Research Article

Fractionation of Rare Earth Elements in Greisen and Hydrothermal Veins Related to A-Type Magmatism

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1. Introduction

The growing demand for rare earth elements (REE) in modern techniques for a sustainable environment has been associated with extensive prospecting for these metals worldwide [1]. Numerous laboratory experiments and genetic models have incorporated fluid-inclusion and isotopic data and/or trace element partitioning calculations to understand how REE are transported, fractionated, and ultimately deposited in ores typically associated to the formation or alteration of carbonatitic and alkaline to peralkaline rocks [2–6]. REE enrichment in these systems is often ascribed to magmatic processes that can concentrate REE in fluoride liquids over silicate melt through liquid immiscibility [7] or in primary REE-rich minerals such as phosphates and zircon crystallizing from magma [8]. In late-magmatic stages, hydrothermal...
mobilization of REE transported via bonding with ligands such as F\textsuperscript{-}, CO\textsubscript{3}\textsuperscript{2-}, Cl\textsuperscript{-}, or OH\textsuperscript{-} can form secondary vein mineralizations from emanating fluids [9]. Fluid-rock interactions may affect REE distribution along the fluid pathway, where the availability of REE-complexing agents and the solubility of REE precipitation products will control REE fractionation through retention and precipitation [10]. REE fluorides, carbonates, and hydroxides are strong complexes that can under certain conditions become unstable and thus act as depositional complexes with limited capability to carry REE [5]. In contrast, REE-chloride complexation, although weaker, can in the absence of insoluble REE-chloride minerals enable extensive REE transportation in many hydrothermal systems [11]. The preferred complexation of LREE by chloride and fluoride and HREE by hydroxide and carbonate ligands experimentally established at low temperatures (generally below 100°C or even at ambient conditions) is not as straightforward at higher temperatures, where pH and redox conditions and coexisting species in the solution have stronger impacts on REE selectivity [5]. Experimental data of REE-carbonate and -hydroxide complexation at elevated temperatures is particularly scarce, although recent simulations predict abundant transportation of REE carbonates in solutions with high CO\textsubscript{2} and low salinity [12]. Fluid-rock and fluid-fluid interactions add further complexity to REE fractionation in natural systems, yet the applicability of experimentally determined partitioning has been implemented in geochemical, physical, and thermal evolution models of magmatic-hydrothermal systems at Strange Lake granitic pluton, Canada [3, 13] and in carbonatites such as those at Bayan Obo, China [14] and Songwe Hill, Malawi [15]. The large discrepancies in the distribution and fractionation of REE into minerals including calcite, fluorite, and apatite demonstrated in these studies appear to outline a virtually site-specific combination of transport and precipitation mechanisms affecting REE deposition. This complexity can be resolved by integrating the components involved at each stage throughout the evolution of the hydrothermal system [10]. We adopt this approach by scrutinizing fluid-inclusion data along with REE abundance and patterns of whole-rock and mineral precipitates from the magmatic source rock and the host-rock reacting with the migrating fluids, as well as from the resulting greisen veins (hereafter named greisen) and distal veins (hereafter named veins) at a site on the Baltic Shield (Laxemar, SE Sweden). The overall aim is to assess mobilization, transportation, fractionation, and ultimate deposition of REE in the hydrothermal system. The temporal relationship between the intrusion and veins is investigated by a newly developed high spatial resolution Rb-Sr dating technique [16], applied for the first time on hydrothermal veins and greisen mineralization.

2. Geological Setting

At the study site (Laxemar, SE Sweden), the country rock is part of the Transscandinavian Igneous Belt (TIB) and is composed of a suite of approximately 1.80 Ga intrusive rocks dominated by granodiorite and quartz-monzodiorite with close temporal and genetic relationships (Figure 1). The TIB formed during several pulses of magmatism between the Svecofokarelian terrains to the north and the younger SW Scandinavian domain in the west [17]. Two alkali- and fluorine-rich granite plutons, the Götemar and Uthammar granites, crop out in the area (Figure 1). These are most likely syntectonic intrusions related to far-field effects of the 1.47 to 1.44 Ga Danopolitan compression event [18, 19], as shown by an overlapping U-Pb zircon age (1433 ± 10 Ma) for the Götemar granite [20]. They are lacholphillshaped and have greater lateral extent in the subsurface than at the ground surface, and are characterized by a narrowing root zone or feeder reaching a depth of approximately 4 km (Figure 1(c)) [21]. The Götemar granite is a horizontally zoned tabular structure with a diameter of 5 km, formed at an estimated crustal depth of 4–8 km [22]. It is geochemically classified as a metaluminous to peraluminous anorogenic granite [23] with high FeO/MgO and Na\textsubscript{2}O+K\textsubscript{2}O/CaO ratios. It is dominated by quartz and feldspars (up to 80 vol.%, where the proportion of K-feldspar:alkali plagioclase≈1:1), and contains muscovite in minor amounts (<5%) [24, 25]. Greisen mineralization constituting cm-thick fluoride-rich quartz-muscovite veins was formed in the country rock at a late stage of the Götemar granite intrusion [26, 27].

Although no economic REE mineralizations occur in the area, a large number of vein generations have formed throughout the geological history of the TIB rocks. The oldest generation is related to the waning stages of the Svecofokarelian orogeny (1750–1620 Ma) and the youngest to low-temperature conditions in the Quaternary [19, 28, 29]. Veins related to the granite plutons (1433 ± 10 Ma), in focus here, are identified by (i) the occurrence of epidote, (ii) nonbiogenic δ\textsuperscript{13}C values of the calcite, that is, in the -7 to -2‰ V-PDB range, (iii) relatively low δ\textsuperscript{18}O (~25 to -10‰ V-PDB) and 87Sr/\textsuperscript{86}Sr values (0.705–0.710) of the calcites [26], and (iv) hydrothermally red-stained wall rock [30] that has secondary mica \textsuperscript{40}Ar/\textsuperscript{39}Ar ages overlapping with greisen mica and the granite [27]. Furthermore, several vein generations were formed during the Palaeozoic, and multiple episodes of ancient microbial activity in the bedrock fractures have been recorded from the Paleozone to the Quaternary [28, 29, 31]. Although the general decrease in temperature over time has had a profound effect on the mineral assemblage of the veins, several minerals, such as calcite, have been formed under both high- and low-temperature conditions and thus exist in most generations [27]. This phenomenon and the spatial restriction of the borehole diameter (approximately 5 cm) mean that it is not straightforward to assign a certain vein to a specific geological event.

