1	Parisite-(La), ideally $CaLa_2(CO_3)_3F_2$, a new mineral from
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ABSTRACT

41 Parisite-(La) (IMA number 2016-031), ideally $CaLa_2(CO_3)_3F_2$, occurs in a hydrothermal 42 vein crosscutting a metarhyolite of the Rio dos Remédios Group, at the Mula mine, Tapera 43 village, Novo Horizonte county, Bahia, Brazil, associated with hematite, rutile, almeidaite, 44 fluocerite-(Ce), brockite, monazite-(La), rhabdophane-(La), and bastnäsite-(La). Parisite-(La) 45 occurs as residual nuclei (up to 5 mm) in steep doubly-terminated pseudo-hexagonal pyramidal 46 crystals (up to 8.2 cm). Parisite-(La) is transparent, yellow-green to white, with white streak, and 47 displays a vitreous (when yellow-green) to dull (when white) lustre. Cleavage is distinct on 48 pseudo-{001}; fracture is laminated, conchoidal, or uneven. The Mohs hardness is 4 to 5, and it is brittle. Calculated density is 4.273 g cm⁻³. Parisite-(La) is pseudo-uniaxial (+), $\omega = 1.670(2)$, $\varepsilon =$ 49 1.782(5) (589 nm). The empirical formula normalized on the basis of 11 (O + F) pfu is 50 51 $Ca_{0.98}(La_{0.83}Nd_{0.51}Ce_{0.37}Pr_{0.16}Sm_{0.04}Y_{0.03})_{\Sigma=1.94}C_{3.03}O_{8.91}F_{2.09}$. The IR spectrum confirms the 52 absence of OH groups. Single-crystal X-ray studies gave the following results: monoclinic (pseudo-trigonal), space group: C2, Cm, or C2/m, a = 12.356(1) Å, b = 7.1368(7) Å, c =53 28.299(3) Å, $\beta = 98.342(4)^{\circ}$, V = 2469.1(4) Å³, Z = 12. Parisite-(La) is the La-dominant analogue 54 55 of parisite-(Ce).

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57 Keywords: Parisite-(La), new mineral, rare-earth fluorocarbonate, Mula mine, Novo Horizonte,
58 Bahia, Brazil

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INTRODUCTION

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64 Parisite-(La) is the La-dominant analogue of parisite-(Ce). Minerals with a chemical composition consistent with that of parisite-(La) were reported in several studies from different 65 66 localities: Třebíč durbachite massif, SW Moravia, Czech Republic (Sulovský 2001); at a 67 metabauxite/marble interface in the eastern part of Samos island, Greece (Theye et al. 2003); 68 alkaline rocks in Romania (Hirtopanu 2006, Hirtopanu et al. 2015); Cerro Boggiani massif, Alto 69 Paraguay Province, Paraguay (Enrich et al. 2010); Bear Lodge carbonatite, Wyoming, USA 70 (Moore et al. 2015). Parisite-(Nd) is also reported in literature as a mineral without IMA approval 71 (Jambor et al. 1988).

Parisite-(La) has been approved as a new mineral species by the Commission of New Minerals, Nomenclature and Classification (CNMNC) of the International Mineralogical Association (IMA 2016-031). The type specimen of parisite-(La) is deposited in the mineralogical collection of the Museu de Geociências, Instituto de Geociências, Universidade de São Paulo, Rua do Lago, 562, 05508-080 - São Paulo, SP, Brazil, with the registration number DR1032 (a part of the holotype), and at the University of Arizona Mineral Museum (RRUFF Project deposition # R130687).

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OCCURRENCE

83 Parisite-(La) occurs in a late-stage hydrothermal vein crosscutting metarhyolites, at the Mula mine, Tapera village, Novo Horizonte, State of Bahia, Brazil (12°48'28"S; 42°10'04"W). 84 The metarhyolites, together with metadacites and meta-andesites, constitute the Rio dos 85 86 Remédios Group, a package of metavolcanic acid rocks formed as a result of peraluminous and 87 alkaline magmatism during a continental rift which opened ~1.75 Ga (Teixeira 2005, Martins et al. 2008). This group crops out in an area of about $35 \times 10 \text{ km}^2$ and is one of the lithologic 88 89 members of a larger package of metasedimentary and locally metavolcanic rocks called Serra do 90 Espinhaço that extends N-S for about 1200 km from the center of Minas Gerais to the northern 91 reaches of Bahia state (Figure 1). The evolution of the sedimentary basin in this region occurred 92 in several stages during the Mesoproterozoic (~1.7-1.2 Ga) and during subsequent

