Rare Earth Elements (REE)-minerals in the Silius fluorite vein system (Sardinia, Italy)

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The Silius vein system located in SE Sardinia (Italy) is analogous to other late- to post-Hercynian mineralization of this type in Europe. The system consists of two main veins, characterized by various generations of fluorite, calcite and quartz, with initial ribbon-like geometries, followed by breccias and cockade-like encrustations. In this study, aimed at investigating the REE concentrations in the Silius vein system, average REE amounts around 800 ppm (locally over 1500 ppm ΣREE), related to the presence of the REE-bearing minerals synchysite-(Ce) and xenotime-(Y), were detected in the carbonate gangue of the fluorite orebody. From the geochemical characteristics of these minerals, and the textural relationships with other gangue phases, it is likely that synchysite-(Ce) and xenotime-(Y) formed at the same P-T-X conditions as the other minerals of the Silius fluorite mineralization. Synchysite-(Ce) and xenotime-(Y) at Silius could be related to a local circulation phenomenon, where the REE are derived from a REE-bearing source rock in the basement of southeastern Sardinia, which has been leached by the same fluids precipitating the fluorite/calcite mineralization. REE minerals could possibly be found elsewhere in association with the carbonate gangue of still unexploited parts of fluorite hydrothermal vein deposits, as well as in dumps and tailings accumulated during fluorite processing.

Keywords: SE Sardinia, hydrothermal veins, REE, fluorite, REE-minerals, carbonates
1. Introduction

Rare Earth Element (REE)-bearing minerals occur in many igneous, sedimentary, and metamorphic rocks, where they may be concentrated in ore deposits both related to igneous and hydrothermal processes, or associated with sedimentary environments and weathering (Chakhmouradian and Wall, 2012). REE-rich ores in carbonatites and peralkaline igneous rocks can be directly derived from magmatic processes, but also from metasomatism and hydrothermal remobilization of magmatic REE minerals (Chakhmouradian and Zaitsev, 2012; Gysi and Williams-Jones, 2013). Other REE-bearing mineral deposits can be associated with hydrothermal quartz- and fluorite-bearing veins of orthomagmatic derivation (Samson et al., 2004; Williams-Jones et al., 2000). Typical weathering-related REE-mineral concentrations occur in placers, and in laterite and ion-absorption clay deposits (Chakhmouradian and Wall, 2012).

REE concentrations have been also detected in several fluorite vein systems, not directly related to magmatism, occurring in the Hercynian basement of Europe. Hercynian fluorite vein systems were emplaced in the European basement between late Paleozoic and early Mesozoic, as a consequence of distal tectonics (e.g. continental extension) associated with the initial opening of the Atlantic Ocean, and circulation of brines within the basement (Muchez et al., 2005; Muñoz et al., 1999, 2005). To date, REE contents of these systems has largely focused on the Paleozoic-Mesozoic hydrothermal veins themselves, focused at defining their occurrence in fluorite, caused by ion-substitution processes (Dill et al., 2011; Möller, 1991; Möller et al., 1976, 1994; Möller and Giese, 1997; Schwinn and Markl, 2005). Several studies, carried out on fluorite veins in south Germany (e.g. Harz Mountains, Schwarzwald, Bohemian Massif), revealed that the measured REE concentrations are associated with the occurrence of specific REE minerals (e.g. Dill et al., 2011; Giere, 1996, and references therein; Haack et al., 1987; von Gehlen et al., 1986). Comparable research was also carried out on Triassic-Jurassic fluorite deposits in the Hercynian terrains of northern Africa (Bouabdellah et al., 2010; Cheilletz et al., 2010).
We present here the results of an investigation aimed at evaluating the REE behavior and fractionation in the Silius fluorite vein system, located in SE Sardinia, Italy. The Silius system is analogous to other late- to post-Hercynian mineralization of this type in Europe (Natale, 1969; Boni et al., 2009). Fluorite and galena reserves were evaluated at 2 million tons of raw material in 2006 (Castorina et al., 2008), when mining operations ceased, but a new exploration campaign has been recently carried out, to find new fluorite resources. The REE contents in the Silius fluorite ore had been already reported by Castorina et al. (2008), in relation to a more general study on the different types of fluorite deposits in Sardinia. These authors used the REE content in fluorite to distinguish between the different hydrothermal F-ore deposits on the island, and compared their data with the REE contents of other Hercynian fluorite mineralizations in Europe.

As for other classical studies on European fluorite deposits (e.g. Dill et al., 2001), we have re-evaluated the REE amounts contained in both fluorite and host rock. Moreover, we have also carried out a thorough mineralogical research on the various gangue components of the vein system. The results of this study, aside from obtaining new data on the nature of the REE minerals in the Silius veins, have shed new light on the characteristics of this kind of fluorite ore and its gangue.

2. Geological Setting

2.1. Regional Geology

The Sardinia-Corsica Paleozoic basement represents a fragment of the Hercynian orogen (Arthaud and Matte, 1977; Carmignani et al., 1994; Crowley et al., 2000) that can be subdivided into three tectono-metamorphic zones (Fig. 1A): i) a “foreland zone”, outcropping in southwestern Sardinia, with low-grade or no metamorphism, ii) a “nappe-zone”, in southeastern and central parts of Sardinia, affected by low- to medium-grade metamorphism, and iii) an “inner zone”, in northern Sardinia, characterized by medium- to high-grade metamorphic signatures (Carmignani et al., 2001).
The “external zone” successions (SW Sardinia), spanning in age from early Cambrian to Devonian and Carboniferous, are represented by clastic and carbonate, mainly shallow water sedimentary rocks (Bechstädt and Boni, 1994; Carmignani et al., 2001).

