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1 Introduction

Rare Earth Elements (REE) are lithophile, form deposits associated with many types of rocks and concentrate preferentially in carbonatites: 5 of the top 10 global REE deposits in exploitation are associated with carbonatites. Thus, carbonatites are the main target in the race to discover new REE reserves which has begun in 2009, when China, the largest producer in the world, announced that would only produce for its domestic market (Stone, 2009; Service, 2010).

The diversity of REE minerals is very large, but few minerals can be processed for the extraction of these elements at low costs. Mineralogy is the most critical problem in the evaluation of the economic viability of a deposit and fluorcarbonates are the most important...
REE ore minerals (Mariano & Mariano, 2012). Fluorcarbonates are exploited in most known mines, such as Bayan Obo (China), where bastnäsite and synchysite are associated with carbonated rocks of unclear origin (Kynicky et al., 2012), and in many REE deposits outside of China, including Mountain Pass (USA), where magmatic bastnäsite is exploited from a carbonatite (Mariano & Mariano, 2012). In most carbonatites, the REE are concentrated in other minerals. In some cases of exploitation, REE are the main product, such as Mount Weld (Australia), where monazite is exploited in lateritic crust (Lottermoser & England, 1988; Lottermoser, 1990), and Kola (Russia), where REE are exploited from a mckelveyite carbohydrothermal deposit (Chakhmouradian & Zaitev, 2012).

In Brazilian carbonatites, REE are (or will be) exploited as byproducts of phosphate, Ti or Nb ores, as in Catalão I (monazite, gorceixite and florencite), Araxá (burbankite, carbocermaite, ancylite, huanghoite and monazite), Serra Negra (rabadofane, florencite and crandallite) and Seis Lagos (Nb-rutile and florencite) (Lapido-Loureiro, 1994; Cordeiro et al., 2011; Rocio et al., 2012). In Brazil, there are many other carbonatites with anomalous concentrations of REE, but in only two of them are the REE concentrated as fluorcarbonates. The first is Barra do Itapirapuã, where the hydrothermal REE mineralization of bastnäsite, parsite and synchysite was studied by Ruberti et al. (2008). The second is Fazenda Varela (Fig. 1), where only petrological studies were carried out (Scheibe, 1979; Scheibe & Formoso, 1982; Menegotto & Formoso, 1998) and the synchysite mineralization was interpreted as being magmatic.

This paper presents the first results of studies on the REE mineralization associated to the Fazenda Varela carbonatite. This research represents a rare opportunity to study mineralization in a carbonatite dome intruded into sedimentsal rocks. The investigations revealed an original type of REE mineralization associated to carbonatite, represented by hydrothermal parsite-(Ce) without fluorite. The study of parsite-(Ce), in addition to the economic interests, represents a contribution to the mineralogical study of fluorcarbonates.

Figure 1. Geological map of the Lages alkaline district (modified from Scheibe, 1979 and Dani, 1998) and location of Fazenda Varela carbonatite, indicated by the arrow.
2 General concepts on REE Fluorcarbonates

Parasite was first described in a Colombian emerald mine and occurs frequently in hydrothermal veins with calcite. In Brazil, this mineral was found in carbonatite (Ruberti et al., 2008) and as a secondary mineral in granites (Botelho & Teixeira, 1995; Pereira et al., 2008).

Parasite belongs to the REE fluorcarbonates group also known as the bastnäsite series (Donnay & Donnay, 1953). The group consists of four minerals: bastnäsite (CeFCO₃), synchysite (CeFCO₃ • CaCO₃), parasite (2CeFCO₃ • CaCO₃), and röntgenite (3CeFCO₃ • 2CaCO₃). Bastnäsite accounts for approximately 90% of the world’s REE production; synchysite occurs subordinately and is associated with bastnäsite. In almost all the cases where bastnäsite is exploited, parasite and röntgenite are comparatively rare. The general mineral formula for the group is nXYCO₃ • mCaCO₃, where X = LREE, Y = (F, OH); m = 0 (bastnäsite) or 1 (synchysite, parasite, röntgenite), n = 1 (bastnäsite, synchysite), 2 (parasite) or 3 (röntgenite). Table 1 presents crystallographic and physical characteristics of these four minerals.

