THE OXIDATION STATE OF EUROPIUM IN HYDROTHERMAL SCHEELITE:
IN SITU MEASUREMENT BY XANES SPECTROSCOPY

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ABSTRACT

X-ray microfluorescence and Eu_L3 micro-X-ray absorption near-edge structure (XANES) spectroscopy were conducted to determine the distribution and valence state of europium in two samples of scheelite (CaWO_4) containing between 40 and 350 ppm Eu. The samples come from hydrothermal gold deposits: the Archean Drysdale deposit in Kalgoorlie, Western Australia (DRY1) and the Caledonian, intrusion-related Stepnyak deposit in Kazakhstan (USP31). Laser-ablation ICP–MS analyses reveal that both samples are characterized by bell-shape chondrite-normalized REE patterns with a maximum enrichment around Eu, and a small positive or no Eu anomaly. USP31 displays a complex zoning between this pattern and a strongly LREE-enriched pattern with a strong positive Eu-anomaly. The XANES spectroscopy reveals the coexistence of Eu^{2+} and Eu^{3+} in both scheelite samples DRY1 and USP31. The oxidation of Eu in USP31 appears to be inhomogeneous within a single crystal, with zones containing a mixture of Eu^{2+} and Eu^{3+} and zones containing more than 90% Eu^{2+}. This study shows that it is possible to obtain semiquantitative element-distribution maps for LREE as well as, in optimal conditions, Eu_L3 XANES spectra on a bending magnet beamline equipped with a solid-state detector.

Keywords: X-ray absorption near-edge spectroscopy (XANES), scheelite, europium, oxidation state, rare-earth elements, elemental mapping, gold deposit.

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AFIN DE CARACTÉRISER LA DISTRIBUTION ET L’ÉTAT D’OXYDATION DE L’EUROPÉIUM DANS DEUX ÉCHANTILLONS DE SCHEELITE (CaWO4) CONTENANT ENTRE 40 ET 350 ppm Eu, NOUS AVONS AVOCÉVÉ DES ANALYSES PAR MICRO-FLUORESCENCE X ET PAR L’ÉTUDE DE LA STRUCTURE FINE DU SEUIL D’ABSORPTION L3 DE L’EUROPÉIUM OBTENU PAR MICRO-RAYONS X. LES ÉCHANTILLONS PROVIENNENT DE GISEMENTS AUFRÊRES HYDROTHERMAUX : LE GÎTE ARCHÉEN DE DRYSDALE À KALGOORLIE, AUSTRALIE OCCIDENTALE (DRY1), ET LE GÎTE CALÉDONIEN LIÉ À UNE INTRUSION DE STEPNYAK, AU KAZAKHSTAN (USP31). LES ANALYSES PAR ABLATION AU LASER – ICP–MS MONTRENT QUE LES DEUX ÉCHANTILLONS POSSÈDENT DES SPECTRES DE TERRES RARES (NORMALISÉS PAR RAPPORT AUX CHONDrites) EN CLOCHE AVEC UN ENRICHISSÉMENT MAXIMUM AU NIVEAU DE L’EU, ET UNE FAIBLE ANOMALIE POSITIVE OU AUCUNE ANOMALIE EN EU. USP31 POSSEDE UNE ZONATION COMPLEXE, LES DOMAINEs VARIANT DE CE SPECTRE À UN SPECTRE FORMENT UNE AUSSIE POSITIVE EN EU. LA SPECTROSCOPIE XANES RÉVÈLE LA COEXISTENCE D’EU2+ ET EU3+ DANS LES DEUX ÉCHANTILLONS. L’ÉTAT D’OXYDATION DE L’EU DANS USP31 SEMBLE ÊTRE INHOMOGÈNE AU SEIN D’UN MÊME CRISTAL, AVEC DES ZONES CONTENANT UN MÈNAGE DE EU2+ ET EU3+ ET DES ZONES CONTENANT PLUS DE 90% D’EU3+. CETTE ÉTUDE MONTRE QU’ELLE EST POSSIBLE D’OBTENIR DES CARTES DE DISTRIBUTION DES TERRES RARES LÉGÈRES ET, DANS DES CONDITIONS OPTIMALES, DES SPECTRES XANES EUL3 EN UTILISANT UNE LIGNE SUR AIDER DE COURBURE AINSI QU’UN DÉTECTEUR OPÉRANT À L’ÉTAT SOLIDE.

**Mots-clés** : structure fine du seuil d’absorption de rayons X (XANES), scheelite, europium, état d’oxydation, terres rares, distribution des éléments, gisements d’or.