The “nappe-zone” area, where the Silius mineralization is located, is characterized by various lithotypes (Fig. 1B). The upper Cambrian-Ordovician successions mostly consist of siliciclastics (metasandstones, phyllites and quartzites), collectively grouped in the “Arenarie di San Vito” (Fig. 1B) and “Solanas” Formations (not outcropping in the study area). These Formations are overlain by the so-called “Ordovician magmatic and volcano-sedimentary complex” (Carmignani et al., 2001), which consists of both effusive and intrusive igneous products, displaying an almost complete arc-related calc-alkaline suite of middle to late Ordovician age (Oggiano et al., 2010; Gaggero et al., 2012). In the Silius area the Ordovician volcano-sedimentary complex, named “Porfiroidi” Formation (Fig. 1B), is represented by rare andesitic lavas and abundant metasediments derived from the reworking of the former, overlain by metarhyolites and metarhyodacites (Carmignani et al., 1994).

The Ordovician magmatic and volcano-sedimentary complex is unconformably overlain by siliciclastic continental to neritic sediments, which were deposited during a period of active intraplate magmatism, resulting in alkali basalt flows, sills and dykes (late Ordovician-Silurian) (Oggiano et al., 2010; Gaggero et al., 2012) (Fig. 1B). In the Silurian to Devonian period, the magmatic activity stopped and pelagic sediments (Graptolitic Shales, or “Formazione degli Scisti a Graptoliti”) were deposited (Fig. 1B). These consist of black shales and pelagic limestones, and are interrupted at the top by the Carboniferous (Culm-type) flysch (Carmignani et al., 2001).

The “inner zone” is represented by a high-grade metamorphic complex, mainly consisting of amphibolites, which are southward bounded by an eclogite-bearing belt, interpreted as the suture zone of the Hercynian orogen (Carmignani et al., 1994).

The Hercynian compressive deformation was concluded by the emplacement of granite batholiths that occurred during the extensional period associated with the collapse of the orogen (310-280 Ma; Di Vincenzo et al., 1994; Carmignani et al., 2001). The entire basement was later intruded by
Permian to Triassic magmatic dykes of bimodal composition (Ghezzo and Orsini, 1982; Atzori and Traversa, 1986; Carmignani et al., 2001). During this period, several hydrothermal vein systems containing fluorite, barite and sulfides were developed in SE Sardinia. The Silius mineralization corresponds to one of these vein systems (Natale, 1969). Other post-Hercynian occurrences in the area are set within cataclastic and mylonitic belts, as the so-called “Filone Argentifero del Sarrabus”, a complex association of stockworks and veins of Pb-Cu-sulfides and silver minerals, as well as the stibnite (±scheelite) vein systems of Su Suergiu and Corti Rosas (Valera et al., 1974; Belkin et al., 1984; Carmignani et al., 1978).

2.2. The Silius hydrothermal veins

The Silius vein system (Fig. 1B) crops out discontinuously for 2-3 km along a NE-SW strike, between the Acqua Frida area (39°30′46″N-9°15′05″E), the Muscadroxiu shaft (39°31′00″N-9°15′18″E) and the Genna Tres Montis shaft (39°31′38″N-9°16′00″E), at an elevation ranging between 600 and 700 m a.s.l. At the 400 m level, the veins reach a maximum length of 4 km. The system consists of two main veins, called “San Giorgio” and “San Giuseppe” (Fig. 2A), which have an almost vertical dip (about 75°), and a strike of N045E along the main sector Acqua Frida - Genna Tres Montis, and N065E northeastward from Genna Tres Montis. These two veins are almost parallel in the topmost levels, with a spacing of 70 m on average, but they coalesce at a depth of 350 m a.s.l. in the NE sector, and at a depth of 450 m a.s.l. in the SW sector of the mineralized area. The San Giorgio vein was the first to be formed, and was displaced and brecciated by the San Giuseppe vein. The San Giorgio vein contains bands of chalcedony, pink fluorite, barite and calcite. The San Giuseppe vein is also banded, and shows several generations of fluorite, calcite and galena (Fig. 2A, B, C). The width of the veins ranges between 7-8 m in outcrop and 15-20 m in depth, where they are interconnected (Fig. 2B). The Silius vein system has been mined from the surface (around 600 m a.s.l.) down to 200 m a.s.l., and explored to 100 m a.s.l. The upper part of the vein system contains
fluorite, barite and galena with grades of 35%, 10% and 3%, respectively, whereas in the lower zones the amount of barite decreases, and the fluorite and galena grades can also reach 50% and 6%, respectively. Quartz and carbonates are the main constituents of the gangue. Two less economically important veins (called San Giovanni and Davide) are also parallel to the other veins, and can be traced to the 500 m level.

All the veins are mainly hosted by the “Porfiroidi” Formation and rarely by Late Ordovician sedimentary rocks of the Graptolitic Shales Formation. The fluorite-barite mineralized veins cut locally the late- to post-Hercynian (Permian to Triassic acid and basic-intermediate) magmatic dykes: this relationship defines a relative minimum age for the mineralization process (Natale, 1969). The veins are characterized by successive generations of fluorite, calcite and quartz, with initial ribbon-shaped geometries, followed by breccias and cockade-like textures (Fig. 2D). The different generations are always accompanied by sulfides (marcasite, sphalerite and galena), and in the late stages by barite (Natale, 1969).