Table 1. Crystallographic data for the mineral series bastnäsite-synchysite (Donnay & Donnay, 1953; Wang et al., 1994).

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Crystal system</th>
<th>Cell Dimensions (Å)</th>
<th>Z</th>
<th>V (Å³)</th>
<th>ρ</th>
<th>Point Groups</th>
<th>Space Group</th>
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</thead>
<tbody>
<tr>
<td>Bastnäsite</td>
<td>hexagonal bipyramidal</td>
<td>7.16 - 9.79</td>
<td>6</td>
<td>434.65</td>
<td>5.02</td>
<td>b</td>
<td>P 2/m</td>
</tr>
<tr>
<td>Synchysite</td>
<td>monoclinic prismatic</td>
<td>12.33 7.11 18.74</td>
<td>12</td>
<td>1,602.75</td>
<td>3.97</td>
<td>3</td>
<td>C 2/c</td>
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<tr>
<td>Parasite</td>
<td>trigonal bipyramidal</td>
<td>7.12 - 14.02</td>
<td>3</td>
<td>616.24</td>
<td>4.34</td>
<td>3</td>
<td>R 3</td>
</tr>
<tr>
<td>Röntgenite</td>
<td>trigonal bipyramidal</td>
<td>7.13 - 69.4</td>
<td>9</td>
<td>3,055.41</td>
<td>4.19</td>
<td>3</td>
<td>R 3</td>
</tr>
</tbody>
</table>

As shown in table 1, only synchysite has the crystallographic a parameter that is different from the other fluorcarbonates. Wang et al. (1994) show that synchysite displays monoclinic symmetry, whereas other species of the bastnäsite group show trigonal or hexagonal symmetry. The bastnäsite minerals group present two common features: (i) the structure can be broadly described as the stacking of three types of layers (CeF, CO₃ and Ca) along the c axis; (ii) the Ce/F ratios in all the phases are 1, indicating that this feature is common in all minerals of the group. Thus, these minerals differ in the orientation of carbonate group within the carbonate layers, the order of stacking of the different type of layers and the length of the crystallographic c axis. According to Donnay & Donnay (1953), in most cases, fluorcarbonates are polycrystals with syntactical intergrowth of two species in contact along an irregular surface or along repeated parallel planes (0001). The a axes of the two species may have the same or opposite directions. In the latter case, the two species are related to each other as two individual twins with a rotation of 60 ° around the c ternary axis. All pairs have been observed, except the bastnäsite-synchysite pair.

According to Caro (1973), parasite and röntgenite contain layers of bastnäsite and synchysite in varying proportions. Van Landuyt & Amelinckx (1975) studied this group of minerals by transmission electron microscopy, revealing that the syntactic intergrowths can be described as mixtures of bastnäsite and synchysite. The authors considered bastnäsite-(Ce) and synchysite-(Ce) as two end-members and that parasite and röntgenite are ordered mixtures of bastnäsite (B) and synchysite (S) in single layers stacked along the c crystallographic axis direction. Parasite can be considered a BS stacking and röntgenite a BS₃ stacking. The structures of the members of the bastnäsite-synchysite series are shown in figure 2.

![Figure 2. Crystal structure scheme of the bastnäsite mineral series (Van Landuyt & Amelinckx, 1975).](image)

The structure of parasite could be described as an alternation of Ca, CeF and carbonate planes, all perpendicular to three-fold axis. According to the orientation of the CO₃ groups within a carbonate plane, two type of carbonate layer can be distinguished, namely e and g (Fig. 2). The planes formed by the CO₃ groups occur between the CeF and Ca layers and their internal structure alternate periodically from e to g. Therefore, the structure of the mineral may be described in terms of two basic layers (Fig. 2): CeF-e-CeF and g-Ca, both constituting the parasite units (BS). The orientation of the units stacked along the c direction and the periodic variation in the orientation of the [CO₃] groups result in complex polytypic structures. Five parasite polytypes could coexist in the same crystal and there are mixtures of structural layers and syntactical intergrowth,
constituting a polycrystal with local order and integral disorder.