3. Materials and Methods

3.1. Sampling. The Site Characterization Database (SICADA) owned by the Swedish Nuclear Fuel and Waste Management Co. (SKB) contains information on 50 cored boreholes with a total length of approximately 25 km in the Paleoproterozoic crystalline bedrock of the Laxemar area, SE Sweden. These boreholes, and the associated SICADA database, were made as part of previous, but now abandoned, investigations for a nuclear waste repository. This database was used
for the identification of granite-associated veins in the Paleoproterozoic bedrock (TIB rocks). The diagnostic features included the presence of the index mineral epidote (Figures 2(a)–2(d) and 2(f)) (± quartz, calcite, chlorite, and fluorite), the presence of alteration haloes (hydrothermally red-stained wall rock), and characteristic stable isotope ratios of calcite (existing information for a few veins) that distinguish them from abundantly occurring younger veins [26]. The relative proportions of the vein minerals were estimated based on macroscopic and microscopic observations, including photo documentation and available petrographic description of specimens being part of a previously published dataset \((n = 26; [26])\). A compilation of samples and the analytical methods applied to each vein sample in this study is provided in Table 1.

Greisen was, in a manner similar to the veins, identified in the SICADA database and sampled from the drill cores. Identification was based on its notable quartz-mica composition (± fluorite, topaz, and sulphides; Figure 2(e)) and coarse-grained appearance (Figures 2(g) and 2(h)) and typical associated wall-rock sericitization-silicification with diffuse contact to the wall rock. Samples of quartz-monzodioritic to granodioritic rocks belonging to the TIB and coarse to medium-grained granite, fine-grained granite, and aplite dykes belonging to the Götemar granite were taken from fresh rock avoiding zones that were red-stained or otherwise altered. A compilation of samples and the analytical methods applied to each greisen, Götemar granite, and TIB rock sample in this study is provided in Table 2.
3.2. Chemical Analyses. Thick sections of veins, greisen, TIB rocks, and Götemar granite were polished and carbon coated, and the individual minerals were analysed by Scanning Electron Microscopy (SEM) and Energy Dispersive X-ray Spectroscopy (EDS) on a Hitachi S-3400N (Department of Earth Sciences, University of Gothenburg, Sweden; hereafter GU) in order to verify pure mineral phases and obtain composition stoichiometry. The acceleration voltage was 20 kV, and specimen current was 1 nA. Oxide calibration standards were linked to a cobalt drift standard and a stable specimen current of 1 nA. X-ray spectrometric corrections were made by an INCA computer system. Spot size was ~5 μm, and the detection limit was 0.1 oxide %. After the SEM-EDS analyses, the identified minerals were analysed for major and trace elements by laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) using either a LSX-200 Cetax or a NWR213 UV laser system followed by analyses on an Agilent 7500 or an Agilent 8800. The LSX-200 laser was run with 4 mJ power at 4 Hz and a 200 μm spot size. The raw counts were reduced using in-house software (LaserCalc). NWR213 was run with 0.355 to 1.015 mJ power at 5–10 Hz and a 10–100 μm spot size. Data processing was carried out using Glitter software. Calibration standard (NIST-612 glass), secondary standard of homogeneous greisen fluorite (only for fluorite), and internal standards (44Ca for calcite and fluorite and 29Si for silicate.

Figure 2: Representative vein (a–d) and greisen (e–h) specimens. (a) Photograph of sealed epidote-fluorite vein; (b) BSE image of epidote and chlorite lining a vein with calcite in the centre and in a veinlet penetrating into the wall rock; (c) photograph of a reactivated (presently open) vein with epidote closest to the wall rock and calcite precipitated after epidote; (d) photograph of a sealed epidote-calcite-fluorite vein; (e) BSE image of topaz intergrown with muscovite, fluorite, and quartz in greisen; (f) photograph of greisen containing epidote, biotite, and calcite; (g) photograph of fluorite-rich greisen with muscovite and quartz; (h) photograph of fluorite-poor greisen composed of quartz, muscovite, and low amounts of fluorite.
minerals, where mineral-specific compositions were determined by SEM-EDS of the same crystals) were used for both laser runs. Accuracy was controlled by analyses of basalt glass standard (JB-1 and BCR-2G) and was generally better than 10%.

Handpicked specimens of calcite and fluorite (cleaned by removing other minerals attached to the grains) of veins and greisen were analysed for Mg, Ca, Mn, Fe, Rb, Sr, Y, and La-Lu by inductively coupled plasma (ICP) mass spectrometry (MS) on an Agilent 7500 ICP-MS at GU. 12 mg of each calcite specimen was dissolved in 47 ml 5% HNO₃ for 1 h, whereas 50–100 mg of fluorite was dissolved in a mixture of 3 ml HNO₃(conc.) and 9 ml HCl(conc.) overnight at 90°C. The dried residue was treated with a mixture of 7.5 ml HNO₃(conc.) and 5 ml H₂O overnight at 90°C. Standards of dolomite (JDo-1), granite (SARM 48), and basalt (W-1) were included. Multielement standard solutions (Merck, VI Certipur, and Agilent diluted to 0.1, 1, 10, and 100 ppb) were used to construct linear calibration graphs. A solution containing equal amounts (800 ppb) of In and Re was used as an internal standard. Precision was generally better than 5%, and accuracy was better than 10%.

