lamination, pits, percussion scars and ruts. They are highly resorbed with glossy lustre suggesting more affinity with corrosive magma, i.e., related to lamproite or another minette rock type, more than to kimberlite (e.g., Kaminsky *et al.*, 2000). More details on their inclusions and possible geological source region need further research.

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