The syntax among minerals of the bastnäsite group is not universal and pure phases without syntax have been found (Mariano, 1989). Syntaxial intergrowth is the most common and thirty-two types of mixed-layer components and polytypes have been observed (Van Landuyt & Amelinckx, 1975; Meng et al., 1995, 1996, 1997, 2001a, 2001b; Wu et al., 1997, 1998a, 1998b, 2000). The formulas of the mixed layer components can be expressed as $(m + n) \text{Ce} \left( \text{CO}_3 \right) \text{F}_{n} \text{CaCO}_3 (m \geq n)$, where the $m$ and $n$ values vary as presented former. The structure of parisite had not been completely determined due to its complexity (Meng et al., 2001b).

3 Study area, materials and methods

3.1 Geological Setting

The Fazenda Varela carbonatite was identified by Scheibe (1979) as part of the Lages alkaline district. This district consists of numerous bodies of ultrabasic and alkaline rocks distributed over an area of 1,200 km$^2$, called the Lages dome. This dome intrudes into the sedimentary rocks and basalts of the Paraná Basin (Fig. 1). The alkaline rocks are divided into two groups (Scheibe, 1986; Dani, 1998): leucocratic alkaline rocks (phonolite, nepheline syenite and analcime trachyte) and ultrabasic rocks (olivine and melilitite lamprophyre). K-Ar datings of the alkaline rocks yielded ages between 63 Ma and 78 Ma, and Rb-Sr dating provided ages of 81 ± 8 Ma (Scheibe et al., 1985; Scheibe, 1986).

The carbonatite occurs as veins that cut the Rio Bonito Formation, a member of Guatá Group in an area of approximately 1,500 x 900 m (Fig. 3). The Rio Bonito sediments in the studied area consist of medium sandstone composed principally of quartz and feldspar that are affected by brecciation and metasomatism related to the carbonatite intrusion.

The main outcrop occurs in the western part of the carbonatite and was mined in the 70's for soil correction. Boulders of pure carbonatite can still be encountered at this place. This carbonatite has granular texture and high porosity. A banded structure of orthoclase-rich and carbonate-rich zones associated with sedimentary rock fragments is common (Fig. 4A). According to Scheibe (1979), a drilling hole was interrupted at a depth of 100 m still within a homogeneous carbonatite.

In most outcrops, the carbonatite occurs as abundant veins vertical to subvertical, without a preferred orientation, and cut the sedimentary rock, forming breccia. The thickness of the veins ranges from few millimeters up to 50 cm (Fig. 4B). The breccia consists of quartz-feldspathic sandstone that was affected by silicification and carbonatation. The pure carbonatite and the carbonatite veins associated with the breccia are cut by small veins filled with calcite and/or quartz. Chronological relations provide evidence that the first events were the carbonatite intrusion and the transformation of the country rock to the metasomatized breccia, followed by the formation of veins with a predominance of calcite and quartz and the development of the siliceous venular system.
3.2 Geochemical data

The Fazenda Varela carbonatite is classified as ferrocarbonatite in the Woolley & Kempe (1989) diagram. According to Scheibe (1979), there are at least two phases of rhombohedral carbonate. Microprobe analyses made by Menegotto & Formoso (1998) show that the most abundant carbonate is Fe-dolomite. According to Scheibe (1979), the mineralogical composition (ankerite, barite and REE fluorcarbonate) associated with the high concentrations of REE and the high Ba/Sr, Nb/Ta and Th/U ratios indicate that carbonatite was formed in the final stages of a fractionation process. The high LREE concentration relative to the HREE in the Fazenda Varela analyses is characteristic of carbonatite rocks whatever of their origin (partial melting of upper mantle, fractional crystallization or immiscibility). According to Scheibe (1979), the Sr\(^{87}/\)Sr\(^{86}\) ratio (0.71) may indicate crustal contamination or derivation from a mantle with high initial Sr\(^{87}/\)Sr\(^{86}\) ratio, but further analyses are required to confirm this interpretation. Based on Taylor et al. (1967) and Deines & Gold (1973), Scheibe (1979) also concluded that Fazenda Varela C and O isotope ratios indicate that this carbonatite formed in a subvolcanic environment.