### Table 1: Summary of vein samples and analytical techniques applied on them.

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bt=biotite, cc=calcite, chl=chlorite, epi=epidote, fl=fluorite, qz=quartz, and py=pyrite. ICP=inductively coupled plasma mass spectrometer. LA = laser ablation inductively coupled plasma mass spectrometer.
Bulk specimens of greisen ($n = 5$) and Götemar granite ($n = 4$ coarse/medium grained, $n = 3$ fine grained, $n = 4$ aplite) were analysed for concentrations of major elements Sr and REE using ICP Atomic Emission Spectrometry (AES) and ICP Quadrupole Mass Spectrometry (QMS) at ALS Scandinavia AB, Luleå, Sweden. The analyses were made
according to EPA-approved procedures documented in EPA methods 200.7 (for ICP-AES) and EPA methods 200.8 (for ICP-QMS). Each granite and greisen specimen was ground, fused with LiBO$_2$, and finally dissolved in 5% HNO$_3$ prior to determination. The accuracy for all metals was generally better than 10%.

3.3. Isotope Analyses. Isotope ratios of carbon ($^{13}$C/$^{12}$C) and oxygen ($^{18}$O/$^{16}$O) for 11 handpicked calcite vein specimens were determined at the Scottish Universities Environmental Research Centre (SUERC). Analysis of 1–2 mg calcite was performed on CO$_2$ released by the phosphoric acid dissolution method [33] in an automated preparation system that includes cryogenic purification of the gas. The pure gas was introduced into a dual-inlet, triple-collector gas source isotope ratio mass spectrometer (Micromass SIRA 10). Calibration of the mass spectrometer standard gas was via NBS 19. Precision was ±0.1‰ at 1σ. Oxygen isotope ratios were derived from the CO$_2$-measured values using a phosphoric acid fractionation factor derived from the literature [34]. The results for carbon and oxygen isotopes are reported as ‰ relative to V-PDB.

 Sulphur isotope analyses of two handpicked pyrite vein specimens were carried out at the Swedish Museum of Natural History, Stockholm. SO$_2$ was released from each pyrite specimen (0.1–20 mg) following a method described elsewhere [35], whereby samples were combusted at 1070°C for 25 min in the presence of excess Cr$_2$O$_3$. SO$_2$ was separated from excess oxygen and other combustion products by standard vacuum line techniques, and the purified gas was analysed for $^{34}$S/$^{32}$S on an isotope ratio mass spectrometer (Finnigan MAT DELTA Plus). Results are reported as ‰ relative to VCDT. Reproducibility was better than ±0.2‰ at 1σ.

Strontium isotope analyses on three handpicked fluorite specimens were carried out at the Institute of Energy Technology, Norway by thermal ionization mass spectrometry (Finnigan MAT 261). Each ground fluorite specimen was reacted with 1–2 M HNO$_3$ at 50°C for 48 hours. The acquired solution was centrifuged, separated from the residue, evaporated, dissolved in 200 μl 3 M HNO$_3$, and then loaded onto an ion-exchange column packed with Sr-spec Eichrom resin. Unwanted elements were washed out with 3 M HNO$_3$, and Sr was eluted with water. The eluate was evaporated and loaded on degassed Re filaments. Calibration was performed via NIST/NBS 987 standards. Precision was ±20 ppm. The total blank of Sr was 50–100 pg.

3.4. Fluid Inclusions. Quartz in one greisen and five vein samples were prepared as doubly polished, 150 μm thick, fluid-inclusion wafers. Microthermometric analyses were undertaken at the Department of Geological Sciences, Stockholm University on a Linkam THMS600 heating-freezing stage mounted on a Nikon microscope. The precision of the measurements was ±0.1°C for temperatures below +40°C and ±0.5°C above +40°C. Solid phases or gaseous fluid components are identified by using a Horiba Jobin-Yvon LabRAM 800 HR Raman spectrometer and an Olympus BX41 petrographic microscope. The analyses were made using 514 nm excitation of an Ar-ion laser adjusted to an on-sample intensity of 5 mW, with a laser beam focused to a spot of 1 μm. Raman acquisition was performed with a spectral range of 150–4200 cm$^{-1}$ and a spectral resolution of about 0.3 cm$^{-1}$, using a 1800 l/mm grating and an air-cooled (-70°C) CCD array detector.

3.5. In Situ Rb-Sr Dating. Three samples of cm-thick greisen consisting of coarse-grained (up to 1.5 cm in diameter) muscovite, biotite, K-feldspar, quartz, and fluorite along with accessory calcite, chlorite, pyrite, and topaz and two samples of 2–3 cm thick veins comprising up to 0.5 cm sized biotite + calcite + epidote were dated by in situ Rb-Sr LA-ICP-MS analysis. Isochrons were achieved by combining muscovite or biotite with K-feldspar, calcite, fluorite, and/or epidote. Rb-Sr analyses were performed at the microgeochemistry lab at GU using an ESI 213NWR laser ablation system connected to an Agilent 8800QQQ ICP-MS with an ORS3 (octopole reaction system) reaction cell installed between two quadrupoles. Following laser warm-up, ablation occurred with a static spot mode in a constant He flow (800 ml/min). The ablated material was mixed with N$_2$ and Ar before entering the ICP-MS torch region, where it reacted with N$_2$O or O$_2$ gas in the reaction cell. This reaction chemically separated $^{87}$Rb from $^{87}$Sr and thereby enabled calculation of $^{87}$Rb/$^{86}$Sr and $^{87}$Sr/$^{86}$Sr ratios. N$_2$O, SF$_6$ [36], or O$_2$ [16] were utilized as reaction gas in separate sessions. The raw $^{87}$Rb/$^{86}$Sr and $^{87}$Sr/$^{86}$Sr ratios were converted by correction factors derived from repeated analysis of standards NIST SRM 610 and mica-Mg-A1. $^{87}$Rb/$^{86}$Sr normalization was performed by using mica-Mg, a reference mineral powder of a phlogopite separated from Bekily, Madagascar [37], pressed to a nanopellet tablet [36]. Sample data reduction and within-run error calculation of selected element and isotopic ratios were performed using an in-house spreadsheet. No error propagation from uncertainties of literature data or within-run errors of standards was applied to sample errors.

