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Distribution of REE-bearing minerals in felsic magmatic rocks and paleosols from Gran Canaria, Spain: intraplate oceanic islands as a new example of potential, non-conventional sources of rare-earth elements

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#### ABSTRACT

Gran Canaria is a hotspot-derived, intraplate, oceanic island, comprising a variety of alkaline felsic magmatic rocks (i.e. phonolites, trachytes, rhyolites and syenites). These rocks are enriched in rare-earth elements (REE) in relation to the mean concentration in the Earth's crust and they are subsequently mobilised and redistributed in the soil profile. From a set of 57 samples of felsic rocks and 12 samples from three paleosol profiles, we assess the concentration and mobility of REE. In the saprolite that developed over the rhyolites, we identified REE-bearing minerals such as primary monazite-(Ce), as well as secondary phases associated with the edaphic weathering, such as rhabdophane-(Ce) and LREE oxides. The averaged concentration of REE in the alkaline bedrock varies from trachytes (449 mg kg<sup>-1</sup>), to rhyolites (588 mg kg<sup>-1</sup>) and to phonolites (1036 mg kg<sup>-1</sup>). REE are slightly enriched in saprolites developed on trachyte (498 mg kg<sup>-1</sup>), rhyolite (601 mg kg<sup>-1</sup>) and phonolite (1171 mg kg<sup>-1</sup>) bedrocks. However, B-horizons of paleosols from trachytes and phonolites showed REE depletion (436 and 994 mg kg<sup>-1</sup>, respectively), whereas a marked enrichment was found in soils developed on rhyolites (1584 mg kg<sup>-1</sup>).

According to our results, REE resources on Gran Canaria are significant, especially in Miocene alkaline felsic magmatic rocks (declining stage) and their associated paleosols. We estimate a total material volume of approximately 1000 km<sup>3</sup> with REE concentrations of 672  $\pm$ 296 mg kg<sup>-1</sup>, yttrium contents of 57  $\pm$ 30 mg kg<sup>-1</sup>, and light and heavy REE ratios (LREE/HREE) of 17  $\pm$ 6. This mineralisation can be considered as bulk tonnage and low-grade ore REE deposits but it remains necessary to develop detailed mineral exploration on selected insular zones in the future, without undermining environmental and socioeconomic interests.

**Keywords:** REE exploration; REE-bearing minerals; weathering processes; ocean intraplate volcanic island; Gran Canaria

#### Highlights:

- Felsic magmatic rocks from Gran Canaria island are potential low-grade REE ores.
- Primary REE-bearing minerals: monazite-(Ce).
- REE-bearing weathering phases: rhabdophane-(Ce), LREE oxides.
- Secondary weathering REE-bearing phases linked to goethite and hematite.

#### **1. Introduction**

According to the Union of Pure and Applied Chemistry (IUPAC), the elements in the REE group are Sc, Y and the 15 elements from the lanthanides group, i.e. La, Ce, Pr, Nd, Pm, Sm, Eu, Gd, Tb, Dy, Ho, Er, Tm, Yb and Lu (Connelly et al., 2005). REE are not as rare as the group name implies. The term "rare" is used because REE are very difficult to find in high enough concentrations for economic extraction. In the Earth's crust, the average concentration of REE ranges from 150 to 220 mg kg<sup>-1</sup> (Rudnick and Gao, 2003), exceeding that of many other metals such as copper (55 mg kg<sup>-1</sup>) and zinc (70 mg kg<sup>-1</sup>), and they are intensely exploited by industrial mining (Long et al., 2010).

REE underpin a broad range of emerging technologies within state-of-the-art green energies, such as efficiency enhancement of solar cells and artificial photosynthesis, based on H<sub>2</sub> generation (Wondraczek et al., 2015). Indeed, REE occupy a very relevant

position (holding the highest supply risk rate) in the list of Critical Raw Materials defined by the European Commission (2014), this includes REE and geological materials with a high economic relevance coupled with a high risk of supply.

The extreme supply risk of REE derives from the fact that China monopolises REE production worldwide (98% of global production in 2010 and around 85% at present) (Gambogi, 2017). Evidence of this is the global alarm in 2010, when the Chinese government drastically reduced export quotas, thus revealing the high vulnerability of the REE global market. Considering their critical economic role, the increasing demand for REE and the significant risks of supply, European governments promote exploration for new REE resources and the study of non-conventional geological environments including sedimentary rocks, soils and other weathering profiles (e.g. laterites and bauxites).

In the recent past, vertical distribution of trace REE was widely applied to control essential geochemical and weathering processes in soils, such as adsorption, solubility and transport factors (Tyler, 2004; Santana et al., 2015; Bern et al., 2017). In addition, the distribution of REE can also be used to determine the occurrence of Fe and Mn oxides in soils, due to the higher adsorptive affinity of LREE in the soil mineral lattice, compared to HREE (Chang et al., 2016).

However, several recent works carried out in different regions worldwide reported that REE contents in weathered rocks could also occur, not only at trace levels but also at concentrations of economic value. The first examples of economically significant REE contents in weathered rocks were reported in bauxites from China (Wang et al., 2000, 2003, 2010; Chai et al., 2001; Li et al., 2005) and, more recently, in Turkey, Greece (Deady et al., 2016) and in the Dominican Republic (Torró et al., 2017). Relevant REE concentrations have also been detected in laterites in regions such as Greece (Eliopoulos, 2014) and the Dominican Republic (Aiglsperger et al., 2016).

Paleosol REE-geochemistry has widely attracted scientific community interest. Studies have been carried out in worldwide locations, such as Southern China (Bao and Zhao, 2008; Wang et al., 2010; Chan et al., 2016), Brazil (Angelica and da Costa, 1991; Marques et al., 2004; Santana et al., 2015), Cameroon (Beyala et al., 2009), South Carolina (Foley and Ayuso, 2015), the Caribbean region (Aiglsperger et al., 2016; Torró et al., 2017), Cape Verde (Marques et al., 2016) and Spain (Fernández-Caliani 2018; Reinhardt et al., 2019), among others. In all of these manuscripts, the study of the geochemical evolution

of each area reflects the occurrence of weathering processes, the corresponding formation of soils and their relation with the concentration of metals (including Al and Fe but also minor elements, such as REE). It has also been noted how supergene products generate a significant variety of REE bearing minerals, such as aluminium-phosphate–sulphate (APS) phases (Dill et al., 1995) and phosphates, commonly monazite (Oelkers and Poitrasson, 2002; Marques et al., 2004; Santana et al., 2015).

