**Abstract**

**Raman and Photoluminescence Spectroscopy in Mineral Identification.** Kuehn, J. Wolfgang, Gemlab Research & Technology, Canadian Institute of Gemmology, P.O. Box 57010, Vancouver, B.C., V5K 5G6, CANADA, gemlab@cigem.ca

**Introduction:** Raman spectroscopy is particularly useful for rapid identification of minerals and gemstones. Searchable data-bases such as in the RRUFF project [1] contain a large number of reference spectra and are in public domain; the data-bases can be modified and expanded to the user’s requirements.

**Raman Applications:** Fig 1 demonstrates the quick identification of Zektzerite which is an extremely rare mineral; there are few gem quality specimen known and they could easily be mistaken for other minerals.

![Figure 1 Zektzerite confirmed with RRUFF database](image)

Gemologists often have difficulty to identify gemstones set in jewellery. Fig 2 shows how a Raman spectrometer can quickly separate gems with overlapping refractive indices (all in the range RI = 1.54 – 1.56).

![Figure 2 Gemstones with overlapping refractive indices](image)

The measurement of refractive index (RI) with a gemmological refractometer requires a flat polished surface; if the RI of a gemstone is above the limit of the refractometer other time consuming gem testing tools and methods have to be used. Fig 3 illustrates the quick separation of diamond from its most popular imitations.

![Figure 3 Separation of diamond from CZ and moissanite](image)

Garnet varieties cover the full colour spectrum and due to their similarity are difficult to identify; Raman based spectral measurements can give a precise and relatively inexpensive solution for these problems [2].

The detection of treatments in gemstones is often challenging; for example in Fig 4 the use of extended Raman spectroscopy can detect polymer treatment in jadeite.

![Figure 4 Raman spectrum of polymer treated jadeite](image)

**Photoluminescence (PL):** In some cases luminescence features are so intense that one can no longer see the Raman peaks associated with the atomic vibrations of the mineral. Luminescence spectroscopy is used to measure the energy levels of luminescence centers [3].

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PL Applications: The GL Gem Raman PL532 TEC is a dual purpose Raman spectrometer with an adjustable 532 nm laser source (200 mW) and is suitable for PL studies in the range from 530 – 750 nm. After reducing the laser output to approximately 30% of full power photoluminescence patterns become visible.

In Fig 5 a synthetic red flux grown spinel is quickly separated from natural red spinel; to arrive at the same conclusion using a microscope and other gem testing tools would have been more time consuming.

Figure 5 Separation of synth. from natural red spinel (685 nm)

In a similar manner emerald types (whether natural or synthetic, also geological provenance) can be determined by observing chromium PL peak locations.

Current Challenges: Synthetic materials, treatments, imitations and fakes are problems facing not only dealers, buyers and collectors but the entire jewellery industry. With the advent of small synthetic diamonds mixed in parcels with natural diamonds scientists are urgently working on methods and instrumentation to properly identify synthetic and treated diamonds.

One area of great interest are photoluminescence studies for diamond type screening and detecting HPHT treatments [4]. However, this method should be supported with other instrumentation and not be used alone.

Fig 6 illustrates a synthetic HPHT treated diamond with a nitrogen-vacancy defect at 637 nm; it also shows the suppression of the 558, 566 and 576nm bands which usually indicate an untreated diamond if the 637 nm peak is missing as well. Strong silicon-vacancy peak at about 737nm may indicate CVD (chemical vapour deposition) grown synthetic diamond.

Crystallographic defects causing the above mentioned PL peaks for natural untreated diamonds are not fully understood.

Figure 6 Characterization of synth. HPHT treated diamond

Conclusion: Several economically priced Raman spectrometer systems are now available and allow for rapid identification of gems and minerals. In general polished surfaces of gemstones produce good Raman scattering with high percentage matches in the RRUFF data-bases. For rough surfaces spectrometers with thermoelectric cooling (TEC) allow for low signal detection at exposure times of 1,000 ms and higher.

For the detection of synthetic diamonds and HPHT (high pressure high temperature) treatments the PL option of the Raman spectrometer is of particular value. With experience it may even provide a semi-quantitative estimate of trace elements such as chromium.

The analysis of Gr1 and SiV centres in CVD-grown yellow diamonds and the comparison with existing published data indicate that other impurities like Xe, Ni or Cr may produce other colours as well; more research is required. [5]

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