**INTRODUCTION**

Tracking the oxidation state of ancient hydrothermal fluids is fundamental for our understanding of the physicochemical conditions of ore transport and deposition in many Cu, Au, Zn, Pb, or U deposits (e.g., Heinrich et al. 1996). Geologists rely mainly on mineral assemblages to estimate redox conditions in ancient hydrothermal systems. However, stability fields of minerals commonly cover large ranges in redox conditions. Moreover, in cases of complex polyphase mineralization, it can be difficult to relate one particular assemblage to a particular stage of mineralization. The oxidation state of trace elements incorporated in hydrothermal minerals may provide an additional tool to estimate the redox state of ancient fluids. In this paper, we explore the use of in situ X-ray absorption near-edge structure (XANES) microspectroscopy to measure the oxidation state of europium in scheelite from gold deposits at 7 × 7 µm² resolution, and show that the samples studied contain a mixture of Eu²⁺ and Eu³⁺.

**RARE-EARTH ELEMENTS AND SCHEELITE IN HYDROTHERMAL SYSTEMS**

Rare-earth elements (REE) occur as trivalent cations in nature, and behave in a consistent manner during geochemical processes such as melting, fractional crystallization, transport in a fluid, phase separation, and coprecipitation. Fractionation among the REE is caused by a slight systematic decrease in ionic radius from La to Lu, and by changes in oxidation states for some REE. In particular, Eu²⁺ exists under reducing conditions; its stability in aqueous solutions increases with increasing temperature (Sverjensky 1984, Haas et al. 1995).

The REE are well established as petrogenetic indicators in magmatic systems (e.g., Lipin & McKay 1989). Europium is used to estimate redox conditions in such systems; however, the ratio Eu²⁺/Eu³⁺ is rarely measured directly, and only the so-called europium anomaly, based on the variation in total Eu relative to neighboring REE, is used. The REE and the europium anomaly are also commonly used in the study of hydrothermal systems; however, their use has been hindered by the complexity of the controls on REE fractionation in such environments (Bau 1991, Gieré 1996). Direct measurement of the Eu²⁺/Eu³⁺ value in tracer minerals may provide a sensitive probe into redox conditions at the time of mineral growth, and would constrain the interpretation of the Eu anomaly in these minerals.

Scheelite (CaWO₄) is a common accessory mineral in various types of gold mineralization (e.g., Uspensky et al. 1998). Scheelite incorporates tens to thousands of ppm REE in substitution for Ca (e.g., Brugger et al. 2000b). The wide distribution of scheelite in Au-bearing systems and its close association with deposition of gold make it an excellent candidate as a tracer of the composition of the Au-bearing fluids and Au-deposition processes (Uspensky et al. 1998, Brugger et al. 2000a, b). Scheelite also is useful for dating ore deposition by means of the radioactive decay of ¹⁴⁹Sm to ¹⁴⁹Nd (half life 106 Ga; e.g., Anglin et al. 1996, Brugger et al. 2002, Eichhorn et al. 1997, Ghadiri et al. 1999).

**EU L₃ XANES SPECTROSCOPY OF MINERALS**

The XANES spectra at the L₃-edge of Eu can be used to determine the valence state of europium in the sample. Well-resolved edge-resonance peaks at 6973.7 and 6981.7 eV are attributed to electron transitions between the 2D₃/2 and the 5d states in Eu²⁺ and Eu³⁺, respectively (Rakovan et al. 2001, and references therein). However, the collection of Eu XANES data (Eu L₃-edge 6977 eV, EuL₅1 5849.5 eV) on natural minerals is made difficult by interferences by manganese (K-edge 6539 eV, Kα₁ 5900.3 eV) and neodymium (L₃-edge 6208 eV; Lα₁ 5227.6 eV, Lβ₁ 5721.1 eV, Lβ₂ 5828.1 eV), which cannot be resolved...
with solid-state detectors (energy resolution \(\sim\) 150 eV). Rakovan et al. (2001) established the oxidation state of Eu in a powdered sample of fluorapatite from Llallagua (Bolivia) containing about 700 ppm Eu in the presence of high concentrations of Mn and Nd, using a high-resolution wavelength-dispersive spectrometer (WDS) to resolve the interferences. They also benefitted from higher fluxes at the sample, in the order of \(10^{10}\) photons per second, by using an insertion-device beamline. For the present experiment, we wanted to test whether a less demanding analytical set-up, including a solid-state spectrometer (where all elements of interest can be investigated at the same time, as opposed to the WDS detector, in which the analyzing crystal and detector have to be scanned to measure a range of wavelengths) and a bending magnet beamline would be sufficient to obtain useful information on a natural mineral such as scheelite. Compared to fluorapatite, scheelite has the advantage that it contains very little manganese, in most cases, less than 20 ppm (Table 1).