Our investigation develops a geochemical exploration survey of the felsic, volcanic and associated sedimentary rocks from the three magmatic stages of the island of Gran Canaria. The first cycle corresponds to the Miocene-aged, alkaline declining stage, whereas the other two correspond to the Plio-Quaternary, volcanic rejuvenation stage. The study was focused on the REE distribution in the soils that developed on the alkaline felsic volcanic rocks from the island of Gran Canaria. The aim of the present study is to contribute towards the understanding of REE behaviour and accumulation in the soil profile and to evaluate the potential of soils developed from felsic volcanic rocks within intraplate oceanic islands as non-conventional sources of REE.

#### 2. Geological setting

The island of Gran Canaria is approximately located at the centre of the Canary Islands archipelago (Fig. 1), which is a Cenozoic, hotspot-derived, intraplate system developed on oceanic Jurassic lithosphere (Schmincke, 1976, 1982; Hollik et al., 1991; Steiner et al., 1998). This island definition agrees with the Walker (1990) model of the intraplate oceanic islands formation, with submarine and subaerial stages. The island growth is similar to that described for other intraplate, oceanic, volcanic islands worldwide (Walker, 1990; Caroff et al., 1995; Le Dez et al., 1996; Carracedo et al., 2002; Hekinian et al., 2004; Schmincke and Sumita, 2010; Staudigel and Clage, 2010). The subaerial growth of Gran Canaria (Fig. 1) is characterised by a sequence of three main magmatic phases: the (i) shield-building, (ii) alkaline declining and (iii) volcanic rejuvenation stages, the last two were separated by an erosional gap of approximately 2 million years, between 7.3 and 5.3 Ma (Fuster et al., 1968; Lietz and Schmincke, 1975; McDougall and Schmincke, 1976; Feraud et al., 1981; Balcells et al., 2004; van den Bogaard, 2013). The shield-building stage records the growth of a complex mafic shield between 14.5 and

14.1 Ma and the subsequent formation of a collapse caldera (Tejeda Caldera) at ca. 14.1 Ma (Schmincke and Swanson, 1966; Schmincke, 1982; Freundt and Schmincke, 1995) and highly explosive post-caldera volcanism (ca. 14.1 to 7.3 Ma). The volume of subaerial, mafic volcanic material erupted in the shield stage was about 1000 km<sup>3</sup> (Schmincke, 1976, 1982; Schmincke and Sumita, 2010). The Miocene mafic shield was overlaid with peralkaline, phonolitic-trachytic to rhyolitic lavas and ignimbrites (Fuster et al., 1968; Schmincke, 1976, 1990), intruded by a felsic cone-sheet swarm, dykes, domes and plugs (Schirnick et al., 1999). The estimated volume of subaerial, mafic and felsic volcanic material erupted at each stage totals about 1000 km<sup>3</sup> (Schmincke, 1976, 1982; Schmincke and Sumita, 2010). During the gap in volcanism, the island underwent strong erosion, which led to the formation of a radially arranged set of ravines and a large volume of sediment was deposited as alluvial and fan-delta conglomerates (fanglomerates), mainly on the S-SW and N-NE coastal platforms (Fuster et al., 1968; Lietz and Schmincke, 1975; Balcells et al., 1992). Rejuvenated volcanism comprised two main phases: (i) Roque Nublo, and (ii) Post-Roque Nublo (Fig. 1). The Roque Nublo phase (5.3 to 2.7 Ma (Guillou et al., 2004)) refers to the evolution of a complex stratovolcano in the central area of the island. The Roque Nublo stratovolcano emitted products with a basanitic and ankaramitic to trachytic and phonolitic composition, such as lava flows, ignimbrites and domes (Balcells et al., 1992; Pérez-Torrado et al., 1995). The volume of subaerial, mafic and felsic volcanic material erupted in this rejuvenation stage was about 200 km<sup>3</sup> (Schmincke, 1976, 1982; Schmincke and Sumita, 2010). The Post-Roque Nublo stage (3.7 Ma to present) is characterised by effusive and strombolian activity from vents aligned mainly along a NW-SE rift (Guillou et al., 2004). Firstly, mafic fissural volcanism (mostly lava flows and fall deposits) occurred with a progressive migration of emission centres to the north. Secondly, the so-called "platforming lavas" were randomly formed on the northern and eastern parts of the island, around 1.5 Ma. Finally, the most recent volcanism (<1 Ma) involved the scattered strombolian and phreatomagmatic eruptions of alkaline, intermediate and mafic magmas over the centralnorthern part of the island. The volume of subaerial, mafic and intermediate volcanic material erupted during this rejuvenation stage was estimated to be 10 km<sup>3</sup> (Schmincke, 1976, 1982; Schmincke and Sumita, 2010).

#### 3. Materials and Methods

#### 3.1. Sampling

This study was undertaken on a total of 57 samples of igneous bedrock, paleosol and sedimentary materials from Gran Canaria (Table 1a-b and Fig. 1). In addition, 12 samples were selected from three representative paleosols, all of them from the Miocene alkaline declining stage: (i) the Fataga profile, developed on a trachy-phonolitic lava bedrock (R-PH; Figs. 2a, 3), (ii) El Queso profile, which comprised a trachy-rhyolitic ignimbrite bedrock (R-RH; Figs. 2b, 3), and (iii) the Culata profile, which is composed of trachytic lava bedrock (R-RH; Figs. 2c, 3). The Fataga profile (PH) consists of 20 to 30 cm of saprolite, a 20 to 30 cm weak red (10R5/3) B-horizon, a 30 to 50 cm very pale brown (10YR7/3) and a laminated B-horizon. The El Queso profile (RH) is formed by metric layers of weak, red rhyolitic ignimbrites (10R5/4) with saprolite, topped by a brown (7.5YR4/4) B-horizon. The Culata profile (TR) is a well-developed and preserved profile consisting of two continuous saprolite levels, from 50 to 70 cm, at about 1 m depth, each being a reddish brown (5YR5/4) B-horizon and a, presumably truncated, 20 cm dark reddish grey (5YR4/2) BA-horizon.