4. Results

4.1. Mineralogy and Chemistry of Bedrock. The TIB granitoid rocks comprise quartz and feldspars (up to 60–65 vol.% proportion of K-feldspar: plagioclase=1:3) with varying amounts of chlorite, biotite, pyroxene, amphibole, and accessory titanite,apatite, and magnetite. The total REE concentrations in these rocks are on average 173 ppm, with a strong enrichment of LREE relative to chondrite (Figure 3(d)). The eight minerals analysed for chemical elements show a large variability both in terms of REE concentrations and fractionation patterns as well as Eu anomalies (Figures 4(c)–4(f)), with the highest concentrations in titanite (mean$\Sigma$REE = 18114 ppm) and apatite (mean$\Sigma$REE = 6307 ppm). The concentrations of the other metals are given for all bulk rocks in Appendix A and for the individual minerals in Appendix B.

The Götemar granite contains rock-forming quartz, K-feldspar, and plagioclase and accessory muscovite, fluorite, monazite, zircon,apatite, and rutile. In this rock, the REE occur in concentrations (mean$\Sigma$REE = 320) that are higher than in the TIB rocks. The coarse-grained granite variety
has strong enrichment in light relative to heavy REE (Figure 3(a)) [38, 39], whereas the fine-grained granite and the aplite dykes have downward concave to relatively flat patterns (Figure 3(b)) [39]. Of the six minerals analysed for chemical elements, the REE concentrations are low in muscovite and K-feldspar (mean $\sum$REE of 1.5 and 12 ppm, respectively), but they are typically orders of magnitude higher in fluorite ($\sum$REE = 2156 ppm), apatite ($\sum$REE = 14080 ppm), monazite ($\sum$REE = 45 wt %), and zircon ($\sum$REE = 3187 ppm, Figures 4(a) and 4(b)). Among the minerals, there is also a large variability in chondrite-normalized Eu anomalies and fractionation patterns, especially in the main accessory minerals that show patterns ranging from strong LREE enrichment in monazite to strong HREE enrichment in zircon, whereas fluorite and apatite display relatively flat patterns (Figures 4(a) and 4(b)).

4.2. Mineralogy and Chemistry of Greisen. The greisen is composed of quartz, muscovite, and fluorite. Minor minerals include biotite, K-feldspar, albite, chlorite, epidote, and pyrite. The main accessory mineral is calcite, whereas there is sporadic occurrence of topaz and fine-grained halite, sylvinite, barite, hematite, rutile, monazite, zircon, chalcopyrite, and sphalerite. Bulk greisen has low REE concentrations (mean = 75 ppm) when compared to the Götemar granite and the TIB rocks, and it has chondrite-normalized patterns that are enriched in LREE (Figure 3(c)). On a mineralogical level, epidote, chlorite, topaz, K-feldspar, and muscovite are all LREE-enriched but with highly variable concentrations with the highest $\sum$REE = 1120 ppm) in epidote (Figures 5(a) and 5(b)). These minerals, except chlorite, also have positive Eu anomalies (Figures 5(a) and 5(b)). Calcite has relatively flat REE fractionation patterns with a
Figure 4: Chondrite-normalized [40] REE concentrations of minerals in (a and b) the Götemar granite and (c–f) the rocks of the Transscandinavian Igneous Belt (TIB).
Figure 5: Chondrite-normalized [40] REE concentrations of minerals in (a–c) greisen and (d–f) veins.
weak MREE depletion, whereas fluorite shows strong progressive enrichment with atomic number throughout the REE series (Figure 5(c)). Both calcite and fluorite are relatively rich in REE (median $\sum$REE = 190 and 210 ppm, respectively, Appendix B).

4.3. Mineralogy and Chemistry of Veins. The mineralogical composition of the veins is heterogeneous, and not all minerals in the assemblage are present in each specimen. Taking these features into account, it is possible to define that the vein paragenesis consists, in decreasing order, of quartz (36%), calcite (23%), epidote (18%), chlorite (16%), pyrite (4%), and fluorite (3%). There is a large variability in the concentrations and chondrite-normalized fractionation patterns of REE among the minerals. Median $\sum$REE concentrations are highest in epidote (61 ppm, range = 30-124) and lowest in chlorite and fluorite (1.5, range = 0.74-30 and 2.8, range = 0.65-2.3, respectively), whereas calcite shows generally low (15.7) albeit highly variable values (4.6-51). Epidote is LREE-enriched (median $\text{La}_{N}/\text{Yb}_N = 17$) and the only vein mineral with a distinctly positive Eu anomaly (Figure 5(d)). Also, fluorite and chlorite are LREE-enriched (median $\text{La}_{N}/\text{Yb}_N = 3.9$ and 6.4, respectively), whereas calcite is HREE-enriched (median $\text{La}_{N}/\text{Yb}_N = 0.14$ and median $\text{Pr}_{N}/\text{Yb}_N = 0.07$) and MREE-depleted (median $\text{La}_{N}/\text{Nb}_N = 1.7$ and median $\text{Tb}_{N}/\text{Yb}_N = 0.06$, Figures 5(d)-5(f)).

Determination of bulk-vein chemistry was challenging because the veins are heterogeneous in terms of the mineralogical composition. Therefore, bulk-vein chemistry was not assessed from measurements on whole-vein specimens, but it was calculated based on the proportion of mineral species and the median metal concentration determined for each of these minerals (Figure 3(c)). Based on this calculation, the veins have concentrations of Fe, Mg, Mn, and Sr that were overall higher (Appendix A) but have $\sum$REE (median = 15 ppm) that are much lower than in the bedrock of the area. The REE in the veins display a concave MREE-depleted fractionation pattern with a positive Eu anomaly (Figure 3(c)).