### TABLE 1. RESULTS OF LASER-ABLATION ICP-MS ANALYSES OF SCHEELITE FROM STEPNYAK AND DRYSDALE

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**DRY1, Drysdale Stockwork, Kalgoorlie, Western Australia**

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Results are expressed in ppm.
an Ar–Ar age of 2602 ± 8 m.y., interpreted as the age of the mineralization (Kent & McDougall 1995). Desulfidation of an \( \text{H}_2\text{S} \)-rich fluid (molality \(< 10^{-5}\) ) by reaction with wallrock Fe-rich silicates and carbonates contributed to the precipitation of gold from aqueous bisulfide complexes (Mikucki & Heinrich 1993, Mernagh et al. 2004). The relatively high grade of the Mt. Charlotte stockwork is interpreted as resulting from unusually sharp temperature-gradients (\(< 220 \) to 300°C) developed (2004). The relatively high grade of the Mt. Charlotte.

Sample USP31 consists of scheelite in a quartz–calcite matrix from the Stepnyak deposit, northern Kazakhstan, which has produced \(< 25 \) t of gold (Spiridonov 1995). The ores at Stepnyak are associated with a Caledonian intrusive complex (diorite, tonalite, granodiorite). The ore-forming fluids were characterized by \( T \approx 370°C, P \approx 3.6 \) kbar, salinity \(< 17\% \) NaCl equivalent, \( \text{CO}_2 \approx 7 \) mol/kg, \( \text{CH}_4 \approx 2.5 \) mol/kg, \( \text{P} (\text{H}_2\text{S}) \approx 2.5 \) kbar; the mineralization shows a strong depth-controlled zoning, from shallow gold ± Ag-sulfosalts ± antimonite and gold – galena – sulfantimonides to gold – galena – tellurides and gold – tellurides in the abyssal zone (pressure from 2 to 3.6 kbar; Spiridonov 1995).

USP31 and DRY1 are vein fragments containing millimetric crystals of scheelite. The samples were embedded in epoxy resin (25.4 mm diameter, 1 mm thick) and freshly polished to 1 \( \mu \)m flatness.

**Experimental Methods**

Laser-ablation inductively coupled plasma – mass spectrometry (LA–ICP–MS) analyses were performed with a NewWave 266 nm UV laser (beam \(< 30 \) \( \mu \)m in diameter) and a Finnigan (ThermoQuest) MAT ELEMENT ICP–MS located at Monash University. Analytical details are provided in Table 1 of Brugger et al. (2000).

The XANES spectra were collected at beamline 2–BM–B at the Advanced Photon Source (APS) at Argonne National Laboratory, USA. Beamline 2–BM–B has a Kohzu double Si (111) monochromator, resulting in an energy resolution \( E/\Delta E \approx 1.4 \times 10^{-4} \). Two Kirkpatrick–Baez (KB) mirrors at an incident angle of 4 mrad suppressed the third-order reflection from the Si(111) monochromator down to 0.3%. The KB mirrors are contained in a He-filled box with 0.5 mm Kapton window in order to reduce the air-induced scatter and prevent surface oxidation (Eng et al. 1998). The KB mirrors focused the beam to \( 7 \times 7 \) \( \mu \)m² at the sample (sitting in air). Two ionization chambers placed before the sample were used to monitor the intensity of the incident beam. A single-channel Si drift diode (SDD) detector (Iwanczik et al. 1996), fitted with a collimator, was located at \(< 90° \) to minimize detection of the scattered X-ray beam. The XANES spectra were measured by scanning the beam energy over the Eu \( L_3 \) absorption edge (6977 eV) while collecting counts in a region of interest centered around the Eu \( L_3 \) peak fluorescence line (5849.5 eV). The scans were performed from 6.9 to 7.1 keV and from 6.96 to 7.0 keV in steps of 0.5 and 0.3 eV, respectively, with counting times ranging from 8 to 30 s per step. At these energies, the incident flux was between \( 10^6 \) and \( 10^7 \) photons/s at the sample. The monochromator energy was calibrated by using synthetic scheelite doped with \(< 6 \) wt% Eu³⁺, synthesized in a NaCl melt according to the method described by Gaft et al. (1999). The XANES spectrum of this sample is similar to that obtained on synthetic Eu₂O₃. The position of the Eu³⁺ edge was obtained by fitting the XANES spectrum with a pseudo-Voigtian peak, a linear background, and an arc-tangent step function (Fig. 1a); a shift of \(< 1.27 \) eV was applied to bring the peak position to the literature value of 6981.7 eV (see below).