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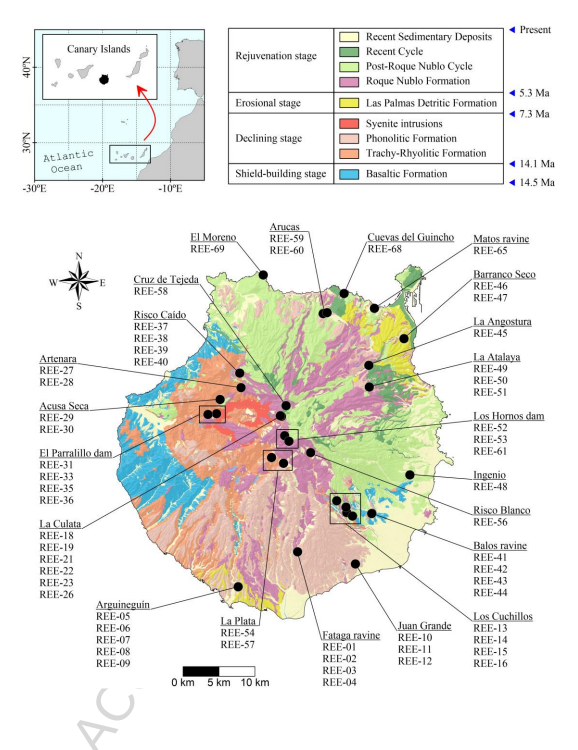


Figure 1. Geographical setting (upper left), description of geological units (upper right) and geological map (main view) of Gran Canaria (modified after Bellido-Mulas and Pineda-Velasco, 2008). Sampling locations are also plotted (black dots). Sample names are shown in toponymical groups for clarity purposes.

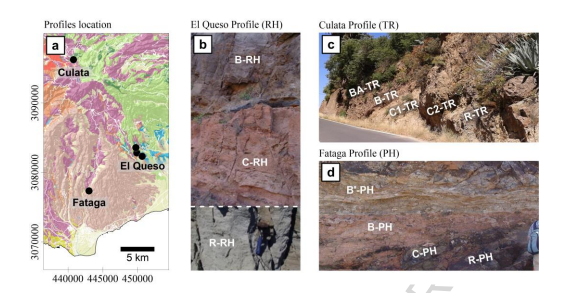


Figure 2. (a) Geological map with the location of the studied samples (in black dots) and the respective name of the profiles, which are located in the south-eastern area of the island (see Fig. 1); (b, c, and d) Field photographs of El Queso, Culata and Fataga profiles with their related horizons (white labels). For more details, see Table 2a and 2b.

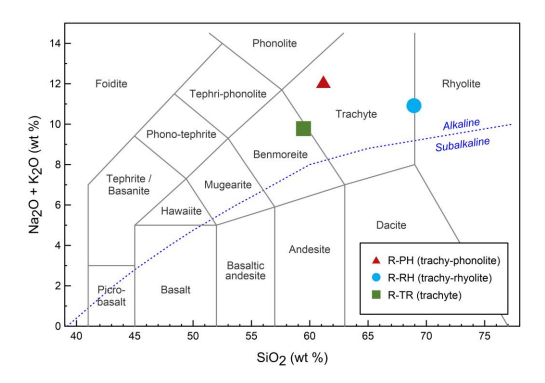


Figure 3. TAS diagram for the volcanic bedrocks of the area studied (diagram after Le Bas et al., 1986; dashed curve from Irvine and Baragar, 1971).

Table 1a. Sample set listed with geographical coordinates (WGS84; UTM zone 28N), elevation (m asl) and location references for the igneous and sedimentary rocks collected within the Gran Canaria study area.

Sample	Easting	Northing	Elevation	Location
REE-01	443025	3077367	472	GC-60 road, Fataga
REE-02	443025	3077367	472	GC-60 road, Fataga
REE-03	443025	3077367	472	GC-60 road, Fataga
REE-04	443025	3077367	472	GC-60 road, Fataga
REE-05	434771	3072482	91	Ceisa quarry, Arguineguín
REE-06	434771	3072482	91	Ceisa quarry, Arguineguín
REE-07	434771	3072482	91	Ceisa quarry, Arguineguín
REE-08	434771	3072482	91	Ceisa quarry, Arguineguín
REE-09	434771	3072482	91	Ceisa quarry, Arguineguín
REE-10	451085	3075661	85	Lopesan quarry, Juan Grande
REE-11	451085	3075661	85	Lopesan quarry, Juan Grande
REE-12	451085	3075661	85	Lopesan quarry, Juan Grande
REE-13	449878	3082826	550	GC-65 road, El Queso
REE-14	449761	3083587	552	GC-65 road, El Queso
REE-15	448475	3084471	568	GC-550 road, Santa Lucía de Tirajana
REE-16	450665	3082333	459	GC-65 road, El Queso
REE-18	440783	3096230	1160	GC-608 road, La Culata
REE-10	440783	3096230	1160	GC-608 road, La Culata
REE-21	440783	3096230	1160	GC-608 road, La Culata
REE-21 REE-22	440783 440783	3096230	1160	GC-608 road, La Culata
REE-22 REE-23	440783	3096230	1160	GC-608 road, La Culata
	440783 440641			
REE-26		3096312	1176	GC-608 road, La Culata
REE-27	435157	3100231	1142	GC-210 road, Artenara
REE-28	435157	3100231	1142	GC-210 road, Artenara
REE-29	432247	3098579	810	GC-210 road, Acusa Seca
REE-30	432247	3098579	810	GC-210 road, Acusa Seca
REE-31	431765	3096604	514	GC-210 road, El Parralillo dam
REE-33	431765	3096604	514	GC-210 road, El Parralillo dam
REE-35	430559	3096485	315	GC-210 road, El Parralillo dam
REE-36	430559	3096485	315	GC-210 road, El Parralillo dam
REE-37	434982	3102241	931	Risco Caído, Artenara
REE-38	434982	3102241	931	Risco Caído, Artenara
REE-39	434982	3102241	931	Risco Caído, Artenara
REE-40	434982	3102241	931	Risco Caído, Artenara
REE-41	453383	3082737	199	Los Corralillos, Balos ravine
REE-42	453383	3082737	199	Los Corralillos, Balos ravine
REE-43	453341	3082680	221	Los Corralillos, Balos ravine
REE-44	453341	3082680	221	Los Corralillos, Balos ravine
REE-45	452936	3103343	355	GC-320 road, La Angostura
REE-46	457838	3107038	93	GC-5 road, Barranco Seco
REE-47	457838	3107038	93	GC-5 road, Barranco Seco
REE-48	458648	3088079	176	GC-189 road, Ingenio
REE-49	452975	3100322	555	La Atalaya parking, Santa Brígida
REE-50	452975	3100322	555	La Atalaya parking, Santa Brígida
REE-51	453018	3100318	548	La Atalaya quarry, Santa Brígida
REE-52	441227	3093525	1609	GC-600 road, Los Hornos dam
REE-53	441227	3093525	1609	GC-600 road, Los Hornos dam
REE-54	439404	3090501	1226	La Lajilla guarry, GC-60 road, La Plata
REE-56	444845	3091174	1031	GC-654 road, Risco Blanco
REE-57	441068	3089684	1237	GC-60 road, La Plata
REE-58	441433	3097709	1538	GC-150 road, Cruz de Tejeda
REE-59	441433	3110526	334	Lomo Tomás de León, Arucas
REE-59 REE-60	440300 447135	3110526	258	Lomo Tomás de León, Arucas
				GC-600 road, Los Hornos dam
REE-61	441830	3092760	1645	
REE-65	453717	3111267	34	Matos ravine, El Rincón
REE-68 REE-69	449505	3113343	85	GC-2 road, Cuevas del Guincho El Moreno headland, Gáldar
	438251	3115919	2	