4.4. Fluid Inclusions in Greisen and Veins. Fluid inclusion analysis of quartz in greisen and veins revealed diverse compositions of several fluid types representing an early high-temperature stage and a later CO₂-bearing stage. The main results are presented below, whereas additional details are given in Appendices C and D.

4.4.1. Early Stage. Primary randomly scattered inclusions (Type 1) in quartz±epidote±calcite±fluorite veins are composed of an aqueous CaCl₂-NaCl-H₂O fluid with a high salinity (33.6-39.8 eq. mass % NaCl) and with halite and/or ferropromsialite ((Fe, Mn)₃Si₂O₇(OH, Cl)₁₀) and hematite as solid phases in coexisting inclusions. The salinity was deduced from the melting temperatures of halite [41] at 233-321°C (mean = 265°C). Total homogenization with the dissolution temperature of ferropromsialite recorded at 423-429°C (mean = 426°C) may be considered close to the formation temperature of the veins.

The majority of the halite-bearing inclusions display a partial (liquid+vapor+solid to liquid+solid) homogenization at 150°C followed by total homogenization at 233-321°C by the melting of halite (liquid+solid to liquid). Applying the method of intersecting isochors and liquidi for the NaCl-H₂O system as discussed by Bodnar [42] and combining the obtained temperature data with known experimental data, the approximate trapping pressure is estimated to be at 2-3.5 kbar.

The same type of inclusion also appears along secondary (Type 1) fluid inclusion planes both in greisen and veins, where they were trapped after the fracturing of the quartz and a gradual decline in pressure to 1 ± 0.5 kbar. The pressure is estimated according to the method of Bodnar [41] from total homogenization temperatures (liquid+solid to liquid) measured at 147-192°C (mean = 176°C) by halite melting, corresponding to a salinity of around 30 eq. mass % NaCl [41] and with a preceding partial homogenization (liquid+vapor+solid) in the range of 100-150°C.

4.4.2. Late Stage. After further fracturing, the early highly saline stage was followed by an inflow of CO₂ in the veins. The pressure initially at this stage is unknown, but molar volumes obtained from the homogenization and melting of CO₂ inclusions in clusters (Type 2A) display a range from 47 to 55 cm³/mole, which suggests a CO₂ flow during a decreasing pressure to a level for the predominant condition of this stage. Secondary CO₂ carbonic inclusions (Type 2A) with molar volumes close to 55 cm³/mole were trapped contemporarily with secondary aqueous-carbonic inclusions (Type 2B) with CO₂ molar volumes of 55-70 cm³/mole and a salinity of 13.5-22 eq. mass % CaCl₂ + NaCl calculated from the gas hydrate dissociation [43, 44] and with secondary aqueous inclusions (Type 2C) with a similar salinity estimated from ice-melting temperatures [41]. The occurrence of these coexisting carbonic and aqueous fluid inclusions indicates that the fluids during the late fluid-stage remained heterogeneous as two immiscible fluids and were trapped simultaneously as Type 2A-C inclusions. Sporadic interaction of the aqueous and carbonic fluids has resulted in the precipitation of nahcolite (NaHCO₃) in aqueous-carbonic inclusions with >20 vol.% CO₂ phase and burbankite ([(Na, Ca)₃(Sr, REE, Ba)₃(CO₃)₅]) or calcite in aqueous-carbonic inclusions with <20 vol.% CO₂. An approximate pressure estimate of 0.3-0.7 kbar can be made using the method by Roedder and Bodnar [45] with the intersection of isochors for the molar volumes 55-70 cm³/mole of the CO₂-phases (secondary Type 2A and Type 2B) with the isochors for the aqueous phase represented by homogenization temperatures to liquid between 100 and 160°C and a salinity of 22 eq. mass % CaCl₂ + NaCl for Types 2B and 2C.

4.5. Isotope Geochemistry and Geochronology. The results of the in situ Rb-Sr geochronology of secondary mineral assemblages are shown in Figure 6. Dating of muscovite±biotite±K-feldspar±calcite±fluorite in three greisen samples yielded ages of 1433 ± 6 Ma, 1432 ± 7 Ma, and 1431 ± 5 Ma. Dating of biotite±calcite±epidote in two vein samples resulted in
ages of 1432 ± 8 Ma and 1432 ± 4 Ma. The radiometric data is contained in Appendix E.

The isotopic systems δ34S, 87Sr/86Sr, and δ18O versus δ13C have previously been used in this kind of settings to distinguish between veins belonging to different geological eras [26, 46]. Consequently, δ34S values of pyrite in veins and greisen, δ18O and δ13C values of calcite in the veins, and 87Sr/86Sr values of calcite and fluorite in the veins and greisen were analysed in some samples and collected from the literature in others (Figure 7). For each of these isotopic systems, the samples showed a small variability (Figure 7). Appendix F contains the stable isotopic data.

5. Discussion

5.1. Genetic Links. The coupling of the hydrothermal greisen and vein mineralization to fluids emanating from the Götemar granite was strongly supported by age determinations, as in situ Rb-Sr age analyses of three greisen and two vein samples representative for these mineralizations have a narrow interval of 1432 ± 8 Ma that overlap with the emplacement age (1433 ± 10 Ma) of the Götemar granite [20]. In addition, the veins and greisen show isotopic ratio signatures (87Sr/86Sr of calcite and fluorite, δ34S_{VCDDT} of pyrite, and δ18O_{VPDB} and δ13C_{VPDB} of calcite) that overlapped and were significantly different to the corresponding signatures of younger veins formed in the Neoproterozoic and Phanerozoic (Figure 7) eras. Thus, there is strong evidence of a genetic link between greisen and vein formation and granite emplacement that allows interpretation of REE transport and fractionation along this hydrothermal system. Geophysical modelling has revealed that the Götemar granite stretches southwards in the subsurface, reaching the northernmost KLX boreholes (Figure 1), where abundant mica-rich greisen veins indicate close proximity to the granite. However, influence from the contemporary Uthammar granite [47] further south of Laxemar (Figure 1) cannot be entirely excluded. Strategies for future studies to distinguish between these two magma sources may include quartz trace element composition, as utilized in [48].