In previous fluid inclusion studies on the Silius mineralization, it was reported that most ore minerals precipitated at temperatures in the range of 120-180 °C from fluids with salinities reaching up to ~18 wt.% NaCl equiv. (Boni et al., 2009). Due to the very low first melting temperatures (~~50 °C) recorded in both fluorite and calcite, the authors suggested the involvement of Ca-rich fluids in the mineralization process. Lead isotopes of sulfides in the Silius mineralization have been determined as intermediate between the compositions of Ordovician metavolcanic rocks (“Porfiroidi” Formation) and Hercynian granites (Boni et al., 2009). Sr and Nd isotopes of Silius fluorite, measured by Castorina et al. (2008), have a signature coherent with a brine circulation within both Paleozoic metasedimentary rocks and Hercynian granitoids.

3. Sampling and analytical methods
The samples analyzed in this study were mainly collected in both the San Giuseppe and the San Giorgio veins, along the two vertical sections of the Muscadroxiu and the Central shafts of the mine (the latter is located between the Muscadroxiu and the Genna Tres Montis shafts) at every 20 m, from 100 m a.s.l. up to 200 m a.s.l. At each sampling site, we collected fluorite, calcite, and fragments of the host rock. Additional sampling was carried out at the Acqua Frida shaft, at 300 m elevation, where we collected fluorite and calcite from both veins. In Table A1 are listed all the analyzed samples.

Fluorite and calcite were cleaned of the impurities (sulfides and Fe-oxides) by handpicking under binocular microscope. The purity of the minerals was tested with X-ray diffraction analysis, by using a Seifert–GE ID3003 diffractometer (Dipartimento di Scienze della Terra, dell’Ambiente e delle Risorse, Università degli Studi di Napoli Federico II), with CuKα radiation, Ni-filtered at 40 kV and 30 mA, 3-80 °2θ range, step scan 0.02°, time 10 sec/step, and the RayfleX (GE) software package; a silicon wafer was used to check the instrumental setting. Sample holder was a zero-background plate of quartz crystal cut and polished 6° of the c-axis.

For polished thin sections preparation, the samples were impregnated with Araldite D and Raku Hardener EH 2950 (OMT Laboratory, Aosta, Italy). Cathodoluminescence microscopy (CL) was carried out with a Hot Cathode Instrument at the Heidelberg University. Secondary electron imagining by scanning electron microscopy (SEM), and energy dispersion spectrometry (EDS) investigations on thin sections were carried out with a Jeol JSM 5310, equipped with the INCA X-stream pulse processor and the 4.08 version Inca software (Oxford Instruments detector) (DiSTAR, Università di Napoli, Italy), operating at 15 kV primary beam voltage, 50-100 mA filament current, variable spot size and 50 s net acquisition time. Reference standards were: albite (Si, Al, Na), orthoclase (K), wollastonite (Ca), diopside (Mg), almandine (Fe), rutile (Ti), barite (Ba), strontianite (Sr), eskolaite (Cr), rhodonite (Mn), pyrite (S), sphalerite (Zn), galena (Pb), fluorite (F), apatite (P), sylvite (Cl), smithsonian phosphates (La, Ce, Nd, Sm, Y), pure vanadium (V) and Cornig glass (Th, U). Analytical errors are 1% rel. for major elements and 3% rel. for minor elements.
Quantitative data sets of selected samples were obtained by wavelength dispersion spectrometry (WDS), using a Cameca SX100 electron microprobe operating at 20 kV, 20 nA, and 10 \( \mu \)m spot size (Natural History Museum, London, UK). All the compositions of the standards used are reported in Table A2. Interference corrections have been carried out prior to matrix correction for the following elements: F (overlapping element, Ce), Nd (Ce), Sm (Ce), Ba (Pr) and La (Nd). A monazite (Manangotry) standard (composition and determinations in Table A3) was used as a quality control sample. WDS detection limits for each element are reported in Table 3, and are mostly in the order of 0.01 wt%.

Major, minor and trace (REE) elements in fluorite, carbonate minerals and host rock samples have been measured at the ACME laboratories (Vancouver, Canada) and at the Activation Laboratories (Ancaster, Canada). In both the laboratories, the same protocol was adopted: the samples were pulverized to 85% -200 mesh, to obtain about 20 g of pulp. Major oxides and several minor elements were analyzed by ICP-OES following a LiBO\(_2\)/Li\(_2\)B\(_4\)O\(_7\) fusion and dilute nitric digestion. Loss on ignition (LOI) was calculated by weight difference after ignition at 1000°C. Rare earth and refractory elements were determined by ICP-MS following a LiBO\(_2\)/Li\(_2\)B\(_4\)O\(_7\) fusion and nitric acid digestion. In addition, a separate volume split was digested in Aqua Regia and analyzed by ICP-MS to detect the precious and base metals. Lower detection limits obtained in the two laboratories are reported in Tables 1, 2 and 4.

4. Mineralogy and major element geochemistry

4.1. Fluorite

The analyzed samples consist of medium-sized (<1cm) white/violet fluorite crystals, associated with calcite, quartz, barite and a few minute sulfide intergrowths. In thin section fluorite appears of different generations, each showing a strong blue luminescence under CL (Fig. 3). As reported in the
literature (Natale, 1969), it is possible to distinguish at least three fluorite generations: a first and more extensive generation consisting of fluorite alone (Fig. 3A, B), a second generation of fluorite with a calcite gangue (Fig. 3C, D), and a third one associated with quartz and barite (Fig. 3E, F).