Table 1b. Sample set sorted by island stage and geological formation, description of rock lithologies (V: volcanic; SV: subvolcanic; P: plutonic; S: sedimentary; VS: volcano-sedimentary) and bulk rock values of LREE (La-Eu), HREE (Gd-Lu), LREE/HREE ratios, total REE and yttrium (in mg kg<sup>-1</sup>).

Stage	Formation	Sample	Туре	Description	LREE	HREE	LREE/HREE	Σ REE	Y
Rejuvenation	Post-Roque Nublo	REE-48	VS	Red paleosol	397	32	13	429	43
(<5.3 Ma)	(3.5 Ma - Present)	REE-58	VS	Red paleosol	584	32	18	616	35
		REE-68	V	Tephrite-phonolite lava flow	357	19	18	376	26
	Roque Nublo	REE-15	V	Trachytic lava flow	422	24	17	446	30
	(5.3 - 2.9 Ma)	REE-29	S	Sandstone	502	29	17	531	39
		REE-30	S	Siltstone	427	25	17	452	35
		REE-37	VS	Lahars	453	26	17	479	35
		REE-38	VS	Lahars	524	27	20	550	36
		REE-39	VS	Lahars	497	28	18	526	37
		REE-40	VS	Lahars	441	21	21	461	28
		REE-45	S	Paleosol	385	19	20	404	25
		REE-52	SV	Phonolitic collapsed dome	502	26	19	528	36
		REE-53	SV	Phonolitic edge dome	471	26	18	496	31
		REE-56	SV	Phonolitic dome	776	50	15	826	68
		REE-61	V	Block and ash ignimbrite	419	25	17	444	31
		REE-69	V	Block and ash ignimbrite	514	33	16	547	44
Erosional	Las Palmas Detritic	REE-46	S	Siltstone	382	23	17	405	26
(7.3 - 5.3 Ma)	Formation (LPDF)	REE-47	S	Siltstone	369	24	15	393	30
Declining	Trachy-phonolitic	REE-1	V	Phonolite lava flow	970	67	15	1036	102
(14.1 - 7.3 Ma	) (13.4 - 7.3 Ma)	REE-2	S	Paleosol	1082	89	12	1171	158
		REE-3	S	Paleosol	931	64	15	994	82
		REE-4	S	Paleosol	1053	86	12	1139	123
		REE-5	V	Ash and pumice ignimbrite	615	39	16	654	54
		REE-6	S	Paleosol	507	35	14	542	49
		REE-7	S	Paleosol	669	29	23	698	36
		REE-8	V	Ash and pumice ignimbrite	785	51	15	836	73
		REE-9	S	Paleosol	640	41	16	681	58
		REE-10	V S	Phonolite lava flow	776	52	15	828	69 62
		REE-11 REE-12	S	Paleosol Paleosol	354 780	34 53	10 15	388 833	63 66
		REE-12 REE-18	V	Ash and pumice ignimbrite	427	53 22	15	633 449	29
		REE-19	s	Paleosol	448	22	20	471	29
		REE-21	s	Paleosol	474	24	20	498	30
		REE-22	s	Paleosol	410	26	16	436	31
		REE-23	S	Paleosol	418	30	14	448	39
		REE-26	S	Paleosol	403	18	22	421	27
		REE-27	SV	Phonolitic dome	388	11	36	399	21
		REE-28	SV	Phonolitic dome top	322	11	31	333	20
		REE-31	SV	Phonolitic felsic dyke	719	38	19	757	60
		REE-33	SV	Trachytic felsic dyke	932	58	16	990	78
		REE-35	Р	Intracaldera syenite	447	27	16	474	35
		REE-36	Р	Intracaldera syenite	550	40	14	590	59
		REE-49	S	Paleosol	808	49	17	857	59
		REE-50	V	Ash and pumice ignimbrite	424	24	18	448	28
		REE-51	V	Ash and pumice ignimbrite	264	22	12	286	45
	V	REE-54	SV	Phonolitic dome	560	17	32	577 576	31
		REE-57	SV V	Phonolitic dome	542 388	34 30	16 13	576 418	46 43
		REE-59 REE-60	V V	Block and ash ignimbrite Block and ash ignimbrite	388 847	30 55	13	418 902	43 68
		REE-60 REE-65	V	Block and ash ignimbrite	647 534	32	15	902 566	42
	Trachy-rhyolitic	REE-13	v	Rhyolitic lava flow	535	53	10	588	42 74
	(14.1 - 13.4 Ma)	REE-14	v	Paleosol	549	52	10	601	73
		REE-16	v	Paleosol	1484	100	15	1584	121
		REE-41	s	Siltstone base	1225	55	22	1280	74
		REE-42	S	Sandstone base	363	31	12	394	40
		REE-43	S	Siltstone roof	718	36	20	753	43
		REE-44	S	Sandstone roof	272	26	10	298	35

#### **3.2. Optical Microscopy Studies**

Selected samples from each horizon profile were prepared as double-polished thin sections for their study under a petrographic microscope (Leitz Orthoplane), observed with both transmitted and reflected light. The microphotographs were taken with a Leica DFC 420 coupled camera. In addition, these thin sections were used for the microprobe analysis.