93 metamorphism, folding and fracturing, during the Brasiliano Cycle of the Neoproterozoic (630 -94 490 Ma, according to Pedrosa-Soares et al. 2011). During the Brasiliano event, swarms of 95 hydrothermal quartz veins were generated. These have been exploited at Novo Horizonte for the 96 production of rutilated quartz, barite and gold. Martins et al. (2008) reported muscovite ages of 97 404 and 490 Ma in altered metarhyolites close to these veins, confirming their formation during 98 the Brasiliano event. Parisite-(La) is associated with almeidaite (its type locality is also the Mula 99 mine: Menezes Filho et al. 2015), hematite, rutile, fluocerite-(Ce), brockite, monazite-(La), 100 rhabdophane-(La) and bastnäsite-(La). The pit of the Mula mine is composed of brecciated quartz 101 veins cemented by chalcedony. The veins show a stockwork pattern, which is typical of hydraulic 102 fracturing processes, and are partially kaolinized; their orientation is NNW-SSE, with a thickness 103 of up to 2 m.

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APPEARANCE and PHYSICAL PROPERTIES

108 Parisite-(La) occurs as residual nuclei (up to 5 mm) in doubly-terminated pseudo-109 hexagonal pyramidal crystals (up to 8.2 cm) with corrugated faces (Figure 2a). The crystals 110 apparently do not differ from what is usually described for parisite-(Ce): acute dipyramids with 111 horizontally striated faces, terminated by pinacoid. The pinacoid faces are cleavage planes. The 112 crystals are prismatic in appearance due to oscillatory combination of steep pyramids, but true 113 prism faces are lacking or very small. However real forms of parisite-(La) can be pedion, 114 pinacoid and sphenoid if the space group is C2, pedion, pinacoid and dome if Cm, pinacoid and 115 prism if C2/m. These crystals were partially replaced by bastnäsite-(La), monazite-(La), and 116 rhabdophane-(La). Crusts consisting of microcrystals of parisite-(La) also occur (Figure 2b).

Parisite-(La) is transparent, yellow-green to white, with white streak, and displays a vitreous (when yellow-green) to dull (when white) lustre. It is non-fluorescent under both shortwave (254 nm) and longwave (366 nm) ultraviolet radiation. Cleavage is distinct on pseudo- $\{001\}$ and parting was not observed; fracture is laminated, conchoidal, or uneven. The Mohs hardness is between 4 and 5, and the mineral is brittle. Density was not measured; calculated density is 4.273 g cm⁻³ using the empirical formula. In transmitted light parisite-(La) is 123 colourless, pseudo-uniaxial (+), $\omega = 1.670(2)$, $\varepsilon = 1.782(5)$ (589 nm). Pleochroism was not 124 observed.

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INFRARED and RAMAN SPECTROSCOPY

In order to obtain infrared (IR) absorption spectra, powdered samples were mixed with anhydrous KBr, pelletized, and analyzed using an ALPHA FTIR spectrometer (Bruker Optics) with the resolution of 4 cm⁻¹ in the wavenumber range from 360 to 3800 cm⁻¹; 16 scans were obtained. The IR spectrum of an analogous pellet of pure KBr was used as a reference. The assignments of IR bands have been made in accordance with Adler and Kerr (1963), White (1974), and Nakamoto (1997, 2009).

135 The IR spectrum of parisite-(La) is close to that of parisite-(Ce) (Figure 3). Weak IR bands in the range $1700-3000 \text{ cm}^{-1}$ correspond to overtones and combination modes. The other 136 bands are assigned as follows (cm^{-1} ; w – weak band, s – strong band, sh – shoulder): 1454s, 137 138 1430sh (degenerate asymmetric C-O-stretching vibrations), 1089w, 1081w (non-degenerate 139 symmetric C-O-stretching vibrations), 871s, 850sh (out-of-plane bending vibrations of the 140 carbonate ion triangles), 746w, 734w (in-plane bending vibrations of the carbonate ion triangles), 679w, 602w (presumably, overtones and/or combination modes), 368 (lattice mode involving 141 Ca–O- or *REE*–O-stretching vibrations). Bands in the range 3000-3800 cm^{-1} are not observed, 142 143 which indicates the absence of OH groups.