No pH estimates exist for the hydrothermal fluids forming the greisen and veins, but several observations suggest that it is unlikely that fluids avoided extensive rock buffering for up to two km in the thin pathways of the veins: (i) primary minerals in the wall rock adjacent to the veins are extensively altered [29]; (ii) the fluid inclusions were characterized by moderate to high salinity, Na+, Ca2+, and K-chloride-dominated compositions, and Na, Ca, Fe, or Si in the solid phases indicating that feldspar and possibly pyroxene, biotite, and magnete contributed with chemical species throughout greisenization and veininig; and (iii) at high fluid/rock ratios, greisen mineral assemblages are dominated by topaz-quartz [49], a feature not observed in the study area where the amount of topaz in the investigated greisen was very low. Boiling, or effervescence, is another process that may either decrease or increase fluid pH depending on the acidity of the gas phase released by unmixing [50, 51]. The lack of a vapor phase in the primary fluid inclusions and the absence

![Figure 6: Isochrons of (a–c) greisen and (d–e) vein samples dated by in situ Rb-Sr dating in this study, with ages in 1 s. (f) BSE microphotograph of epidote and calcite with albite and biotite in greisen sample KLX06: 593 m. bt=biotite, ms=muscovite, kf=K-feldspar, cc=calcite, fl=fluorite, and epi=epidote.](Image 388x477 to 519x567)
Figure 7: Isotopic composition of Mezoproterozoic minerals compared with isotopic composition of minerals of Neoproterozoic, Paleozoic, and post-Paleozoic minerals occurring at the site, collected in this study, and collected previously in Drake and Tullborg [26].
of a salinity increase with a decreasing homogenization temperature suggest that the formation depth of greisen and veins at 4–8 km [21] was too deep for boiling to occur. Taken together, this implies a system with a low fluid/rock ratio where fluid pH was extensively buffered by the wall rock at both greisen and vein stages. The pH buffering was possibly accompanied by increasing oxygen fugacity, $f_{O_2}$, as the fluids dispersed into the veins, as indicated by Ce and Eu anomalies in calcite and fluorite (Appendix H).

Maximum temperatures of 430°C in primary fluid inclusions in the veins represent the lower end of the decreasing thermal gradient from the magmatic crystallization stage through greisenization and subsequent vein formation. The crystallization temperature of the Götemar granite is here tentatively estimated at about 945°C by Ti-in-zircon geothermometry [52] using the trace element composition of one zircon grain (Appendix B). This single-spot estimate is rough as it cannot account for intra- and intercrystalline Ti variation that can be substantial in zircon [53], but it is still reasonable given that A-type granite (with the exception of alkaline granites) crystallization temperatures are normally above 900°C [54]. The greisen formation temperature is estimated at about 460–480°C by two-feldspar thermometry [55] based on the $K_2O$, $Na_2O$, and CaO composition of coexisting K-feldspar and plagioclase (Appendix G) and a pressure set to 2 kbar representing the established granite intrusion depth of 4–8 km [22]. The uncertainty of the temperatures yielded by this feldspar geothermometer has been estimated at ±30°C [55]. Moreover, the sensitivity for compositional differences is high. The low concentrations of $Na_2O$ and CaO in K-feldspar and K$_2O$ and CaO in the albite plagioclase end member approaching detection limits of the SEM-EDS analysis imply that the uncertainties are likely higher and that the temperature range thus should be treated with caution. A difference of one order of magnitude for either of these compounds causes temperature shifts of up to 100°C, with $Na_2O$ and CaO in K-feldspar being the most sensitive. The indicated temperatures are nonetheless in the range of 400–600°C required for greisenization [56].

5.3. General REE Behavior. The REE concentration estimated for the bulk vein was only 15 ppm, which is three times lower than that for greisen, more than an order of magnitude lower than that for the TIB wall rock, and approximately 20 times lower than that for the anorogenic granite. This is a reflection of the low ΣREE content of all the REE-bearing minerals composing the veins, including calcite (median = 15.7 ppm), epidote (61 ppm), fluorite (2.8 ppm), and chlorite (1.5 ppm). The low REE concentrations of these vein minerals become particularly obvious when comparing with the concentrations in the corresponding minerals in granite (median fluorite: 2156 ppm) and greisen (calcite: 190 ppm, epidote: 1120 ppm, fluorite: 210 ppm, and chlorite: 59 ppm, Appendix A; Figures 8(a)–8(d)). Although these features may be partly explained by transport into the veins of REE-poor fluids such as meteoric water or host-rock brines, such a mechanism seems to have been minor or insignificant considering that (i) fluid mixing was sparsely encountered, (ii) no clear trends in the REE patterns can be linked to wall-rock influence, and (iii) the diverging REE patterns between the different deposition stages and among the minerals at each stage suggest that fluid transport and deposition mechanisms dominated REE fractionation. Hence, the low REE content in the veins strongly suggests that the conditions during intrusion of the granite did not favor extensive REE partitioning into the fluids that were separated and transported into fractures of the country rock, despite that these fluids contained ligands with a capacity to complex and transport the REE including (i) fluoride as evidenced by the occurrence of fluoride and topaz in greisen and fluorite in the veins (Figure 2), and that fluorine is enriched (on average by 11%) in the altered wall-rock zone around the veins (typically red stained) relative to fresh rock [30]; (ii) carbonate as evidenced by calcite in both greisen and veins (Figure 2); and (iii) chloride as evidenced by the highly saline fluids and the presence of halite and sylvite in greisen.