3.3 Methods

Seventh rock samples were collected in the western part of the carbonatite body (Fig 3). Optical microscopy of thin sections was used to identify different paragenesis. The same samples were also analyzed by X-ray diffractometry, energy dispersion spectroscopy (EDS) and electron microprobe. The X-ray diffractometry were performed using Siemens D-5000 equipment, with Cu-K\(\alpha\) radiation and Ni filter, in the range of 2\(°\) and 80\(°\), with velocity of 2\(°\)/20 per second. A concentrate of heavy minerals (sample CFV16) containing parsite, barite and apatite was also analyzed by this technique. This concentrate was obtained using diiodomethane (d = 3.325) and the Frantz isodynamic magnetic separator (15\(°\) of lateral inclination and 25\(°\) of vertical inclination, with a current of 0.4A). EDS analyses were made using a JEOL-JSM5800 scanning electron microscope (SEM) with a voltage of 20 kV and spot size of 5 \(\mu\)m. These analyses were performed at Centro de Microscopia Eletrônica (CME-UFRGS). The electron microprobe data were obtained in thin sections (CFV16 and CFV17 samples) at the CNRS/Université d’Orléans ISTO Laboratory (France) with a Cameca SX50 device, using a voltage of 15kV with spot size of 1 \(\mu\)m.

The whole rock analyzes, using ICP-MS, were performed at ACME Analytical Laboratories Ltda. (Canada). The analytical data were processed using Geoplot, Excel and Minfile programs. As there are many geochemical data on carbonatites in the literature, we analyzed three representative samples (CFV14, CFV16 and CFV17), only for our control. In addition, 14 samples of the homogeneous breccia (without carbonatite veins) were analyzed in order to investigate their potential as REE ore. Representative breccias analyzes (BFV02, BFV05, BFV06, BFV07, BFV10 e BFV15) are presented in table 2.

4 Results

4.1 Mineralogy and REE contents of Fazenda Varela carbonatite and associated breccia

The carbonatite, even in the veins of small thickness, has a granular texture that essentially consists of carbonates that crystallize in two stages. The carbonates from the first stage are magmatic and form rhombohedral grains, with approximately 0.4mm (Fig. 4C-D). These grains are zoned and their inner bands are more ferrous (ankerite and Fe-dolomite) than those on the edge (Fe-dolomite, dolomite or, subordinately, calcite). Fe and Mn bearing impurities highlight the carbonate zonation and clouded their core crystals. Orthoclase, pyrite, and quartz are also magmatic minerals from the first stage. Orthoclase is often euhedral or subhedral (Fig. 4D), rarely twinned, with dimensions between 0.1 and 0.4 mm. Orthoclase is included in the carbonates of the two generations (Fig. 4C-4D) and is frequently hydrothermalized, sometimes rounded and with corroded edges. Pyrite forms euhedral crystals with 0.5 to 2 mm, which are often pseudomorphosed by iron oxide. The quartz crystals occur principally in cavities or veins associated with apatite and carbonate from the second stage, but sometimes they are included in the carbonate from the first stage. Rare crystals of pyrochlore and monazite were observed with SEM.

The carbonatite was strongly affected by late- and post-magmatic processes. Fluids percolated through the fractures and grain boundaries, forming hydrothermal domains constituted by carbonates of the second generation, barite, quartz, and apatite, all of which have thin size. The second generation of carbonate is composed of calcite, Fe-dolomite and parsite-(Ce). The first two occur (a) as small rhombohedral crystals in the interstices between the carbonate crystals of the first stage (Fig. 4D), (b) on the rims of the carbonate crystals of the first stage (Fig. 4D), and (c) in the hydrothermal domains (Fig. 4C, between the ankerite crystals). In this case, the second stage minerals display irregular contours and contain many mineral inclusions. The parsite-(Ce), described by Scheibe (1979) as synchysite, was the last carbonate to crystallize. Its crystals are light brown in natural light and present bi-refringence of the third order. They are better developed than the other two carbonates and are frequently 0.4mm in size. They occur with fibrous habits and elongated forms (Fig. 4C), crystallized either partially following the boundaries of the ankerite crystals and partially penetrating into the matrix of hydrothermal minerals. They can be also totally in the hydrothermal...
matrix, where they often have fibroradiated habits and seem to replace and/or encompass the thinner sized minerals. These characteristics, which are related to the pervasive hydrothermalism, covering/replacing various minerals, can be clearly observed in the SEM images (Fig. 4G). The parisite-(Ce) also occurs in veins with calcite and hematite, which cut the carbonateir-regularly.