Minor amounts of SiO₂, or various values of LOI or tot-C, detected by bulk chemical analyses (Table 1), are related to intergrown quartz or carbonate minerals in fluorite, also revealed by XRD. The most contaminated samples have not been considered when discussing the relationships between REE sorption processes and genetic conditions.

4.2. Calcite and dolomite

Mineralogical analyses have been conducted on several samples of the carbonate gangue, in order to verify if this consisted solely of calcite, as reported in the literature. X-ray diffraction clearly confirm that the carbonates of both the San Giorgio and San Giuseppe veins consist of a mixture of calcite and dolomite/ferroan dolomite, not yet reported for this vein system. The carbonate gangue can locally contain also small radial aggregates of newly formed kaolinite and quartz.

Calcite has yellow-orange colors under CL (Fig. 3D), and generally appears zoned. This indicates that variable amounts of Mn²⁺ have substituted for Ca²⁺ in the calcite lattice (Götze, 2012). Under microscopic observation, dolomite replaces calcite in patches (Fig. 3D). Replacive dolomite shows an extremely low luminescence under CL and consequently, when associated with calcite, it appears generally dark (Fig. 3D, F). Where the replacement is strongly pervasive, the dolomite colors under CL are dark-reddish. According to Götze (2012), the red CL color of dolomite is generally due to variable amounts of Mn²⁺ ⇌ Mg²⁺ substitution in its structure. The low luminescence intensity of the Silius dolomite is related to the presence of discrete Fe²⁺ amounts in the cationic sites, which quench the luminescence effects related to Mn²⁺ (Götze, 2012).

SEM analyses confirmed that dolomite is not pure, but rather consists of the ferroan variety and ankerite, with Fe contents up to 23 wt.% FeO, and Mn up to 4 wt.% MnO (Fig. 4). Fe-bearing
dolomite has a mottled texture (Fig. 4A). Two replacive generations have been detected: an initial almost pure dolomite phase (<5 wt.% FeO, ~1 wt.% MnO), which is followed by a high-Fe dolomite phase (>6 wt.% FeO, >1 wt.% MnO) and eventually by ankerite (Fig. 4B, C).

Plotted on a diagram defined by CaMg(CO$_3$)$_2$-CaFe(CO$_3$)$_2$-CaMn(CO$_3$)$_2$ (Fig. 5), the analyzed phases have a composition covering the field between pure dolomite and ferroan dolomite, with an almost continuous substitution between Ca$_{1.08}$Mg$_{0.86}$Fe$_{0.03}$Mn$_{0.03}$(CO$_3$)$_2$ and Ca$_{1.03}$Mg$_{0.42}$Fe$_{0.46}$Mn$_{0.09}$(CO$_3$)$_2$.

Only two outlier data show a more definite ankeritic composition: Ca$_{1.01}$Mg$_{0.19}$Fe$_{0.68}$Mn$_{0.12}$(CO$_3$)$_2$ and Ca$_{1.01}$Mg$_{0.14}$Fe$_{0.73}$Mn$_{0.13}$(CO$_3$)$_2$.

Chemical analyses on the carbonate gangue corroborate the Fe-rich composition of the dolomite. In fact, several samples from the gangue of the fluorite mineralization (Table 2) contain variable Mg and Fe amounts, which result from the mixture of different phases, as calcite, dolomite, Fe-rich dolomite and ankerite. These samples have a very similar composition, though they were collected in different zones of the mine.

4.3. REE-minerals

Abundant REE-minerals have been detected within the carbonate gangue of both Silius mineralized veins (Fig. 4C, D, E, F, G, H). The mineralogy of these phases is quite simple and is mostly represented by synchysite-(Ce) (Table 3), a Ca-REE-fluorcarbonate [CaREE(CO$_3$)$_2$F], and by rare xenotime-(Y) (YPO$_4$). These minerals have been observed only under electronic microscope, because their amounts is below (or very near to) the XRD detection limits adopted in this study (< 1 wt.%).

In the analyzed samples, synchysite-(Ce) occurs as acicular and tabular crystals (Fig. 4C, D), with an average size below 100 µm (Fig. 4E, F). In detail, synchysite-(Ce) has been observed both as single crystals and as composite aggregates in two main locations: firstly within the dolomitized calcite and Fe-bearing dolomite (Fig. 4C, D), and secondly within quartz veinlets and in cavities occurring among the calcite and dolomite patches (Fig. 4E). In the first setting, synchysite-(Ce) is distributed
throughout the irregularly dolomitized carbonate gangue, while it is almost absent in unaltered calcite (Fig. 4D). When synchysite-(Ce) occurs in the quartz veinlets, it is paragenetically followed by the third fluorite generation (Fig. 4F, G). No REE minerals have been found as inclusions in the fluorite crystals of the main generation that have been selected for ICP analyses.

Table 3 shows the chemical composition of synchysite-(Ce). The total REE content is on average around 45-46 wt.%. The formula of synchysite-(Ce) has been built on the basis of $(\text{CO}_3)_2\text{F}$ and 5 negative charges, and is given in Table 3. The average synchysite-(Ce) formula is

$$\text{Ca}_{1.07}(\text{La}_{0.19},\text{Ce}_{0.36},\text{Pr}_{0.04},\text{Nd}_{0.15},\text{Sm}_{0.03},\text{Gd}_{0.03},\text{Y}_{0.13})(\text{CO}_3)_2\text{F}.$$  

Xenotime-(Y), similarly to synchysite-(Ce), occurs as subhedral crystals filling cavities within calcite and dolomite (Fig. 4H). The few analyzed specimens (EDS) are characterized by Dy and Yb contents of ~3 wt.%.