Fluorescence images were collected at both 2–BM–B (10.0 keV) and 2–ID–E (9.9 keV) for 6.3 hours and 3.5 hours, respectively. The excitation energy was chosen to be below the W \( L_2 \) edge (10.207 keV) in order to minimize background in the region of the spectrum that hosts the REE. Beamline 2–ID–E is a side-branch beamline that uses a silicon crystal beam-splitter as a single-bounce monochromator. For these experiments, a zone plate 320 \( \mu \)m in diameter with an outermost zone-width of 100 nm was used (focal length of 24.75 cm at 9.9 keV in the first-order diffraction), resulting in a focused beam-spot of \( 0.7 \times 0.4 \) \( \mu \)m². The sample chamber was flushed with He gas mixture to reduce absorption of X-ray fluorescence and to decrease the background fluorescence from Ar in air. An order-sorting aperture intercepted higher-order diffraction maxima and, in conjunction with a central stop, blocked the unfocused beam. A three-element Ge energy-dispersion detector (Canberra) was located at \(< 90° \) to minimize detection of the scattered beam and fitted with a conical collimator to suppress scattered X-rays and background fluorescence not coming from the sample.

The fluorescence images and spectra collected on USP31 were analyzed using GeoPIXE II (Ryan 2000). GeoPIXE II, as the name suggests, was originally developed to analyze proton-induced X-ray emission (PIXE) data and uses the Dynamic Analysis (DA) method to project quantitative images of element distribution. The DA technique is a matrix-transform algorithm that unfolds overlaps and subtracts background, escape peaks and other detector artifacts. GeoPIXE II has been extended for use with energy-dispersion synchrotron X-ray fluorescence (SXRF) data (Ryan et al. 2005). To make the transition to SXRF, it was necessary to modify the program to calculate accurate theoretical X-ray yields generated by incoming photons for a given composition of sample, experimental geometry and beam energy. X-ray yields and relative intensities reflect sub-shell ionization cross-sections, Coster–Kronig transition rates for the L shell, and fluorescence yields. Sub-shell absorption cross-sections use...
RESULTS

The fluorescence spectra collected at 2–BM–B and 2–ID–E show the presence of the rare earths, with the 2–ID–E spectra featuring a somewhat higher energy-resolution than 2–BM–B (Fig. 2). In addition to elastic (10 keV) and Compton scattered radiation, the spectra show a prominent feature around 8 keV, which arises from resonant Raman scattering on the major element W in the sample (e.g., Briand et al. 1981, Surić et al. 1991). This area of the spectrum and the peaks due to the scattered beam were excluded in fitting the spectrum in GeoPIXE. Calcium was used as an internal standard to quantify the flux (stoichiometric value of 13.9 wt%). The following elements were included in the fitting procedure: Si, Ar, Ca, Mo, La, Ce, Pr, Nd, Sm, Eu, Gd, Dy and Er. Apart from La and Ce, all REE overlap to some extent, making the quantitative determination of REE in natural samples delicate, even with the use of high-resolution WDS spectrometers (Williams 1996). As can be seen in Figure 3, the concentrations obtained by the GeoPIXE spectral fitting are of the right order of magnitude, and the changes observed in the quantitative DA maps appear to be real. The errors provided by GeoPIXE for the concentrations of the individual REE arise from counting statistics only and are estimated from the error matrix generated in the fit using the method of Bevington (1969), and scaled by the square root of reduced chi-squared of the major line for each element to account for residual systematic deficiencies in the fitting model (Ryan et al. 1990). They vary between 10 and 20% (1σ); these values are an underestimation of the error for the different REE, owing to the strong overlap among the different REE and the resulting colinearity among the concentrations of the different REE. The GeoPIXE II software hence enables semiquantitative analysis of the samples for the REE, despite the low concentrations of the REE and the low spectral resolution. This capability is important for selecting areas in the sample with different REE compositions for XANES analysis.