Barite (Fig. 4C) occurs as anhedral crystals that are often rounded, with sizes ranging from 0.05 to 100 mm. Apatite forms a crypto- to microcrystalline mass, with euhedral or acicular crystals with a size of approximately 0.01 mm, which are frequently associated with anhedral crystal of hydrothermal quartz filling veins. It is noteworthy that EDS analyses show the presence of chlorine in apatite. Rare crystals of aluminum strontium phosphate (goyasite?) were observed in the SEM analyses. This phase probably belongs to the hydrothermal stage.

In metasomatized breccia, even in the most homogenous parts (i.e. without carbonatite veins), it is possible observe an intense silicification (Fig. 4F), which sometimes completely erases (Fig. 4E) the grain boundaries of the quartz and feldspar crystals of the original sedimentary rock. Well-developed carbonate crystals from the first stage (Fig. 4E) that displays zones well-marked by impurities than those observed in carbonate samples, are commonly observed in the siliceous mass. As well as the carbonate, the breccia is cut by hydrothermal veins with calcite, hematite, and parisite-(Ce).

The sample CFV17 (Tab. 2) displays a REE pattern distribution (Fig. 5) typical of carbonate and characterized by high LREE contents and strong fractionation (LaN/YbN= 1.343). The samples CFV14 and CFV16 are also rich in LREE, but have much lower fractionation (LaN/YbN = 72 and 22, respectively). These two samples have relatively flat HREE distribution pattern, and GdN/YbN = 2.02 (CFV14) and 2.22 (CFV16). The origin of anomalous concentrations of P and HREE of these samples (Tab. 2) are most likely related to the presence of hydrothermal apatite.

Table 2 shows representative analyses of the metasomatized breccia (BFV02, BFV05, BFV06, BFV07, BFV10 and BFV15). Four samples have REE + Y ranging from 2,486ppm to 4,139ppm, whereas two samples show low values (506 and 696ppm). As well as in the carbonatite, La and Ce are the most abundant REE in the breccia. The concentrations of HREE and Y are very low, which is corroborated by the petrographic evidence (Fig. 4F) that the same hydrothermal fluid that concentrates REE in carbonatite formed the breccia REE-mineralization.

Table 2. Whole rock chemical analyses (wt %) of Fazenda Varela carbonatite (CFV14, CFV16 and CFV17) and associated metasomatized breccia (BFV02, BFV05, BFV06, BFV07, BFV10, BFV15).
Figure 4. Fazenda Varela carbonatite: (A) Banded carbonatite sample; (B) Metasomatized breccia with portions of silicified sedimentary rock (white) and carbonatite matrix (brown); (C) Photomicrograph of carbonatite (TLCP) with rhombohedral crystals of magmatic ankerite. The fine minerals in the central part are carbonate from the second generation, barite and parisite-(Ce) with fibrous habit. All have hydrothermal origin, and parisite-(Ce) was formed on the boundary of ankerite grain; (D) Photomicrograph of carbonatite (TLCP). Note on the left side, two magmatic carbonate crystals (cbn 1) and hydrothermal interstitial carbonate (cbn 2). The central and right side is a "hydrothermal domain", note the fibroradiated parisite-(Ce) and the subhedral crystals of magmatic orthoclase included in hydrothermal minerals; (E) Photomicrograph of metasomatized breccia (TPPL) with zoned rhombohedral carbonate showing Fe-oxides inclusions highlighting the growth bands; (F) Photomicrograph of metasomatized breccia (TLCP) cut by hydrothermal veinlet fills with calcite, hematite and parisite-(Ce); (G) SEM image showing a hydrothermal domain with parisite-(Ce); (H) SEM image of banded parisite-(Ce) with syntaxial intergrowth between phases richer in Ca (gray band) and REE (white band). Abbreviations: Ank: ankerite, Brt: barite, Cbn1: rhombohedral magmatic carbonate Cbn 2: hydrothermal carbonate, Or: orthoclase, Par: parisite-(Ce) and Qtz: quartz. TPPL: transmitted plane-polarized light, TLCP: transmitted light with crossed polars.
4.2 Study of the REE florcarbonate of Fazenda Varela carbonatite