#### **3.3. Powder X-ray Diffraction (PXRD)**

Samples of the selected profiles were crushed in an agate mortar, obtaining a particle size below 40  $\mu$ m. The experimental profiles were performed using a PANalytical X'Pert PRO Alpha1 powder diffractometer at the Scientific and Technological Centre of the University of Barcelona (CCiTUB). The diffractometer equipment was used with monochromatised incident Cu K $\alpha$ 1 radiation at 45 kV and 40 mA and was equipped with a PS detector with an amplitude of 2.113°. The patterns were obtained by scanning random powders from 4° to 80° (20). Data sets were obtained by a scan time of 50 seconds at a step size of 0.017° (20) and a variable divergence slit. Mineral identification and semi-quantitative results were facilitated using the X'Pert search-match software with Powder Diffraction File, version 2 from JCPDS. When the occurrence of quartz was detected in the sample, it was used as an internal standard to correct diffraction patterns for instrumental shifts in the 20 position. Quantitative mineral phase analyses were obtained by full refinement profile using TOPAS V4.2 software.

#### 3.4. Scanning Electron Microscopy (SEM and FE-SEM)

A selection of these samples was examined on a scanning electron microscope (SEM) equipped with an energy-dispersive spectrometer (EDS), at the CCiTUB. The operating conditions were 15 to 20 kV accelerating voltage and 5nA beam current. In addition, we carried out a more detailed study of REE mineral phases to determine their textural and qualitative compositional features with a JEOL JSM-7100 field emission scanning electron microscope (FE-SEM) at the CCiTUB.

#### 3.5. Electron probe microanalysis (EPMA)

Further identification of REE mineral phases was started using EPMA spot analyses. EPMA was performed on a JEOL JXA-8230 electron microprobe with 5 wavelengthdispersive spectrometers (WDS) and a silicon-drift detector EDS, at the CCiTUB.

Three X-ray intensity maps of Ce, La, Fe, Al and P (plus backscattered electron images (BSE)) were collected using the EPMA operated in WDS mode by stage scanning, with an accelerating voltage of 20 kV and a beam current of 50 nA. The mapped areas were 92 x 92  $\mu$ m, 280 x 420  $\mu$ m and 112 x 140  $\mu$ m, with dwell times of 10 ms, 50 ms and 40 ms per point, respectively. Pixel sizes (point spacing) were 0.18  $\mu$ m, 0.70  $\mu$ m and 0.28  $\mu$ m, respectively. Spectrometers were peaked to the specific X-ray line positions using appropriate standards. Background maps, acquired with the WDS tuned to a background position on the right of the corresponding peak, were subtracted from the corresponding peak intensity maps. Total map acquisition times were 2.0 h, 4.5 h and 7.0 h, respectively. Additional X-ray intensity maps for C, Al, Si, Cl, K, Ca, Ti, Mn and Ni were simultaneously collected in EDS mode.

Spot analyses were carried out with an accelerating voltage of 15 kV, a beam current of 10 nA and a focused beam, in order to achieve the best lateral resolution. Background positions were carefully adjusted. Analytical standards included natural and synthetic silicate, oxides and REE glasses as follows: apatite (Ca and P), UO<sub>2</sub> (U), ThO<sub>2</sub> (Th), LaB<sub>6</sub> (La), Ce<sub>2</sub>O<sub>3</sub> (Ce), REE glasses (Pr, Nd), NiO (Ni), hematite (Fe), yttrium-aluminium garnet (Y), ZrO<sub>2</sub> (Zr), Gd<sub>3</sub>Ga<sub>5</sub>O<sub>12</sub> (Gd), and rhodonite (Mn). The correction procedure 'XPP' was used to convert specimen intensity ratios into concentrations. Counting times ranged from 20 to 40 s for both peak and background. Detection limits were ~ 300-400 mg kg<sup>-1</sup> for Ca, P, Ni and Fe, ~500 mg kg<sup>-1</sup> for Y, La and Ce, and ~1100-1300 mg kg<sup>-1</sup> FOR U, TH, PR, GD AND ND.

#### 3.6. Bulk Rock Geochemistry

Major, minor and trace element analyses of 57 samples of rock, sediment and paleosol were determined at the ACTLABS Activation Laboratories Ltd., Ontario (Canada). The major elements SiO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, Fe<sub>2</sub>O<sub>3</sub>, TiO<sub>2</sub>, K<sub>2</sub>O, MgO, Na<sub>2</sub>O, CaO, P<sub>2</sub>O<sub>5</sub>, MnO, Ba, Be, Sc, Sr, V, Y, and Zr were analysed by FUS-ICP, in addition to loss on ignition (LOI). Trace elements, including Ag, Bi, Cr, Cs, Co, Cu, Ga, Ge, Hf, In, Mo, Ni, Nb, Pb, Rb,

REE, Sb, Sn, Ta, Th, Tl, U, W, Zn were analysed by FUS-MS; As, Au, Br, Hg, Ir, Sc, Se were analysed by INAA; Cd and S were analysed by TD-ICP. Major element detection limits ranged from 0.01 to 0.10 wt% (1 wt% equals 10,000 mg kg<sup>-1</sup>) and trace element detection limits ranged from 0.01 to 1.00 mg kg<sup>-1</sup>, except for gold (Au) in ppb, which had a detection limit of 1  $\mu$ g kg<sup>-1</sup>. Au, Hg and Ir concentrations are systematically below their respective detection limit for the used analytical methods.