144 Raman spectrum of parisite-(La) (Figure 4) was collected on a randomly oriented crystal at 145 100% of 150 mW on a Thermo Almega microRaman system, using a 532 nm solid-state laser, 146 confocal Olympus optics with a 10x objective, and a thermoelectrically cooled CCD detector. The laser was partially polarized with 4-cm⁻¹ resolution and a spot size of 1 μ m. The peak at 1428 147 cm^{-1} corresponds to the v₃ asymmetric stretching mode of CO_3^{2-} anions. Symmetric C–O 148 stretching modes are represented by the bands at 1081, 1091, and 1098 cm⁻¹. The bands at 737 149 and 871 cm⁻¹ correspond to in-plane and out-of-plane vibrations of CO_3^{2-} , respectively. The 150 bands at 600 and 970 cm⁻¹ may be due to any vibrations involving F⁻ anions. Bands with 151 wavenumbers below 500 cm⁻¹ are attributed to lattice modes. The tentative assignment of Raman 152 153 bands is based on previous studies (Frost & Dickfos 2007, Guastoni et al. 2010, Frost et al. 154 2013). All peaks above 1500 cm-1 are due to fluorescence. This effect is typical for REE 155 compounds (see e. g. Betancourtt 2003) 156 157 158 CHEMICAL DATA 159 160 Chemical analyses (25) were carried out using a Cameca SX100 microprobe (WDS mode, 161 25 kV, 40 nA, beam diameter 5 µm) at the Department of Geosciences, University of Arizona, 162 Tucson, Arizona, U.S.A. H₂O was not analysed because of the absence of bands corresponding to 163 O-H vibrations in the IR spectrum. The thermogravimetric (TG) curve of parisite-(La) in 164 nitrogen atmosphere is given in Figure 5. The second step in the TG curve may be due to the 165 decomposition of an intermediate carbonate. Fluorine is not lost during the heating because CaF₂ 166 melts above 1400°C without decomposition and *REE*F₃ do not decompose into elements below 167 1200°C (Ranieri et al. 2008). The formula corresponding to CO₂ content of 23.68% obtained 168 experimentally from TG data is is not charge balanced. This weight loss is slightly less than 169 24.50 wt% CO₂ calculated for formula neutrality. Mean analytical results are given in Table 1. 170 normalized on the basis of 11 (O + F) pfuThe empirical formula is 171 $Ca_{0.98}(La_{0.83}Nd_{0.51}Ce_{0.37}Pr_{0.16}Sm_{0.04}Y_{0.03})_{\Sigma=1.94}C_{3.03}O_{8.91}F_{2.09}$, where CO₂ was constrained to 172 24.50% for charge neutrality. The idealized formula is $CaLa_2(CO_3)_3F_2$, which requires: La_2O_3 173 60.80, CaO 10.46, CO₂ 24.63, F 7.09, O=F -2.98, total 100.00wt.%. 174 An additional sample was measured at the Instituto de Geociências, Universidade Federal 175 de Minas Gerais, Belo Horizonte, Minas Gerais, Brazil. Chemical analyses (7) were carried out 176 using a Jeol JXA8900R electron microprobe (WDS mode, 15 kV, 20 nA, beam diameter 5 µm). 177 Mean analytical results are given in Table 1. The empirical formula normalized on the basis of 11 178 (O + F) pfu is Ca_{0.91}(La_{0.82}Nd_{0.47}Ce_{0.43}Pr_{0.26}Sm_{0.04}Y_{0.02})_{\Sigma=2.04}(CO₃)_{3.03}F_{1.91}, where CO₂ was 179 constrained to 24.70% for charge neutrality. 180 181 182 CRYSTALLOGRAPHY 183

184 Single-crystal X-ray diffraction studies and Convergent-Beam Electron Diffraction 185 (CBED) analysis performed under TEM indicated monoclinic symmetry (pseudo-trigonal) and 186 systematic absences compatible with space groups C2(#5), Cm(#8), C2/m (#12). Single-crystal 187 X-ray diffraction studies carried out using a Bruker APEX2 CCD automated diffractometer with 188 graphite-monochromatized MoKa ($\lambda = 0.71073$ Å) radiation gave the following results: a =12.356(1) Å, b = 7.1368(7) Å, c = 28.299(3) Å, $\beta = 98.342(4)^{\circ}$, V = 2469.1(4) Å³, Z = 12. The a: 189 190 b: c ratio calculated from the unit cell parameters is 1.7313: 1: 3.9652. Gladstone-Dale 191 compatibility is -0.046 (good) for the chemical analysis made in Arizona and -0.032 (excellent) 192 for the chemical analysis made in Minas Gerais.

We confirmed that the space groups C2/c(#15) and Cc(#9) cited by Ni et al. (2000) were not consistent for parisite-(La). The space group C2/c(#15) analysis show 816 reflections satisfying systematic absence conditions with an average I/s (I) of 2.07 (21518 reflections read) and for the space group Cc(#9), there are 789 reflections with an average I/s(I) of 2.11 at the same conditions (20612 reflections read).