The X-ray diffraction analyses performed on heavy minerals (Tab. 3) present parsite and synchysite peaks. The SEM images of parsite-(Ce) (Fig. 4H) show that its crystals have gray and white phases that are oriented approximately in parallel without regular distribution patterns. Gray phases are dominant but the proportion of white and gray phases varies with magnification (Fig. 4H and Fig. 6). The chemical profiles (Fig. 6) show that the white phases are richer in REE and poorer in Ca in relation to the gray phases. In the white bands, Ca ranges from 3.45 % to 19.67 % (average 12.78 %), and the REE content varies from 80.33 to 96.55 % (average 87.22 %). In the gray bands, the Ca content varies from 15.40 to 35.81 % (average 21.12 %), and the REE vary from 64.19 % to 84.60 % (average 78.88 %). Due to semi-quantitative character of these analyses, the given contents have just an indicative value, but these variations are significant. However, according to the relative proportions between Ca and REE, the white bands tend to have parsite compositions, and the gray bands tend to have intermediate compositions between parsite and synchysite. A few analyses display very low Ca concentrations, indicating a composition similar to that of bastnäsite.

As the banding and compositional variation exist in different scales, the electron microprobe analyses represent an average of the bands compositions under the electron beam spot size. To verify the differences in crystal compositions, cross-sections analyses were performed. Depending on the crystal dimensions each section is composed by 30 to 60 analyses. Along the profiles, variations in the composition were observed. However, these variations are generally small and there were no systematic differences between the edge and core of a crystal. The same was true among crystals of the same sample and among crystals from different samples. Table 4 shows representative analyses of the parsite-(Ce). They are consistent with those given by Ni et al. (2000).
As the composition is similar with that of parisite, the structural formulas were calculated based on three cations. The CO\(_2\) was calculated based on the charge balance. In all the analyses, Ce was the dominant cation, thus requiring the use of the suffix "-\(\text{Ce}\)" (Bayliss & Levinson, 1988). The preferential order for REE incorporation was Ce > La > Nd >>HREE ~ Y. The concentration of the LREE (La, Sm) ranged from 45 to 52.5 wt%. An ideal parisite-(Ce, La) has a LREE concentration of ~ 60 wt%. The value of REE + Y in the formula was generally between 1.50 and 1.60, lower than the ideal value (REE + Y = 2).

Calcium was the dominant divalent cation, and its value in the structural formula exceeded the expected value (Ca = 1) in practically all analyses. Thus, the analyses with lower Ca values tended to have higher REE values. However, there was no statistical correlation between Ca and REE. Therefore it is not possible to interpret the occurrence of substitution of REE by Ca. Ba and Sr occurred in small quantities, and the F concentrations varied widely, from 3.79 to 6.06 wt%, corresponding to a structural formula of 1.03 to 1.65 (the value in the ideal formula is F = 2).

The distribution pattern of REE in parisite-(Ce) (Fig. 5), although increased by a factor of approximately 100 times, was very similar to the REE pattern of the carbonatite, especially for the LREE, which are the main REE in parisite and carbonatites. In parisite with higher HREE concentrations, the distribution pattern of the HREE was very similar to that of the carbonatite.