5. REE geochemistry

5.1. Fluorite

Excluding the fluorite samples contaminated by intergrown gangue minerals, the REE amounts determined by ICP on bulk fluorite are solely related to REE diffused in its crystal structure. In fact, no discrete REE minerals have been detected within the analyzed fluorite crystals in contrast to the carbonate gangue.

The REE contents of the Silius fluorite (Table 1) have been normalized to the post-Archean Australian Shale (PAAS; Taylor and McLennan, 1985), in order to compare their distribution with that observed in other deposits of the same type. REE in Silius fluorite are generally lower than PAAS. It is possible to observe that the REE concentrations normalized to PAAS have a roof-shaped pattern, characterized by positive Eu and Y anomalies (Fig. 6A). The Eu anomaly is also reflected in the Eu/Eu* ratio, which is always above 2, except in samples FL-06 and FL-11 (contaminated by
quartz and carbonates), and in FL-05 and FL-12, which have no contamination. The patterns are asymmetric, because the majority of the samples have HREE contents very near to PAAS, whereas the LREE are strongly depleted; in fact, several samples have a La/Yb$_n$ ratio $< 1$. The sample FL-3 is the only not-contaminated specimen to have the MREE and HREE more enriched than PAAS, and the LREE more depleted than PAAS (La/Yb$_n$ ratio $\approx 0.03$). Other samples (Fl-06, Fl-09, FL-10) have a La/Yb$_n$ ratio slightly $> 1$, which is too small to be considered as proof for LREE/HREE fractionation. Moreover, these samples are characterized by normalized MREE values, which are higher than the LREE and HREE values.

5.2. Carbonates

Significant amounts of REE-bearing mineral inclusions, trapped in calcite and dolomite, clearly influence the REE concentrations determined by ICP on the bulk samples of the carbonate gangue. However, the amounts of REE possibly hosted in the calcite and dolomite structure are lower than the WDS detection limits.

The bulk carbonate gangue is characterized by REE contents notably higher than those in fluorite, (Table 2); in two of the considered samples, the total REE amounts are over 1500 ppm. In the diagram of PAAS-normalized REE-contents (Fig. 6B), it is possible to notice that the average pattern is almost flat for the carbonate samples, with positive drops in the Eu and Y positions. In detail, the CA-08 sample is the only one where HREE are more enriched than LREE, and LREE are less abundant than PAAS. The other samples are characterized by an enrichment of both LREE and HREE alike (La/Yb$_n$ ratios around 1), or by a LREE enrichment higher than HREE (La/Yb$_n$ ratios are around 2; only the sample CA-06 has a La/Yb$_n$ ratio slightly $> 4$). The Eu positive anomaly is marked by Eu/Eu* ratios between 1.3 and 2.92.

5.3. Vein system host rocks
Two measured samples from the “Porfiroidi” Formation, which is the main host rock of the Silius deposit, have a REE content around 100 ppm ΣREE (Table 4). One of these samples, P6 is clearly contaminated with fluorite. Among the other types of host rocks, the highest REE contents have been detected in the samples consisting of late-Hercynian porphyritic dykes, whereas the metasedimentary black shales have very low REE values. All the host rocks have PAAS-normalized REE patterns depleted in REE, in comparison to the adopted reference (Fig. 6C). Only a sample of basic-intermediate porphyry has REE concentrations that are on average higher than PAAS. In general, two types of patterns can be observed: flat patterns and LREE-slightly depleted patterns (La/Yb_n<1). All of them do not show spikes or anomalies, either positive or negative. This behavior is different from that of fluorite, which has roof-shaped patterns with Eu and Y positive anomalies, and from the carbonate gangue, which is generally characterized by La/Yb_n ratios > 1 and by Eu and Y positive anomalies (Fig. 6).

5. Discussion

5.1. New insights on the mineralogy of the Silius vein system: are REE-bearing minerals compatible with this mineralization type?

In the previously published studies on the Silius mineralization (Natale, 1969; Castorina et al., 2008; Boni et al., 2009), the deposit was described as a simple association of fluorite, barite, and mixed sulfides in a gangue consisting of calcite and quartz. The dolomite-ankerite phases detected in the present study are paragenetically younger than the other minerals of the veins because they replace earlier calcite. This event was then followed by, or is partly contemporaneous with a minor fluorite-barite-quartz mineralization stage, and by the precipitation of a late calcite phase. Consequently, the formation of both dolomite and Fe-dolomite can be considered as evidence for a short and abrupt
change in the chemical composition of the mineralizing fluids, which were temporarily enriched in Mg and Fe, in order to dolomitize the previously deposited calcite in both San Giorgio and San Giuseppe veins. The occurrence of synchysite-(Ce) and xenotime-(Y), that are paragenetically associated with dolomitized calcite, also marks a sharp temporary change in the chemistry of the fluids that for a while became REE-bearing.

Synchysite (the main REE-carrier in the Silius fluorite veins) is generally associated with orthomagmatic-hydrothermal or hydrothermal-metasomatic REE-mineralizations, but it has been also found in Alpine-type veins (Augé et al., 2014; Chakhmouradian and Wall, 2012; Föster, 2001; Giere, 1996, and references therein; Williams-Jones et al., 2000). This mineral was detected in post-Hercynian veins in Germany (e.g. Harz Mountains, and Schwarzwald), where it is associated with bastnäsite (Giere, 1996, and references therein; Haack et al., 1987; von Gehlen et al., 1986). Small occurrences of synchysite-(Ce) in Sardinia are only recorded in association with molybdenite within Hercynian leucogranites in the small prospects of Su Senargiu (Sarroch) and Perda Pibera (Gonnosfanadiga) (Brizzi et al., 1994; Starà et al., 1996; Boni et al., 2003; Orlandi et al., 2013).