198 For TEM studies, the sample was ground in an agate mortar and dispersed in a small 199 volume (1 mL) of isopropanol. The suspension was dropped onto the carbon-coated Cu-TEM 200 grid (300 mesh), and placed in a desiccator before TEM analysis. Convergent-Beam Electron 201 Diffraction (CBED) and Nano-Beam Electron Diffraction (NBD) were performed by using a 202 Tecnai G2-20 TEM (FEI), with LaB₆ filament, operated at 200 kV, at the Center of Microscopy 203 at UFMG, Belo Horizonte, Brazil. The diffraction patterns were recorded by using a side-204 mounted ES500W Erlangshen CCD camera (Gatan). The CBED patterns were performed using a 205 convergence semi-angle of 1.3 mrad, with a beam size of approximately 20 nm, and the NBD 206 conditions were achieved by slightly spreading the C2 condenser lens to obtain a nearly parallel 207 narrow beam.

In order to deduce the crystal system, the Bravais lattice, and the possible space group, electron diffraction experiments were carried out following Morniroli and Steeds (1992), Redjaïmia and Morniroli (1994), Jacob et al. (2012), and Morniroli (2013). A suitable area of the sample was illuminated by a focused narrow beam under the TEM. Observations of CBED patterns were done while the sample was being tilted around a chosen Kikuchi line, or a row containing reflections, which passed through the origin of the reciprocal lattice. Several zone axis patterns (ZAPs) were recorded in the CCD until the ZAP of highest "net"-symmetry had been found. The "net"-symmetry takes into account the position of the reflections observed in the diffraction pattern, not their intensity. It is defined as the symmetry of the spots "net" displayed in the zeroth order Laue zone (ZOLZ) and the whole pattern (WP), which includes the ZOLZ and first order Laue zone (FOLZ) (Morniroli and Steeds 1992; Jacob et al. 2012). Following Morniroli and Steeds (1992), the "net"-symmetry is here referred as "(ZOLZ)WP" symmetry – the (ZOLZ) symmetry is put in brackets henceforth in order to distinguish it from the WP's.

The identification of the monoclinic crystal system was achieved by observing the ZAP of highest "net"-symmetry shown in Figure 6. The two perpendicular axes drawn in the pattern (Figure 6) indicate 2 mirrors, and also 2-fold axes, for the spots net shown in the ZOLZ, therefore the symmetry is (2*mm*). The vertical axis shown in Figure 6, however, is not a mirror for the WP, but the horizontal one. So the "net"-symmetry observed in this ZAP is (2mm)m. Comparisons with the Atlas of Electron Diffraction Zone-Axis Pattern (Morniroli 2013) showed that the ZAP shown in Figure 6 corresponds to the [u0w] zone axis for the monoclinic system.

228 The non-primitive mS (mA or mB) Bravais lattice was deduced by comparing the features 229 displayed in the diffraction pattern shown in Figure 7 with the typical ZAPs for the monoclinic 230 crystal system (Morniroli 2013). The Figure 7.a shows the ZOLZ reflections of the either [010]_b or [001]_c zone axis, on which two symmetry axes are drawn – the subscripts b and c stand for 231 232 unique axis b and c, respectively. These non-perpendicular axes (beta* or gamma* equal 89.18°) 233 indicate the 2-fold symmetry (2) for the spots net seen in the ZOLZ. The FOLZ reflections were 234 not easily observed at first, so the electron beam was tilted along each axis until the FOLZ 235 reflection could be seen at the microscope screen as shown in Figures 7.b and 7.c. From the 236 observation of the FOLZ reflections the symmetry of the WP was taken as 2-fold. So the "net"-237 symmetry of this ZAP concluded as (2)2. This is the exact zone axis pattern required for the 238 identification of the Bravais lattice for the monoclinic system (Morniroli 2013).

The narrow parallelograms drawn in the ZOLZ and FOLZ reflections shown in Figure 7 correspond to the "unit" cell in the reciprocal space. This "unit" cell indicates the periodicity of reflections in the spots net in the diffraction pattern. As can be seen in Figure 6, there is a relative shift of the "unit" cell in the FOLZ compared to the ZOLZ. It means that the Bravais lattice of the Parisite-(La) crystal is non-primitive (Morniroli and Steeds 1992; Jacob et al. 2012). Therefore, the Bravais lattice is *mS* (*mA* or *mB*).