### Table 4. Chemical composition (wt %) and structural formulas of parisite-(Ce).

<table>
<thead>
<tr>
<th></th>
<th>Parisite 1</th>
<th>1a</th>
<th>1b</th>
<th>1c</th>
<th>Parisite 2</th>
<th>2a</th>
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*CO\(_2\) calculated

In figure 7, the fluorcarbonate from Fazenda Varela is compared with the parisite from Muso, Colombia (Ni et al., 2000), synchysite from the Main Sayan Fault in Russia (Savel’eva & Karmanov, 2008), ceric synchysite from Markersbach, Germany ( Förster, 2001) and hydrothermal parisite and synchysite from Barra do Itapirapuã, Brazil (Ruberti et al., 2008). This comparison highlights the similarity of the Fazenda Varela parisite to other natural parisisites, as well as its richness in Ce.
Figure 7. Ca-Ce-La (wt%) data, comparing the Fazenda Varela parsite with Ni et al. (2000) parsites; Savel’eva & Karmanov (2008) syn-
chysites; Förster (2001) ceric synchysites and with the hydrothermal parsite and synchysite from Barra do Itapirapuã (Ruberti et al.,
2008).

5 Discussion

The most important Fazenda Varela parsite-(Ce) feature is the variation in its chemical composition, notably by Ca and REE. These variations have been described for bastnäsite, synchysite and parsite and are associated with the hydrothermal carbonatite mineralization of Barra do Itapirapuã (Ruberti et al., 2008). These authors found that these minerals are composed of mixed layers with different proportions of bastnäsite (CeFCO$_3$) and synchysite (CeFCO$_3$ • CaCO$_3$). In Fa-
zenda Varela, the EDS analyses confirmed the existence of syntactic growth, but with some characteristics distinct from those described by Ruberti et al. (2008), such as the existence of bands that repeats with every increase in the observed SEM scale. Thus, there are not recognizable homogeneous phases. The existence of the bastnäsite component (a mineral without Ca) indicates a composition with Ca <1, which does not occur; instead, there is a tendency of excess Ca in the mineral. A bastnäsite component also indicates the tendency of increased contents of REE in respect to parsite or syn-
chysite, which also does not occur; the REE contents are lower than expected. The contents of the main cations range from a composition (Ca = 0.96; REE = 1.79) close to an ideal parsite (Ca = 1; REE = 2) to an intermediate composition between parsite and synchysite. Thus, emphasizing the coherence among the XRD, EDS and electron microprobe data, the mineral is classified as parsite-(Ce) with particular characteristics represented by an excess of Ca and deficiencies of REE (related to syntactic growth with synchysite) and F.

Carbonatites are typically anomalous REE-bearing rocks, with preponderant LREE. Composition characterized by relative high LREE content and trace quantities of HREE and Y occur: in early crystallized, independent REE minerals; in minerals originating from hydrother-
mal events; and, less frequently, in supergene products of weathering. In many cases, the distinction between what constitute primary mineralization and hydrothermal REE mineralization in carbonatites is debatable (Mariano & Mariano, 2012), such as in Mountain Pass, where parsite is interpreted as hydrothermal (Chakhmournadian & Wall, 2012). In cases where the mineralization is well characterized as hydrothermal, such as in Barra do Itapirapuã (Ruberti et al., 2008), the REE fluorcarbonates paragenesis is more complex and occurs with associated fluorite. There is no record in the literature of REE mineralization associated with carbonatite solely with parsite, whatever the origin of this mineral i.e. magmatic or hydrothermal. The only case of REE hydrothermal mineralization where parsite is the main REE mineral is Snowbird mine (USA), where it is a byproduct of exploitation of hydrothermal fluorite veins related to granite (Metz et al., 1985; Mariano & Mariano, 2012). Fazenda Varela is, therefore, a special case of hydrothermal REE mineralization associated with carbonatite. The characteristics found in Fazenda Varela parsite-(Ce) certainly reflect the conditions of a particular environment.

Some classical concepts about the behavior of REE in a hydrothermal environment were corroborated by thermodynamic calculations; however, these calculations did not take into account changes in the dielectric
behavior of water as a function of temperature. Because there is commonly a close association between hydrothermal REE mineralization and fluorite, one of the concepts is that in environments in which the REE are concentrated to economic or potentially economic levels, they are invariably transported as fluoride complexes. Another concept is the lesser LREE mobility relative to HREE. These ideas are being reviewed by experimental studies on the speciation of REE in fluoride-and chlorine-bearing aqueous solutions in temperatures up to 300°C (Williams-Jones et al., 2012). These experiments established that: (i) the LREE are more hydrothermally mobile than the HREE, (ii) the REE form stable complexes with fluorine and chlorine ions at high temperatures and (iii) the LREE complexes with these ligands are more stable than their HREE equivalents. The same authors reproduced the ore deposition conditions in systems where aliquots of REE-bearing fluid (with 10 wt% NaCl, pH 4.5, cooling from 400° to 200°C) passed through and reacted with a phosphorous-bearing nepheline syenite. As the reaction starts, monazite, containing equal proportions of each REE, precipitated. With progressive fluid-rock interaction, the pH buffering capacity of the rock decreased. Consequently, the LREE were leached and precipitated farther along the path of the fluid at lower temperatures.