Sample USP31 has a very inhomogeneous distribution of the REE, with chondrite-normalized patterns

Fig. 1. Europium L\(_3\)-edge XANES data for (a) synthetic scheelite doped with Eu\(^{3+}\), (b–d) three different locations on sample USP31 and (e) scheelite DRY1. The fits were obtained using two pseudo-Voigtian peaks (50% Gaussian and 50% Lorentzian contribution, a value obtained from the synthetic scheelite doped with Eu\(^{3+}\) centered at 6973.7 (Eu\(^{2+}\)) and 6981.7 (Eu\(^{3+}\)) eV, a linear background and an arc tangent step-function.
plotting between two end-member compositions (Fig. 3): very steep enrichment in the light REE (LREE; La to Eu), with a highly positive Eu anomaly [anal. (1) in Fig. 3] and enrichment in medium REE-enriched pattern (MREE; Eu to Lu) with a maximum enrichment at Eu with a small positive Eu anomaly [e.g., anal. (2)] or none at all [anal. (3) in Fig. 3]. The absolute concentrations can also vary by almost an order of magnitude [e.g., anal. (2) and (3) in Fig. 3]. The geometry of the zoning cannot be understood from the LA–ICP–MS
profile. The elemental maps (Fig. 4) reveal a patchy pattern of zoning, with regions that can be related to the REE patterns measured by LA–ICP–MS (cf. Figs. 3, 4): (1) high Ce and Nd, low Eu, (2) high Ce and Eu, low Nd, and (3) low Ce, Nd and Eu.

DRY1, on the other hand, shows only one type of REE pattern, enriched in the MREE with a maximum between Eu and Gd, and a small positive Eu anomaly or none at all. The total concentration of the REE varies over almost an order of magnitude (Table 1).

The XANES data obtained for the natural minerals are noisy, a result of the low concentration of Eu and a relatively low incident flux (Figs. 1b–e). In addition, the NdLβ interference results in peak-to-background ratios not exceeding ~2. Among the many samples of scheelite analyzed by LA–ICP–MS, USP31 and DRY1 are characterized by unusually high Eu/Dy values. This anomaly results in high Eu concentration and relatively reduced Nd interference, and only these two samples were suitable for measurement with the apparatus used in the present study. Two XANES spectra collected on different regions of USP31 (Figs. 1b, d) are very similar. The first peak clearly appears at the position of the Eu²⁺ resonance-peak. As the intensity of the Eu³⁺ edge resonance-peak is 1 to 1.5 times larger than that
of Eu\textsuperscript{2+} (Rakovsk et al. 2001), most Eu exists in the 2+ oxidation state at these two spots. However, there is a clear shoulder at the position of the Eu\textsuperscript{3+} resonance-peak, indicating the presence of significant (>10%) amounts of Eu\textsuperscript{3+} as well. Other regions seem to lack this shoulder (Fig. 1c), and hence contain in excess of 90% Eu\textsuperscript{2+}. Sample DRY1 shows a XANES spectrum relatively similar to that of point E on USP31 (Figs. 1b, c), indicating a predominance of Eu\textsuperscript{2+} coexisting with significant amounts of Eu\textsuperscript{3+}.

**DISCUSSION**

The study of REE distribution and speciation in hydrothermal minerals is made difficult by low concentrations, spectral overlaps and complex zoning at the scale of 1 to 10 \(\mu\)m. However, with synchrotron-based XRF and XANES techniques, it is possible to obtain semiquantitative information about the concentration, pattern of zoning, and oxidation state of REE in situ in hydrothermal scheelite. In the two samples studied, Eu exists in a mixed oxidation state (Eu\textsuperscript{2+} and Eu\textsuperscript{3+}). Similar evidence for mixed-valence Eu was obtained by Rakovan et al. (2001) for a sample of powdered hydrothermal fluorapatite, suggesting that hydrothermal minerals commonly contain a mixture of Eu\textsuperscript{2+} and Eu\textsuperscript{3+}. The apparently inhomogeneous distribution of Eu\textsuperscript{2+} and Eu\textsuperscript{3+} in USP31 suggests that Eu\textsuperscript{2+}/Eu\textsuperscript{3+} may be preserved in scheelite over hundreds of millions of years, and that diffusion does not affect this ratio severely, at least for the conditions prevailing during the geological history of USP31.

This study hence paves the way for systematic studies of the oxidation state of Eu in hydrothermal minerals. Quantitative data on a large variety of samples can be obtained by using an insertion-device beamline (higher flux) and a WDS spectrometer to minimize the Nd\textsubscript{1/2}B\textsubscript{1/2} interference. Such studies will help in the interpretation of REE data in complex hydrothermal environments. For example, the range of REE distribution in scheelite from Archaean gold deposits has been interpreted both in terms of open-system change in the oxidation state of the ore-forming fluids (Ghaderi et al. 1999), and in terms of closed-system fractional crystallization (Brugger et al. 2000a, b, 2002). These two contrasting interpretations could be discriminated by measuring the ratio Eu\textsuperscript{2+}/Eu\textsuperscript{3+} in the finely (1–10 \(\mu\)m) zoned grains of scheelite.

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