245 The lack of periodicity differences between ZOLZ and FOLZ reflections is an evidence 246 that there are no glide planes parallel to $[010]_b$ or $[001]_c$ (Morniroli and Steeds 1992; Jacob et al. 247 2012) as reported by Ni et al (2000) for parisite-(Ce). It was not possible to deduce the screw axis 248 perpendicular to the zone axis pattern [u0w] shown in Figure 6. The possible 2₁ screw axis 249 perpendicular to that particular zone axis could be related to the forbidden reflections in the row 250 defined by any of the symmetry axes shown in Figure 6. The forbidden nodes would appear, 251 however, due a double diffraction effect. This could be checked out by tilting the beam around 252 the symmetry axis shown in Figure 6 in order to decrease the intensity of the likely forbidden 253 reflections (Morniroli and Steeds 1992), but it has been not verified during measurements. 254 Moreover, the absence of glide planes parallel to $[010]_{\rm b}$ or $[001]_{\rm c}$ are consistent with the possible 255 extinction symbols, A1-1_b or B11-c. From the comparisons of the features observed in the 256 experimental ZAPs with the typical theoretical diffraction patterns (Morniroli 2013), the possible 257 space groups are A121 (#5), A1m1 (#8), A12/m1 (#12), B112 (#5), B11m (#8) or B112/m (#12), 258 alternative settings of the space groups C2(#5), Cm(#8), C2/m (#12).

Unfortunately, despite determining monoclinic unit-cell parameters from the singlecrystal pattern, we could not obtain single-crystal data suitable for the structure refinement. Thin inclusions are visible under the microscope. Possible they are the main cause of the absence of single crystals suitable for structural investigations. Inclusions of synchysite, röntgenite or bastnäsite microblocks are typical for parisite in general.

264 X-ray powder diffraction data (Table 2) were obtained using a Rigaku R-AXIS Rapid II 265 single-crystal diffractometer equipped with cylindrical image plate detector using Debye-Scherrer 266 geometry (d = 127.4 mm; CoK α -radiation). Parameters of monoclinic unit cell refined from 267 powder data are: a = 12.323(7), b = 7.121(2), c = 28.28(1) Å, $\beta = 98.33(4)^{\circ}$, V = 2456(3) Å³.

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DISCUSSION

The second most abundant *REE* in parisite-(La) from Mula mine is Nd, not Ce. On a chondrite-normalized plot there would be a marked negative Ce anomaly. One hypothesis about the conditions of formation could be that the fluids that transported *REE* had leached them in a mildly oxidizing environment, where Ce was partially oxidized to Ce^{4+} and thus remained

276	immobile. The resulting solution would be depleted in Ce. For the other five parisite-(La)					
277	occurrences quoted in the literature, neither crystallographic nor chemical data are presented. The					
278	only exception is a partial chemical analysis for parisite-(La) from Třebíč durbachite massif, SW					
279	Moravia, Czech Republic (Sulovský 2001): La ₂ O ₃ 28.68, Ce ₂ O ₃ 24.07, Nd ₂ O ₃ 4.01, CaO 10.22,					
280	SO ₃ 0.98, F 6.38, $O=F$ -2.67, total 71.64 wt.%. The calculated formula is					
281	$Ca_{1.01}(La_{0.97}Ce_{0.81}Nd_{0.13})_{\Sigma 1.91}(CO_3)_{2.89}(SO_4)_{0.07}F_{1.84}$. In this case Ce is the second most abundant					
282	REE and the conditions of formation differ from that of parisite-(La) from Mula mine.					
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	1					
constitu ent	wt.%	range	stand. dev.	probe standard	xtal	line
CaO	10.10	10.05-10.15	0.02	wollastonite	PET	Κα
Y_2O_3	0.52	0.47-0.55	0.02	YAG	TAP	Lα
La_2O_3	24.77	24.54-24.96	0.11	REE3	LIF	Lα
Ce ₂ O ₃	11.16	11.03-11.32	0.07	REE3	LIF	Lα
Pr ₂ O ₃	4.73	4.63-4.91	0.06	REE3	LIF	Lβ
Nd ₂ O ₃	15.82		0.09	REE2	LIF	Lβ
Sm ₂ O ₃	1.25		0.03	REE2	LIF	Lβ
Eu ₂ O ₃	0.07		0.03	REE1	LIF	Lβ
F	7.30		0.18	MgF ₂	ТАР	Κα
CO ₂	(24.50)			(calculated)*		
-O=F	- 3.07	-	-			
Total	97.15					

Table 1. Chemical data for parisite-(La).

400		2					
401 402 403	constitu ent	wt.%	range	stand. dev.	probe standard	xtal	line
404	CaO	9.45	9.28 - 9.73	0.16	wollastonite	PETJ	Κα
405 406	Y_2O_3	0.51	0.48 - 0.54	0.02	YAG	TAP	Lα
407	La ₂ O ₃	24.82	24.31 - 25.32	0.41	monazite-(Ce)	PETJ	Lα
408 409	Ce_2O_3	12.99	12.86 - 13.24	0.16	monazite-(Ce)	PETJ	Lα
410	Pr ₂ O ₃	7.95	7.27 - 8.83	0.53	monazite-(Ce)	LIF	Lβ
411 412	Nd ₂ O ₃	14.77	14.23 - 15.26	0.38	monazite-(Ce)	LIF	Lβ
413	Sm ₂ O ₃	1.24	1.22 – 1.27	0.02	monazite-(Ce)	LIF	Lβ
414 415	Eu ₂ O ₃	0.07	0.02 - 0.09	0.03	monazite-(Ce)	LIF	Lβ
416	F	6.71	6.01 - 7.58	0.48	MgF ₂	TAP	Κα
417 418	CO ₂	(24.70)	-	-	(calculated)*		
418	-O=F	- 2.82	-	-			
420 421	Total	100.39					

Analyses at the Department of Geosciences, University of Arizona, Tucson, Arizona,
 U.S.A.