Textural evidences and parasite-(Ce) chemical composition, with respect to F and Ca, indicate that this mineral crystallized in an advanced stage of hydrothermalism from a fluid with low F activity and high Ca and CO$_3^{2-}$ activities. Because of the late character, it is not expected that this fluid matches to the residual magmatic volatiles without any or with very little interaction with the rock. Field and petrographic data evidences show that the mineralizing fluid is late and have interacted intensively with the rock during the rise. This interaction should have resulted in preferential partitioning of LREE for the fluid in the form of chloride complex. The parasite deposition may have occurred through the following reaction (1):

$$Ca^{2+} + 2REECl^{2-} + 2HF + 2HCO_3^- = CaREE_2(CO_3)_2F_2 + 2H^+ + 2Cl^-$$  \hspace{1cm} (1)

Such reaction is favored by pH and HCO$_3^-$ activity increase and a decrease in the Cl activity. In Bayan Obo, the bastnäsite deposition was induced by the dissolution of dolomite (Williams-Jones et al., 2012). By analogy, in Fazenda Varela, a similar mechanism could be the dissolution of ankerite by the following reaction (2):

$$2H^+ + CaFe(CO_3)_2 = Ca^{2+} + Fe^{2+} + 2HCO_3^-$$  \hspace{1cm} (2)

Alternatively, the mechanism could be the result of a pH increase by an alkaline water supply obtained from the enclosing sedimentary rock. It is a very plausible hypothesis given the geological context and the fact that the parasite-(Ce) formation occurred at a late stage of hydrothermalism, when external fluid input could be very effective.

6 Conclusions

The Fazenda Varela ferrocarbonatite has ankerite and Fe-dolomite as its main magmatic minerals; orthoclase, pyrite and quartz occur subordinately. The apatite concentrations are observed in some samples, but require further study. This ferrocarbonatite had been strongly affected by late- and post-magmatic hydrothermal events. The fluids percolated through fractures and grain boundaries and formed domains composed of barite, apatite, quartz, calcite, ferrodolomite and parasite-(Ce). The parasite-(Ce) is composed of well-developed fibrous to fibroradiated crystals that occur into the hydrothermal domain.

The REE mineralization (Ce and La) in carbonatite is hydrothermal and consists solely of disseminated parasite-(Ce), without associated fluorite. The country rock is a feldspathic quartz sandstone from the Rio Bonito Formation that was strongly affected by brecciation and metasomatism due to the carbonate intrusion. In the portions closest to the intrusion, the mineralization consists of the ferrocarbonatite that constitutes the breccia matrix. In the more distant parts, without carbonatite veins/matrix, the mineralization occurs as hydrothermal veins of calcite, hematite and parasite-(Ce), and most of the samples display REE concentrations in the range of 2,500 to 4,200 ppm.

The Fazenda Varela parasite-(Ce) shows an excess of Ca and a deficiency of REE relative to an ideal composition due to the growth of syntaxial synchysite. The REE pattern of Fazenda Varela parasite-(Ce) is similar to the Fazenda Varela ferrocarbonatite. This mineral was formed in an environment with low F activity and high Ca and CO$_3^{2-}$ activities from hydrothermal fluid rising, which interacted strongly with carbonatite, resulting in the preferential partitioning of LREE probably as chloride complexes. The hydrothermal mineral crystallization may have occurred due to increase activity of HCO$_3^-$ by the dissolution of magmatic carbonate or by the pH increase due to the water supply from the quartz-feldspar host rock.

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