The data thus point to a large retention of REE in the granite. This is explained mainly by the abundant incorporation of these metals into the accessory minerals fluorite, monazite, and apatite (Figures 4(a) and 4(b)). The high concentrations in monazite and apatite indicate that the retention was primarily a consequence of the high uptake by phosphates [6], whereas the high concentrations in fluorite also show that this mineral constituted a significant trap at the magmatic stage, in line with observations elsewhere that fluorite can incorporate significant quantities of REE from a melt [57]. Unlike the phosphates, fluorite was also abundant in greisen and veins, with an overall pattern of decreasing REE concentrations in the sequence granite>greisen>veins (Figure 8(a)).

5.4. Fractionation Patterns of REE in Greisen and Veins. The REE displayed diverse and distinct fractionation patterns among the vein minerals. Calcite was MREE depleted and HREE enriched (Figures 5(e) and 5(f)), epidote was enriched in the LREE (Figure 5(d)), and fluorite was on average MREE depleted (Figure 5(f)). This shows that albeit overall tiny fluxes of REE into the veins, the vein minerals had in a distinct manner captured and fractionated the REE.

The morphological features suggested that epidote occurred typically, although not exclusively, along the borders of the veins (Figures 2(a)–2(d)). There is thus evidence that of the two major REE-carrying minerals in the veins—epidote and calcite—the former was generally precipitated earlier than the latter. If assuming that the calculated REE pattern of the whole vein (Figure 3(c)) represented the bulk fluid REE composition, epidote precipitation caused a preferential removal of LREE (~85% of the LREE pool, Figure 8(f)), reflecting the consistent enrichment of LREE over HREE in the epidote specimens (Figure 5(d)). Subsequent calcite precipitation, preferentially in the middle of the veins (Figures 2(a)–2(d)), consequently occurred from residual fluids depleted in REE in general and in LREE in particular, causing the calcite to attain low REE concentrations characterized by progressive HREE enrichment (Figures 5(e) and 5(f)) with up to 92% of the HREE pool being trapped in calcite (Figure 8(f)). This fractionation phenomenon was most likely intensified by the tendency of
Figure 8: Comparison of chondrite-normalized REE concentrations in different sample types in (a) fluorite, (b) epidote, (c) calcite, and (d) chlorite. (e) Chondrite-normalized [40] REE concentrations calculated for bulk vein, the residual remaining after the hypothetical withdrawal of epidote, and the contribution by calcite after epidote withdrawal (the epidote withdrawal and calcite contribution were calculated by multiplying the average REE concentrations for the mineral with its estimated mode in the veins), and (f) REE composition of vein minerals normalized to the estimated mineral mode of the veins.
LREE to preferentially become incorporated in epidote as compared to calcite, that is, the REE partitioning for epidote/calcite is typically high for LREE (approximately 100) and frequently decreases across the REE series and reaches values of approximately 1 for HREE [58, 59]. The remaining two REE-carrying minerals in the veins—fluorite and chlorite—were less abundant and had considerably lower \( \sum \)REE concentrations than calcite and epidote (Figure 5(f)), and therefore were negligible in terms of the REE budget of the veins (Figure 8(f)).

The REE patterns of vein calcite and fluorite were opposite to those in greisen. In the veins, calcite was a major REE depositional agent and displayed HREE enrichment, whereas fluorite showed no HREE enrichment as compared to LREE (Figures 5(e) and 5(f)). In greisen, fluorite was a major REE depositional agent and displayed HREE enrichment, whereas calcite had a flat pattern with a slight MREE depletion (Figure 5(c)). In addition, the HREE enrichment of greisen fluorite (Figure 5(c)) was also in contrast to the flat REE trend of fluorite within the Götemar granite (Figures 4(a) and 8(a)). This shows that mineral-specific REE partitioning at the hydrothermal temperatures prevailing at the greisen and vein formation was not a major control of REE fractionation patterns within individual minerals. At the specimen scale for both greisen and veins, the REE patterns of individual minerals were not correlated with coprecipitating mineral paragenesis. For example, there was a similar HREE enrichment of calcite both in vein samples with and without epidote and/or fluorite. The most likely explanation for this feature is the sampling limitations set by the drill-core diameter, that is, only a small part of the entire vein was collected in a single sample. Consequently, major minerals in terms of both relative abundance and REE capture, such as calcite and epidote, most likely occurred in all vein systems although they were not identified in every single sample.

5.5. REE Speciation in the Hydrothermal Fluids. Assessment of the complexation of REE transported in the hot and saline late-magmatic hydrothermal fluids emanating from the granite intrusion is challenging. The existence of abundant calcite and to some degree fluorite in the veins (and opposite proportions in the greisen) and NaCl- and water-rich inclusion types in the greisen and veins means that a variety of transporting and/or depositional REE ligands such as chloride, fluoride, hydroxide, and carbonate existed and may have contributed to the ultimate fate of REE [5, 10]. Additionally, the low \( \sum \)REE concentrations in greisen and particularly the veins indicate low REE concentrations in the source fluids, further complicating the evaluation of REE complexation during transport and deposition. Furthermore, there is the decreasing stability of REE complexes with decreasing temperatures [5]. The temperature constraints indicate a decreasing gradient of at least 300–400°C from granite to greisen and an additional lowering by 30–50°C towards the initial vein stage. This means that simple cooling may have been a significant depositional mechanism as well. Despite this, general assessments of REE complexation can be made.

Chloride is typically the most abundant REE transporting ligand in hydrothermal systems due to the high solubility of chloride minerals [11]. In neutral to slightly acidic hydrothermal solutions, chloride complexes are more stable for LREE than HREE, while under strongly acidic conditions the complexation strength is reversed with more stable complexes for HREE [5, 60]. The strong indications that the Cl-rich greisen and vein fluids most likely had a circum-neutral pH mean that they could have contributed to keep LREE preferentially in solution as Cl⁻ complexes. The predominance of chloride complexation typically increases to higher pH with increasing temperature [5], which suggests that the most substantial REE-chloride aqueous transportation occurred close to the granite, as corroborated by high salinities in the high-temperature fluid inclusions [12].