- 426
- 427 *calculated from the idealized formula.
- 428 Probe standard composition:
- 429 YAG = Y : 44.93%, Al: 22.73%, O: 32.34%
- 430 REE1 = Si: 12.60%, Al: 16.15%, Ca: 17.98%, Eu: 3.8%, Gd: 3.87%, Tb: 3.78%, Tm: 3.81%, O:
- 431 38.01%
- 432 REE2 = Si: 12.65%, Al: 16.21%, Ca: 18.05%, Nd: 3.65%, Sm: 3.67%, Yb: 3.74%, Lu: 3.75%, O:
- 433 38.27%
- 434 REE3 = Si: 12.69%, Al: 16.26%, Ca: 18.1%, Y : 3.21%, La: 3.65%, Ce: 3.42%, Pr: 3.79%, O:
- 435 38.88%
- 436
- 437 438

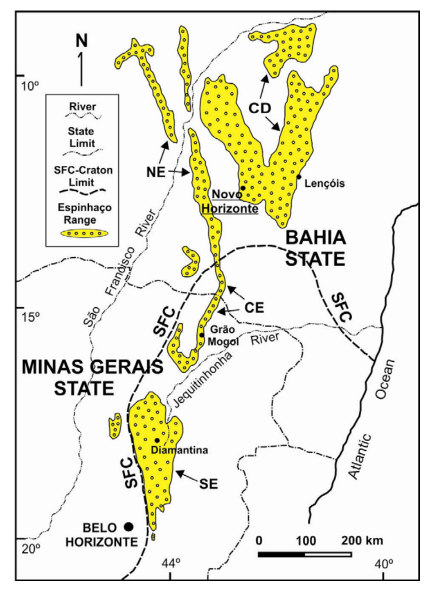
	7 9	* 8	1 1 14
I _{obs}	$d_{\rm obs}, {\rm \AA}$	d_{calc} , Å	h k l*
55	13.95	13.991	002
20	6.98	6.995	004
37	4.655	4.664	006
88	3.555	3.561, 3.558	020, -311
13	3.446	3.451, 3.449, 3.448	022, 311, -313
3	3.323	3.326, 3.325, 3.324	023, 312, -314
18	3.169	3.173, 3.172, 3.171	024, 313, -315
12	3.000	3.004, 3.003, 3.002	025, 314, -316
100	2.827	2.830, 2.829, 2.828	026, 315, -317
2	2.655	2.659, 2.658, 2.657,	027, 316, -318, 404
		2.657	
2	2.495	2.495, 2.495, 2.494	028, 317, -319
8	2.331	2.332	0.0.12
1	2.274	2.275, 2.274, 2.271	-133, -424, -229
2	2.241	2.246, 2.245	133, 422
10	2.199	2.200, 2.200, 2.199	0.2.10, 319, -3.1.11
58	2.055	2.055, 2.054	-331, -602
4	2.034	2.033, 2.032, 2.032	-333, 600, -604
6	1.971	1.972, 1.972, 1.971,	333, -335, 602, -606
		1.971	
38	1.950	1.951, 1.951, 1.950	0.2.12, 3.1.11, -3.1.13
36	1.880	1.881, 1.881, 1.880,	335, -337, 604, -608
		1.879	
9	1.780	1.780, 1.779	040, -622
2	1.767	1.772, 1.772, 1.771	337, -339, -6.0.10
1	1.749	1.749	0.0.16
3	1.725	1.725, 1.724, 1.724	044, 622, -626
23	1.663	1.663, 1.662, 1.662	046, 624, -628
3	1.570	1.570, 1.570, 1.569	0.2.16, 3.1.15, -3.1.17
12	1.542	1.542, 1.542, 1.542,	3.3.11, -3.3.13, 6.0.10, -6.0.14
		1.541	
2	1.502	1.502, 1.502, 1.501	0.4.10, 628, -6.2.12
9	1.425	1.425, 1.425, 1.424	0.2.18, 3.1.17, -3.1.19
9	1.415	1.415, 1.415, 1.414	0.4.12, 6.2.10, -6.2.14
9	1.346	1.346, 1.345, 1.345	-351, -642, -913
3	1.332	1.332, 1.332, 1.332	3.3.15, 6.0.14, -3.3.17
· · · · ·			

*The *hkl* indices are chosen taking into account intensities of reflections of the powder X-ray
diffraction pattern calculated from the structure data for monoclinic parisite-(Ce) reported by Ni
et al. (2000).