In hydrothermal environments REE are transported mainly as non-fluoride complexes (e.g. chloride species) in fluorine-bearing solutions (Williams-Jones et al., 2012; Migdisov and Williams-Jones, 2014). These authors proposed a model in which fluoride ions would work as a binding ligand that promotes REE mineral deposition, rather than as an agent for REE transport. With this background, the co-genetic association of REE minerals and fluorite would be explained by the low solubility of both fluorite and REE minerals. Then, the ore-forming hydrothermal fluids carry appreciable concentrations of REE (and fluorine in the form of HF) in relatively acid conditions. The REE deposition may occur when the neutralization of the acidity of the solution induces the precipitation of fluoride-bearing minerals that in this model are represented only by bastnäsite. This happens, for example, when fluids interact with carbonate host rocks (e.g. dolomite or limestone), which buffer the pH to higher values, leading to the precipitation of the REE fluorocarbonate bastnäsite, according to the following reaction (Williams-Jones et al., 2012):
REECl\(^{2+}\) + HF + HCO\(_3^-\) = REECO\(_3\)F + 2H\(^+\) + Cl\(^-\) \hspace{1cm} (1)

Any mechanism that produces an increase in pH and/or HCO\(_3^-\) activity, or a decrease in Cl\(^-\) activity will lead to the deposition of this mineral. If the host rock of the mineralization consists of dolomite, the ore fluid will react with the carbonate, providing an effective mechanism to precipitate the ore minerals (Williams-Jones et al., 2012):

\[ 2H^+ + CaMg(CO_3)_2 = Ca^{2+} + Mg^{2+} + 2HCO_3^- \] \hspace{1cm} (2)

This reaction produces a sharp increase in both pH and HCO\(_3^-\) activity. Concerning synchysite-(Ce), we may propose, by analogy with bastnäsite, a possible reaction for its formation based on the chemical reaction (1):

\[ \text{REECl}^{2+} + \text{Ca}^{2+} + \text{HF} + 2\text{HCO}_3^- = \text{Ca(REE)(CO}_3\text{)}_2\text{F} + 3\text{H}^+ + \text{Cl}^- \] \hspace{1cm} (3),

where the acid solution is buffered as in reaction (2).

Xenotime is a common accessory phase in granites and pegmatites, and is a common detrital mineral in siliciclastic sedimentary rocks; however, it may also form during hydrothermal processes (Brown et al., 2002; Kositcin et al., 2003; Schaltegger et al., 2005; Cook et al., 2013). In Sardinia, hydrothermal xenotime was found in the same leucogranite-hosted, molybdenite-related vein occurrences of Su Senargiu and Perda Pibera (Orlandi et al., 2013, 2014). In the Silius paragenesis, xenotime-(Y) has been detected as a phase strictly associated with synchysite-(Ce). Considering that synchysite preferentially hosts LREE, whereas xenotime is the common HREE carrier, this is probably a consequence of the different fractionation of LREE and HREE between the two minerals from the REE-bearing hydrothermal fluid. Gysi et al. (2015) investigated experimentally the thermodynamic properties of xenotime-(Y) and other HREE phosphates in a hydrothermal environment, and in aqueous HClO\(_4\)-H\(_3\)PO\(_4\) solutions, at temperatures up to 250 °C. From these experiments, the authors obtained that the solubility of HREE phosphate solids increases sharply with decreasing temperature. At Silius, most of the ore minerals precipitated at temperatures in the range of 120-180°C from a dominant fluid consisting of a NaCl + CaCl\(_2\)-rich brine (Boni et al., 2009).

Hence, we may assume that the mineralizing fluid that precipitated xenotime-(Y), together with...
synchysite-(Ce), can be similar to the model of Gysi et al. (2015) for xenotime-(Y) precipitation from $\text{H}_3\text{PO}_4–\text{HCl–HF}$-fluid at 150 °C temperature. In these experimental conditions, xenotime is stable in the entire pH range that is higher than ~2. This wide stability range can be also compatible with the conditions of precipitation of Ca-bearing REE phases, which, as previously reported, are conditioned by the buffering reactions between the acid REE-bearing fluids and the carbonate host rock (Williams-Jones et al., 2012).

The occurrence of both synchysite-(Ce) and xenotime-(Y) at Silius is compatible with the genetic environment and the temperatures determined in previous studies by Castorina et al. (2008) and Boni et al. (2009).