Gains or losses of each element were measured as the percentage change of the elemental ratio between such elements in the weathering product and in the parent rock (Nesbitt, 1979) as follows:

% change of ratios = 
$$100 [(Ri - Rp)/Rp]$$
 [1]

Ri and Rp represent the element ratio in weathered samples and fresh bedrock respectively, assuming that volume does not change during weathering (Nahon and Merino, 1996). To compare the weathering degree of the profiles, we selected the Vogt's Residual Index (V), which is sensitive to subtle geochemical changes such as alteration at the water table and/or hydrothermal alteration along faults, using the molecular proportions as follows (Price and Velbel, 2003):

$$V = (Al_2O_3 + K_2O) / (MgO + CaO + Na_2O)$$
[2]

Moreover, the Chemical Index of Alteration (CIA) was calculated from the chemical analysis of the weathered materials and parent rocks, using the molecular proportions as follows (Nesbitt and Young, 1989; McLennan et al., 1993; Price and Velbel, 2003):

$$CIA = [Al_2O_3 / (Al_2O_3 + K_2O + Na_2O + CaO)] \times 100$$
[3]

Major element oxides are divided by their molar mass. The amount of CaO represents calcium in the silicate fraction. The TR and RH are carbonate free whereas, in the PH profile, the carbonate content is 1 to 3%. In the latter cases, CaO<sub>silicate</sub> was calculated as (McLennan et al., 1993, Zhao et al., 2017):

$$CaO_{silicate} = CaO_{total} - (CaCO_3 \times 56.1/100.1)$$
[4]

#### 3.7. Iron oxides: sequential extraction and diffuse reflectance analysis

Iron oxides of the RH profile, which presents the highest REE concentrations, were measured at the Department of Agronomy of the University of Córdoba. Three types of Fe oxides were sequentially extracted. Firstly, ferrihydrite (Fh), corresponding to the Fe

dissolved with acid ammonium oxalate by the Schwertmann method (1964) (Fe<sub>o</sub>); secondly, maghemite (Mgh), corresponding to the Fe extracted by sulphuric acid (Fe<sub>s</sub>); and thirdly, the Fe in hematite (Hm) plus goethite (Gt), which were extracted using sodium dithionite in a hot (75°C) sodium citrate and sodium bicarbonate solution (Mehra and Jackson, 1960) (Fe<sub>d</sub>). If we assign Fe<sub>d</sub> to the combination of Fe in stoichiometric Hm ( $\alpha$ -Fe<sub>2</sub>O<sub>3</sub>) and Gt ( $\alpha$ -FeOOH), the Gt content can be estimated as follows:

$$Gt = 1.59 \text{ x} (Fe_d - Hm/1.43)$$
 [5]

The Hm/Gt ratio was estimated by means of diffuse reflectance spectroscopy (DRS; Scheinost et al., 1998; Torrent and Barrón, 2002; Torrent et al., 2007). The DRS is a precise and non-destructive method used to quantify soil properties and can detect the presence of either hematite or goethite at < 0.1% in mixtures with other soil minerals. The DRS spectra of the studied samples were recorded at a scan rate of 30 nm min<sup>-1</sup> and from 380 to 770 nm in 0.5 nm steps, using a Varian Cary 5000 UV-Vis-NIR spectrophotometer equipped with a diffuse reflectance attachment. The relation between Hm and Gt was estimated from the amplitude of the Hm and Gt peaks of the second derivative spectrum of the Kubelka-Munk function (A<sub>Hm</sub> and A<sub>Gt</sub>), according to the equation given by Scheinost et al. (1998):

$$Hm/(Hm + Gt) = 0.068 + 1.325(A_{Hm}/(A_{Hm} + A_{Gt}))$$
 [6]

Atomic Absorption Spectroscopy (AAS), using a Perkin-Elmer 420 instrument, was used to determine iron concentration in all extracts. The total Fe content (Fe<sub>t</sub>), expressed as ion weight percentage, was calculated from the Fe<sub>2</sub>O<sub>3</sub> weight percentage. REE concentration of the B-RH sample from the three extractions was measured by ICP after a digestion procedure, as described by Chang et al. (2016). The sample was mixed with 6 ml of concentrated HNO<sub>3</sub>–HClO<sub>4</sub> (87:13, v/v) and 6 ml of concentrated HF (mass fraction 40%). The mixture was digested and then dissolved in 15 ml of 2% HCl solution.

#### 4. Results

#### 4.1. Mineralogy and textures

#### 4.1.1. Bedrocks

The petrographic study of the trachy-phonolitic lava (Fataga profile, Fig. 2a) shows a fluidal trachytic texture, with microcryptocrystaline mesostasis surrounding the phenocrysts. The phenocrysts and microphenocrysts are mainly anorthoclase and, in minor amounts, nepheline, aegirine, kaersutite, biotite with oxidised rims, and haüyne. The mesostasis is composed of micro/crypto crystals of anorthoclase and, in minor proportions, nepheline, aegirine, kaersutite (occasionally with agpaitic texture), iron oxides, biotite, haüyne and clay minerals (Fig. 4a).

The trachy-rhyolitic block and ash ignimbrite (El Queso profile, Fig. 2b) present a tuffy texture of decimetric trachytic, syenitic and olivine-pyroxene basaltic rock fragments, devitrified pumice fragments, oligoclase and anorthoclase crystals and occasionally microcrystals of augite-aegirine, iron oxides, slightly weathered kaersutite, biotite, and devitrified cineritic mesostasis (Fig. 4b).

The trachytic ash and pumice ignimbrite (La Culata profile, Fig. 2c) presents a finely fragmented texture of ash, pumice, lithic and mineral fragments, typical of ignimbrite rocks (MacKenzie et al., 1984) with an irregular and discontinuous banding of felsic and ferromagnesian microcrystals, pumice and lithic fragments. This rock is formed of millimetric trachytic rock fragments, rare basaltic rock fragments, pumice fragments, anorthoclase millimetric crystals and, occasionally, microcrystals of iron oxides and biotite, as well as devitrified and recrystallised cineritic mesostasis (Fig. 4c).