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- 445

1		
	Parisite-(La)	Parisite-(Ce)
Formula	$CaLa_2(CO_3)_3F_2$	$CaCe_2(CO_3)_3F_2$
Space group	C2, Cm, C2/m	<i>C</i> 2/ <i>c</i> or <i>Cc</i>
<i>a</i> , Å	a = 12.3563(13)	a = 12.305(2)
b, Å	b = 7.1368(7)	b = 7.1053(5)
<i>c</i> , Å	c = 28.299(3) Å	c = 28.250(5)
Z	$\beta = 98.342(4)^{\circ}$	$\beta = 98.257(14)^{\circ}$
	Z = 12	Z = 12
Strong lines of the X-	2.827 (100), 3.555	3.565 (100), 2.838
ray powder-diffraction	(88), 2.055 (58), 13.95	(100), 2.060 (80),
pattern: d , Å (I , %)	(55), 1.950 (38), 4.655	1.938 (60), 1.882 (50),
	(37), 1.880 (36)	1.658 (50), 14.03 (40)
Optical data:		
ω	1.670(2)	1.6718-1.6767
3	1.782(5)	1.7664-1.7729
Optical sign	(+)	(+)
Density,	4.331	4.3915
g·cm ⁻³		
Mohs hardness	4 - 5	4 1/2
References	This study	Cell data: Ni et al.
		(2000); XRPD:
		Cheang (1977);
		optical data, density,
		hardness: Flink (1901)

Table 3. Comparative data for parisite-(La) and parisite-(Ce).



450 Figure 1. The Espinhaço Range in Eastern Brazil (states of Bahia and Minas Gerais), showing its
451 geographic/geotectonic domains Southern Espinhaço (SE), Central Espinhaço (CE), Northern
452 Espinhaço (CE), Northern

452 Espinhaço (NE) and Chapada Diamantina (CD) in relation to the São Francisco Craton (SFC).

453 The study area near Novo Horizonte (Bahia) is underlined.



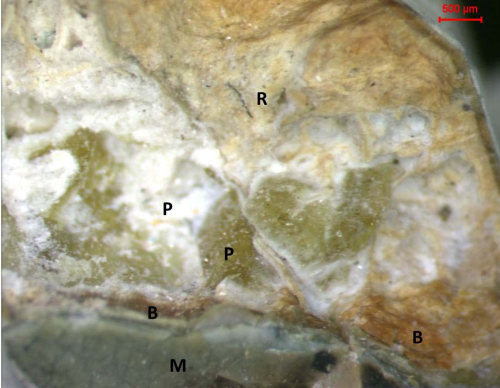
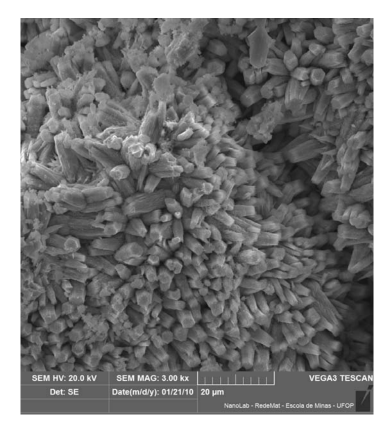


Figure 2a – Partial pseudomorphs of monazite-(La) [M], bastnäsite-(La) [B], and rhabdophane-(La) [R] after parisite-(La) [P].