5.2. Geochemistry of fluorite and gangue minerals: a comparison with previous studies

The REE occurrence in fluorite has been largely sought in the Paleozoic-Mesozoic hydrothermal veins of Europe, as a way to investigate their genesis (e.g., Möller, 1991; Möller et al., 1976, 1994; Schwinn and Markl, 2005; Dill et al., 2011). It is well known that REE are hosted in the fluorite structure due to ion-substitution processes (Möller et al., 1998):

\[
2 \text{Ca}^{2+} = \text{REE}^{3+} + \text{Na}^+ \\
3 \text{Ca}^{2+} = 2 \text{REE}^{3+} + (\square) \\
\text{Ca}^{2+} = \text{REE}^{3+} + \text{F}^{-}
\]

Möller et al. (1998) also suggested that Eu, Ce and Y anomalies in fluorite REE-PAAS-normalized patterns depend on the environmental conditions (mainly T) during the precipitation of this mineral. In particular, the presence of Eu anomalies is strongly temperature-dependent, because at temperatures > 200 °C, Eu$^{3+}$ is reduced to Eu$^{2+}$, which is very mobile and difficult to be incorporated into the fluorite structure, and therefore tends to remain in the fluid. This produces negative Eu anomalies in fluorites precipitated from a fluid with T > 200 °C. Instead, a Eu anomaly is lacking or slightly positive in fluorite precipitated from a fluid with a temperature < 200 °C (Möller et al., 1998).
Another possibility is that the Eu anomaly was inherited from the REE source rock (Schwinn and Markl, 2005). Silius fluorite shows a clear positive Eu anomaly, which probably indicates precipitation temperatures < 200 °C, in agreement with the homogenization temperatures of the fluid inclusions measured by Boni et al. (2009).

The presence or absence of a cerium anomaly is controlled by the oxygen fugacity of the fluid, which controls the redox reaction between Ce³⁺ and Ce⁴⁺. Ce⁴⁺ does not migrate in the fluid together with other REE, because it is typically retained in Fe and Mn-hydroxides (Möller and Bau, 1993). The absence of Ce anomalies in the REE patterns of Silius fluorite confirms that the fluids precipitating the main fluorite generation were relatively oxygen-poor, a fact that is also confirmed by the presence of sulfides (galena and pyrite > sphalerite) associated with fluorite.

Bau and Dulski (1995) suggested that the Y/Ho ratio varies during the migration of the hydrothermal fluid through the rocks: in particular, a progressive migration of the fluids produces a negative correlation between the Y/Ho ratio and the La/Ho ratio. If we exclude from the discussion the fluorite samples FL-04, FL-06 and FL-11 (Table 1) clearly contaminated by other minerals (quartz, barite, calcite), it appears that the Silius fluorite is characterized by a positive correlation between the La/Ho and Y/Ho ratios (Fig. 7). This would exclude a prolonged migration through the host rock before fluorite precipitation, though PAAS-normalized patterns showing positive Y anomalies (as in this case) generally form when the distance between the sites of REE mobilization and deposition is big (Bau and Dulski, 1995).

REE geochemistry of carbonates in Hercynian fluorite veins has been widely investigated, for example in the deposits of the Harz mountains in Germany (e.g. Möller et al., 1984), in the small occurrences of the Valle de Tena in Spain (Subías and Fernández-Nieto, 1995), and in the El Hammam orebodies in Morocco (Cheillez et al., 2010). More specifically, the REE occurrence in the structure of carbonate minerals, i.e. calcite, siderite, and magnesite, has been described by Bau and Möller (1992) for the German ores. However, in our case study, it is necessary to consider that the REE amounts in the carbonates are mostly related to the occurrence of interspersed synchysite-(Ce)
and xenotime-(Y) mineral inclusions, and that they do not depend on substitution processes of REE within calcite or dolomite lattices. Consequently, it is challenging, for example, to explain why all REE patterns of the bulk carbonates at Silius are characterized by Eu positive anomalies, similarly to those of pure fluorite. In our opinion, the different possibilities accounting for the observed Eu positive anomaly are:

i) the Eu anomaly of the carbonate gangue depends on the physico-chemical conditions during precipitation of the minerals; after Bau and Möller (1992), a positive Eu anomaly in calcite indicates precipitation temperatures of at least 200-250 °C, reducing conditions, and Eu reduction to Eu$^{2+}$, which is easily incorporated into the calcite structure;

ii) the Eu positive anomaly in the carbonate gangue is inherited from the REE source rock: considering the high contribution of synchysite-(Ce) to the total REE concentration in the gangue, it is probable that the REE were not fractionated on the basis of their oxidation state when entering the mineral, but were incorporated in toto from the hydrothermal fluid, thus preserving the footprint of their original source. Considering also that fluorite and its carbonate gangue precipitated together, also the Eu positive anomaly in fluorite could be partly inherited from the REE source, not being only an effect of a temperature-dependent process.

Considering the geological, textural, and paragenetic characteristics of the Silius mineralization, option (i) is highly likely, but considering the high contribution of synchysite-(Ce) to the REE concentration in the bulk gangue, option (ii) can be also valid. Consequently, it is not possible to totally exclude that the Eu anomaly was inherited from the REE composition of an original source rocks. A comparison between the REE patterns of fluorite and carbonate gangue with those of several host rocks at Silius shows that both fluorite and carbonates have patterns completely different from those of their host rocks (Fig. 6). This point would be enough to exclude a provenance of the REE from the leaching of the same rocks hosting the vein system. The association of REE-fluorcarbonates with Fe-bearing dolomite and quartz may indicate that the hydrothermal fluids have leached Mg-Fe
silicatic rocks from the basement, which should have contained discrete REE amounts from the start. Moreover, the presence of a positive Eu anomaly in the normalized REE pattern of both fluorite and carbonates (if partly inherited from the original REE source), may suggest that the REE source was characterized by high feldspar (specifically plagioclase) contents. On this point, for example, Castorina et al. (2008), referring to the experimental work of Schwinn and Markl (2005), suggested a possible contribution of a gneiss-derived hydrothermal fluid to the formation of the Silius vein system. On the other hand, the upper Ordovician alkali basalts occurring in Southeastern Sardinia have many characteristics, which fit the above constraints and make these rocks a potential REE-source for the Silius mineralization. In fact, they are Mg-Fe rich, contain discrete amounts of REE, and also show positive Eu anomalies in the REE chondrite-normalized pattern (Gaggero et al. 2012). The occurrence of xenotime-(Y) at Silius suggests also the presence of phosphorus within the source rocks. Phosphorus could have been leached from the Silurian black shales and pelagic sediments of the Graptolitic Shales Formation, which locally host the fluorite mineralization.