#### 4.1.2. Paleosol horizons

The PXRD analysis of the paleosols horizons identified mainly anorthoclase and oligoclase with minor accessory phases (i.e. diopside, hornblende, feldspatoids, titanite, biotite and fluorapatite). Weathering processes promoted depletion of the primary minerals and the formation of secondary phases, such as clay minerals (smectite, illite and kaolinite), calcium carbonates and iron and manganese oxyhydroxides. Also, we identified the occurrence of exotic quartz grains (Fig. 5) because the silica subsaturated nature of potential protoliths in the Canary Islands (Carracedo et al., 2002; Vera, 2004), precludes the occurrence of quartz in the volcanic paragenesis that built the islands.

The concentration of REE is conspicuously low and mainly concentrated in the upper part of the weathering profiles, in-filling the pores and small cavities and covering mineral

surfaces (Fig. 6). These scarce REE minerals are mostly LREE-bearing phosphates that occur as euhedral to subhedral grains up to  $10 \mu m$  in size (Fig. 7a-c).

Because of the small grain size and heterogeneity, accurate determination of the chemical composition of LREE-bearing phases is difficult. Reliable data were only obtained for a limited number of phases, such as monazite-Ce and rhabdophane-Ce. Representative EPMA analyses, along with the corresponding structural formulae of monazite-Ce and rhabdophane-Ce phases, are presented in Table 2. Fe-Mn and REE oxides were identified by SEM-EDS (semi-quantitative).

Monazite-(Ce) [(Ce,La,Nd)PO<sub>4</sub>] is the most common REE phosphate and occurs as euhedral to subhedral isolated grains, exhibiting irregular borders. It is generally altered and presents light rims and lobate reaction fronts as described by Williams et al. (2011) for primarily altered domains (Fig. 7a). It also presents dissolution-precipitation textures where the monazite is transformed to rhabdophane-Ce (Fig. 7d-h). REE silicates (probably of the britholite group) have also been identified, associated with monazite-(Ce) and forming anhedral, subrounded grains up to 5  $\mu$ m in size (Fig. 7d). Around these primary anhydrous phosphates, alteration halos often develop, made up of aggregated acicular crystals of rhabdophane-(Ce) [(Ce,La,Nd,Sm)(PO<sub>4</sub>)·H<sub>2</sub>O] (Fig. 7d-f) and mostly accompanied by concentric layers of REE oxides (Fig. 7g-i).

#### 4.2. Mineral chemistry of paleosols

In order to further examine the distribution of REE and their relation with other significant elements in the studied paleosol profiles, two background-corrected X-ray intensity maps, obtained on thin section from two different areas with previously detected occurrence of REE minerals, are presented in Figs. 8 and 9. A detailed inspection of the X-ray maps showed the relationships between textural position, crystallisation sequence and mineral composition of the samples containing REE. Note that we acquired X-ray maps for several LREE (e.g. La, Ce, Py, Nd, Pm and Sm) but only La and Ce were detected and, consequently, are presented in Figs. 8-9. We also scanned for significant major elements related to the paleosol's mineralogy (i.e. C, Mg, Al, Si, P, Cl, K, Ca, Ti, Mn, Fe, and/or Ni).

The first map (Fig. 8) shows complex and variable textures. In general, the concentration of REE is mostly controlled by the occurrence of primary, isolated grains of LREE-rich phosphates, which were also detected during the previous mineralogical and textural studies using SEM and FE-SEM. The LREE-rich phosphates are mainly grains of monazite-(Ce) that may also contain significant amounts of La (Fig. 8, Table 2). We also found a second Ce-rich phosphate that forms thin concentric layers around the monazite-(Ce) grains (Fig. 9). This second phosphate does not contain La and it is associated with REE oxides and Mn oxyhydroxides (Fig. 8). Figure 8 shows that the distribution of REE is mainly controlled by phosphate phases. REE contents are below detection limits or negligible in the silicates as well as in the carbonate phases present in the samples (Fig. 8).

In the second mapped area (Fig. 9), the distribution of REE shows similar features. Small isolated grains of Ce-rich phosphates, presumably identified as monazite-(Ce), are spread around the analysed area and control the distribution of LREE. However, in this case, the La contents are negligible (Fig. 9). Neither the carbonates nor the main silicates contain REE. The occurrence of Mn oxyhydroxides with very significant Ni contents are also highlighted (Fig. 9).

Table 2. EPMA analyses: composition of REE phosphates (monazite-Ce and rhabdophane-Ce) from Gran Canaria weathering profiles. (b.d.l. means below the detection limit).

		Monazite	9	Rh	Rhabdophane					
analysis №.	C-16	C-34	C-35	C-15	C-32	C-33				
, wt.%										
FeO <sub>t</sub>	0.30	1.73	3.49	1.36	0.30	1.45				
CaO	0.71	0.57	0.17	1.03	0.86	0.96				
$P_2O_5$	29.31	29.11	28.27	27.49	28.26	27.28				
$Y_2O_3$	0.10	0.07	0.08	0.23	b.d.l.	0.24				
ThO <sub>2</sub>	b.d.l.	0.20	0.07	b.d.l.	0.27	0.15				
UO <sub>2</sub>	0.03	0.09	b.d.l.	b.d.l.	b.d.l.	0.15				
$La_2O_3$	17.31	17.34	16.54	16.03	16.1	15.81				
$Ce_2O_3$	35.34	34.48	35.34	32.54	33.79	33.02				
$Pr_2O_3$	3.05	3.32	3.06	3.02	2.92	3.18				
$Nd_2O_3$	11.13	10.67	11.83	10.19	10.6	10.81				
H <sub>2</sub> O				6.98	7.17	6.92				
Total	97.27	97.58	98.85	91.89	93.09	93.05				
apfu										
Fet	0.010	0.058	0.117	0.048	0.010	0.051				
Ca	0.030	0.024	0.007	0.047	0.038	0.044				
Р	0.995	0.987	0.961	0.986	0.999	0.975				
Y	0.002	0.001	0.002	0.005	b.d.l	0.005				
Th	b.d.l	0.002	0.001	b.d.l	0.003	0.001				
U	b.d.l	0.001	b.d.l	b.d.l	b.d.l	0.001				
La	0.256	0.256	0.245	0.250	0.248	0.246				
Ce	0.519	0.505	0.520	0.504	0.516	0.511				
Pr	0.045	0.048	0.045	0.047	0.044	0.049				
Nd	0.159	0.153	0.170	0.154	0.158	0.163				
∑REE	0.981	0.964	0.981	0.961	0.967	0.974				