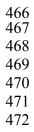
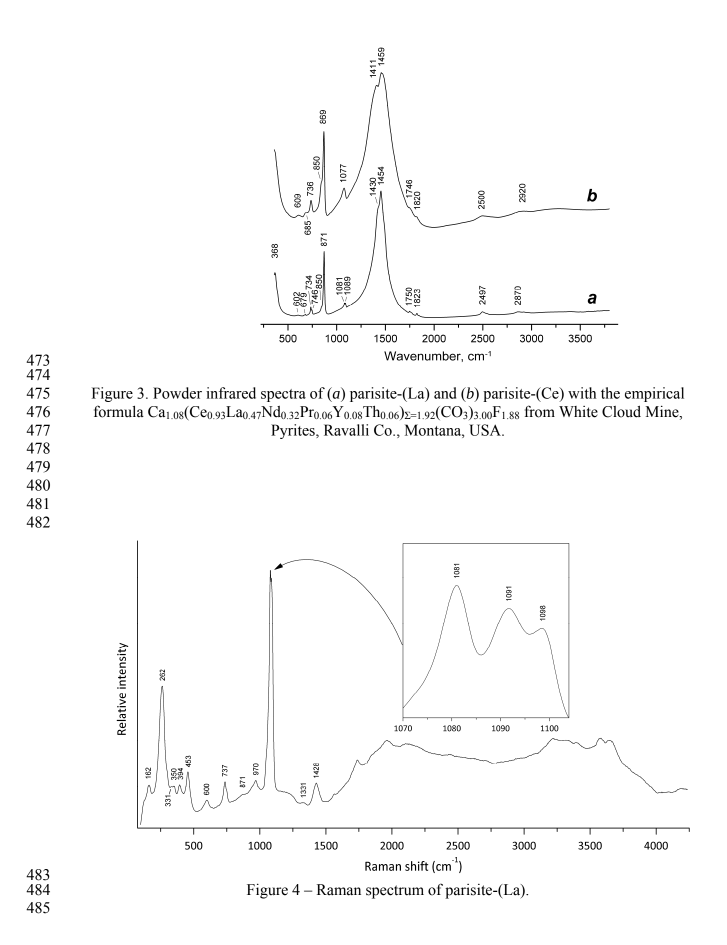


Figure 2b – Crust consisting of microcrystals of parisite-(La).



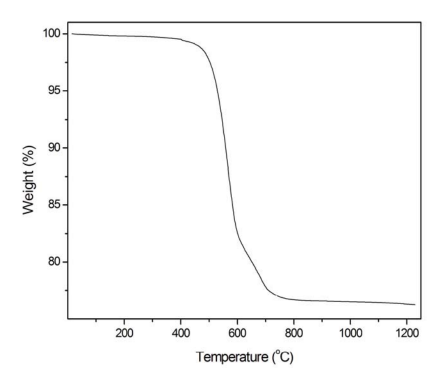
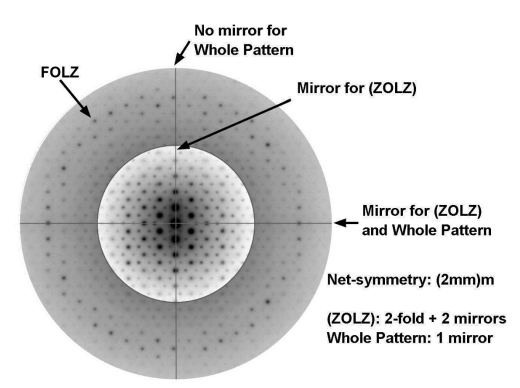
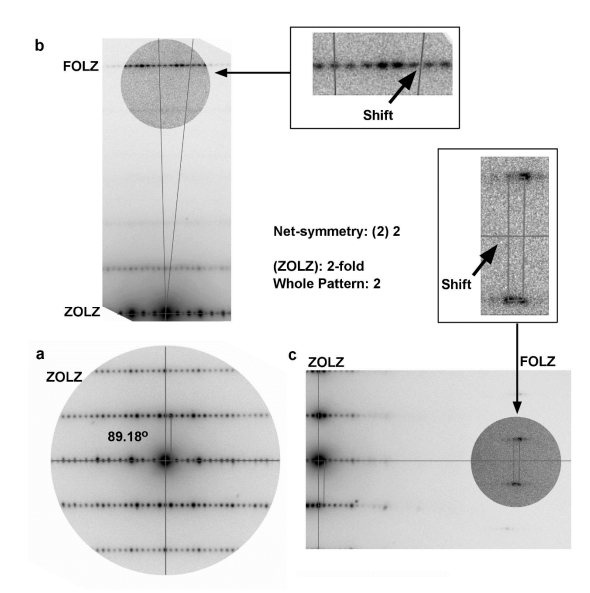


Figure 5. TG curve of parisite-(La).



492
493 Figure 6 – Electron diffraction zone axis pattern of highest "net"-symmetry found in parisite494 (La), required for the identification of the crystal system.



497 Figure 7 – Electron diffraction of the either $[010]_b$ or $[001]_c$ zone axis required for the identification of the Bravais lattice for the monoclinic crystal system. (a) Only the ZOLZ 498 499 reflections are visible. The lines indicate the symmetry axis in the pattern. (b-c) Both ZOLZ and FOLZ reflections are visible after the electron beam had been tilted along each axis shown in (a). 500 501 The narrow parallelogram drawn in each figure corresponds to the "unit" cell in reciprocal space 502 in the ZOLZ and FOLZ. The relative shift of the "unit" cell in the FOLZ, along each symmetry 503 axis, compared to the ZOLZ is an evidence for a non-primitive crystal system. There is no 504 periodicity difference between ZOLZ and FOLZ reflections.

505