6. Conclusions

This study has shown that REE concentrations in the carbonate gangue of the Silius fluorite deposit (SE Sardinia, Italy), are related to the presence of the REE-bearing minerals synchysite-(Ce) and xenotime-(Y). From the discussion of the data and the comparison with the existing literature, it is likely that both synchysite-(Ce) and xenotime-(Y) formed at the same P-T-X conditions considered for the other minerals of the Silius fluorite mineralization. Rare earth elements are likely derived from a REE-bearing source rock in the basement of southeastern Sardinia, which has been leached by the same fluids precipitating the fluorite/calcite mineralization.

In our opinion, the discovery of REE-minerals in the Silius fluorite mine may open interesting perspectives for the exploration of sub-economic REE concentrations in this type of deposits, where
REE could be recovered as by-product of the fluorite exploitation. Moreover, REE minerals could possibly be found also in dumps and tailings accumulated during the fluorite concentration process.

**Acknowledgments**

We would like to thank Ing. G. Mura, Director of the Silius mine, for having allowed us to sample in the underground levels, and R. de’ Gennaro (DiSTAR, Napoli) for the SEM-EDS analyses. Thanks are also due to the anonymous reviewers, whose comments have highly enhanced the quality of the manuscript, and F. Pirajno for editorial handling. N. Mondillo would like to thank B. Lehmann, for the positive discussion on the Harz fluorite, and R. Herrington for suggestions and help. This work was partially financed by the Università degli Studi di Napoli Federico II, grant RDIP2014 to M. Boni and N. Mondillo.

**References**


Gysi, A.P., Williams-Jones, A.E., Harlov, D., 2015. The solubility of xenotime-(Y) and other HREE phosphates (DyPO₄, ErPO₄ and YbPO₄) in aqueous solutions from 100 to 250 °C and p_{sat}. Chem. Geol. 401, 83–95.


**Figures captions**

Figure 1. A) Major tectonic and metamorphic zones of the Hercynian basement in Sardinia (Italy) (Carmignani et al., 1991, modified); yellow star = Silius area. B) Geological sketch map of the Silius area (Nuova Mineraria Silius s.p.a., modified).

Figure 2. A) Silius Mine - production gallery at 235 m a.s.l.: ribbon structure (fluorite, calcite, galena) of the San Giuseppe vein (Boni et al., 2009, modified). B) Green-purple banded fluorite of the San Giorgio vein. C) & D) San Giuseppe vein (zoned fluorite, calcite/dolomite, galena) cutting and partly replacing the San Giorgio vein: chalcedony-rich fragments of the San Giorgio vein occur as breccia clasts cemented by San Giuseppe calcite (Boni et al., 2009, reinterpreted).
Figure 3. A) & B) First fluorite generation, luminescing blue under CL. C) & D) Cubic crystal of the second fluorite generation in carbonate gangue: calcite (yellow-orange under CL) is replaced by dolomite (dark under CL). E) & F) Third fluorite generation consisting of microcrystals associated with quartz and barite in a veinlet, cutting the carbonate gangue (dark-reddish under CL).

Figure 4. A) Fe-bearing dolomite crystal showing a mottled texture. B) Calcite replaced by dolomite, which locally reaches ankerite composition. C) & D) Acicular and tabular crystals of synchysite irregularly distributed throughout the dolomite, Fe-bearing dolomite and ankerite gangue. E) Acicular crystals of synchysite within quartz veinlets and in cavities between calcite and dolomite. F) Synchysite across dolomitized calcite, quartz and fluorite of the third generation. G) Synchysite aggregate of crystals, clearly cemented by fluorite. H) Subhedral crystal of xenotime cutting across calcite and dolomite. Backscattered images (BSE) from SEM-EDS.

Figure 5. Dolomite and Fe-bearing dolomite chemical compositions plotted in the CaMg(CO$_3$)$_2$-CaFe(CO$_3$)$_2$-CaMn(CO$_3$)$_2$ diagram (EDS analyses).

Figure 6. A) REE concentrations in fluorite bulk samples, normalized to PAAS (post-Archean Australian Shale; Taylor and McLennan, 1985). B) REE concentrations in carbonate gangue bulk samples, normalized to PAAS. C) REE concentrations in host rock samples, normalized to PAAS. ICP-OES-MS analyses.

Figure 7. Y/Ho vs. La/Ho in bulk samples of fluorite. Two samples are clearly contaminated by carbonates (see Table 2).
Table 1 - Major (wt.%) and minor (ppm) elements compositions of Silius fluorite (ICP-OES-MS on bulk sample).

Table 2 - Major (wt.%) and minor (ppm) elements compositions of the carbonate gangue of Silius veins (ICP-OES-MS on bulk samples).

Table 3 - Chemical compositions of synchysite in the Silius veins.

Table 4 - Major (wt.%) and minor (ppm) elements compositions of Silius host rocks (ICP-OES-MS on bulk samples).

Table A1 - Description of the analyzed samples from the Silius vein system.

Table A2 - Chemical composition of standards used for WDS analysis of synchysite-(Ce).