High Y to Ho ratios in fluorite in the veins (70–146, median = 85), granite (72–79, median = 75), and greisen (24–57, median = 44) as compared to the corresponding ratio in chondrite (28) indicate REE-fluoride complexation in the source fluids [61] because of the stronger partitioning of Y than Ho (and other REE) in complexes with fluoride [62, 63]. However, the increase in Y/Ho in the veins may also have been influenced by F-input via biotite alteration [64] and, in particular, increased stability of REEF\(^{2+}\) at the lower temperature stage [65]. The extent of REE complexation with fluoride can therefore not be directly assessed, but as fluoride typically is a depositional rather than a transporting ligand [11] and as the fluorite contents in the veins was low, it is likely that fluorite did not contribute significantly to transport REE beyond the greisen stage.

The low REE concentrations in the studied hydrothermal system and the naturally low hydroxide contents in acidic fluids suggest that hydroxide was not a significant transporting agent at early greisen stages where the combination of low pH and temperatures above 350°C [11, 12] disables complex stability. Hydroxyl REE complexation was likely also insignificant during vein formation since the hydroxyl species is only considered to impact REE mobility below 200°C [12].

Carbonate ligands have been shown to form strong complexes with MREE and HREE at both high (600–800°C; [66]) and low (25°C; [67]) PT conditions, whereas data for REE partitioning of carbonate complexation is absent in the moderate hydrothermal regime [5]. Modelling of REE speciation and fluid/calcite partitioning in H\(_2\)O-NaCl-CO\(_2\) solutions in temperatures ranging between 100 and 400°C predicts bicarbonate or carbonate complexation at low salinity but suppresses carbonic species in favor of chloride complexes at high salinity [12]. In our vein system, with fluids rich in NaCl and with temperatures up to 430°C, calcite precipitation was impeded despite probable calcite saturation [12] and high CO\(_2\) pressure until fluid temperature and pressure successively decreased enough to enable H\(_2\)CO\(_3\) dissociation into the aqueous species [68]. This feature is consistent with the low abundance of calcite in greisen and also implies that carbonate complexation had a negligible influence on REE fractionation at the high-temperature hydrothermal stages. The presence of burbankite in vein fluid inclusions shows, however, that REE accompanied carbonates in the vein.
system, at least towards the distal (lower PT) parts of it. High fluid salinity makes HREE-carbonate complexes progressively less stable compared to LREE-carbonate complexes and thus causes preferential HREE enrichment in calcite [12], which is consistent with the typical HREE enrichment measured in our vein calcites (Figures 5(e), 5(f), and 8(c)). Hence, this mechanism coupled with the preferential accumulation of LREE in epidote (Figures 5(f), 8(b), and 8(f)) explains the characteristic HREE enrichment in the vein calcites (Figures 5(e), 5(f), and 8(c)). HREE enrichment of the kind shown here for vein calcite has only in rare occasions been reported in sedimentary [69] and igneous settings [70]. Our findings show that coupled complexation-driven REE retention and preferential REE partitioning into pre- or coprecipitating phases (in this study epidote) may constitute important processes in precipitating calcite with high HREE to LREE ratios from REE-deficient hydrothermal greisen and vein systems.

6. Conclusions

We have performed a detailed study on transport and fractionation of REE by fluids generated by a Mesoproterozoic granite intrusion and ultimately producing greisen dispersed in thin veins in Paleoproterozoic granitoids. The genetic link between granite, greisen, and veins was determined by high spatial resolution Rb-Sr dating and stable isotopic signatures. Fluid-inclusion microthermometry revealed that multiple pulses of moderately to highly saline aqueous to carbonic solutions caused greisenization and vein formation at temperatures ranging from 430°C at the early hydrothermal stage in the veins to above 200–250°C throughout. Although the veins may have received REE-poor fluids such as meteoric water or host-rock brines, there is support for a major portion of hydrothermal fluids dispersed from the granite intrusion into the veins. Hence, the low \( \sum \)REE concentrations of the veins (15 ppm) and also of the greisen (75) as compared to the granite (320) were explained by limited REE partitioning into the hydrothermal fluids that were separated and transported from the granite intrusion into fractures of the country rock. The data thus point to a large retention of REE in the granite, explained mainly by abundant REE incorporation into the accessory minerals fluorite, monazite, and apatite.

Despite the low REE concentrations in the veins and greisen, the minerals composing these materials displayed diverse and distinct REE-fractionation patterns. In particular, the REE features of vein calcite and fluorite were opposite to those in greisen. In the veins, calcite displayed HREE enrichment and fluorite showed no HREE enrichment as compared to LREE, whereas in greisen calcite had a flat pattern with a slight MREE depletion and fluorite displayed HREE enrichment. In addition, the HREE enrichment of greisen fluorite was in contrast to the flat REE trend of fluorite within the Götemar granite. This shows that mineral-specific REE partitioning at the hydrothermal temperatures prevailing at the greisen and vein formation was not a major control of REE-fractionation patterns within individual minerals. The HREE enrichment of vein calcite is a relatively rare feature. It is explained largely by initial epidote precipitation that trapped much of the LREE leaving a residual fluid enriched in HREE from which calcite precipitated, with a likely contribution from carbonate complexation at the vein stage compared to suggested chloride complexation prior, or up to, greisenization.

Data Availability

The major and trace element data, as well as fluid-inclusion microthermometric, radiometric, and stable isotope data of the bulk materials and minerals used to support the findings of this study are included within the article or within the supplementary information files in Appendices A–H.

Conflicts of Interest

The authors declare that there is no conflict of interest regarding the publication of this article.

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Supplementary Materials

Supplementary materials include major and trace element compositions of bulk rocks in Appendix A, major and trace element compositions of minerals in Appendix B, fluid-inclusion microthermometric data in Appendices C and D, radiometric Rb-Sr geochronological data in Appendix E, stable isotope data in Appendix F, feldspar thermometric calculations in Appendix G, and mineral Ce and Eu anomalies in Appendix H. (Supplementary Materials)

References