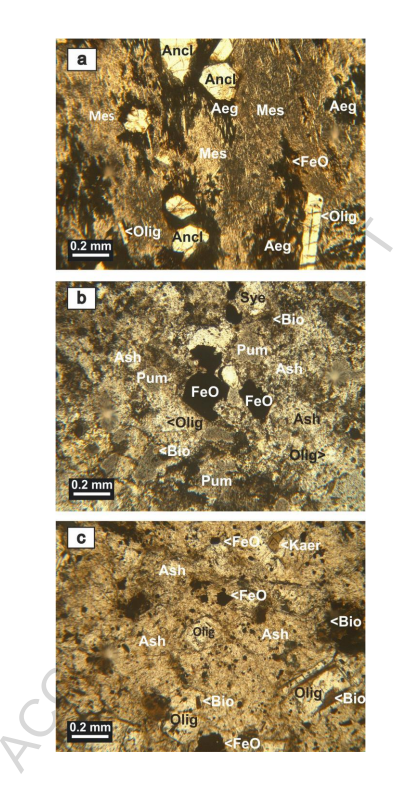


Figure 4. Plane-polarised light microphotos (PPL, 40x) of Fataga (A, phonolitic lava flow), El Queso (B, block and ash ignimbrite) and Culata (C, ash and pumice ignimbrite) bedrocks. (Abbreviations - Ancl: anorthoclase; Aeg: aegirine; Olcl: oligoclase; Mes: mesostasis; FeO: iron oxides; Sye: syenite rock fragment; Bio: biotite; Pum: pumice; Kaer: kaersutite).

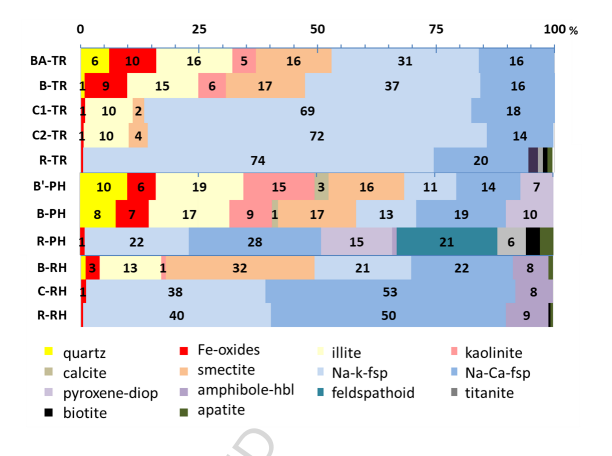


Figure 5. Mineralogical composition of samples estimated by PXRD, except those for bedrocks by petrographic microscope analysis. X-axis refers to semi-quantitative mineral abundances expressed as rounded percentages. Y-axis legend: C, B, B' and BA refer to soil horizons. PH: phonolite profile; RH: rhyolite profile; TR: trachyte profile.

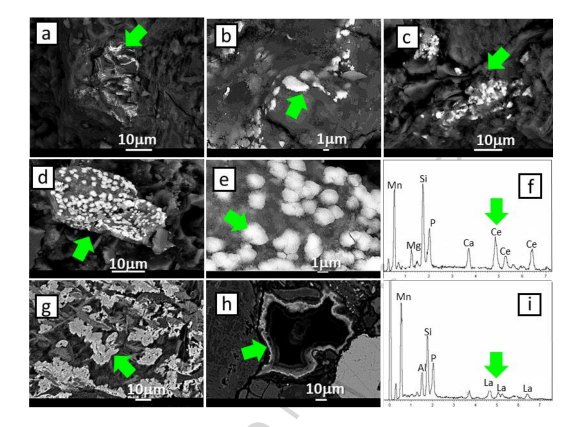


Figure 6. SEM Backscattered-electron (BSE) images of REE bearing phosphates (green arrows) from a B-Horizon sample; (a-e) REE phases from the trachy-phonolite profile (PH); (f) semi-quantitative spectrum of REE phosphates identified from trachy-phonolite; (g-h) backscattering images from the trachy-rhyolite profile (PH); (i) semi-quantitative spectrum from trachy-rhyolite (RH).

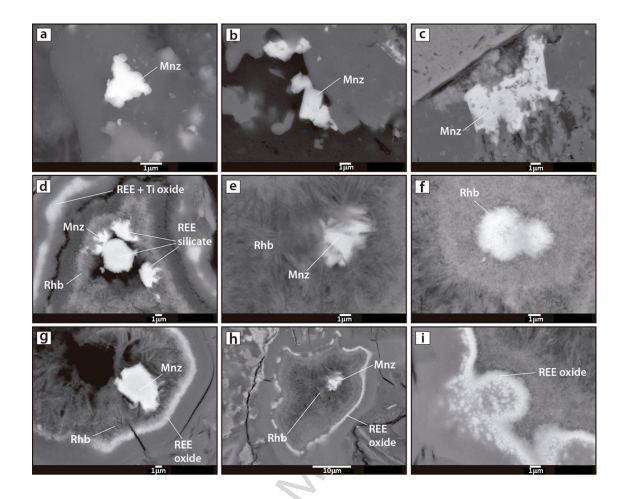


Figure 7. FE-SEM Backscattered-electron (BSE) images of REE bearing minerals found in the upper parts of the studied weathering profiles from Gran Canaria; (a, b) individual monazite-Ce (Mnz) subhedral crystals with slightly rounded outlines; (c) monazite-Ce (Mnz) subhedral grain showing a sharp, irregular outline and abundant scratches; (d) anhedral grains of REE silicate accompanied by subhedral monazite-Ce (Mnz) crystals, surrounded by secondary fibrous rhabdophane-Ce (Rhb) and concentric REE oxides; (e) anhedral monazite-Ce (Mnz) grain floating in an alteration rim of fibrous rhabdophane-Ce (Rhb); (f) typical popcorn texture generated by foamy rhabdophane-Ce (Rhb) growing over a precursor monazite grain; (g, h) anhedral monazite-Ce (Mnz) grain embedded by fibrous rhabdophane-Ce (Rhb) surrounded by a concentric rim of REE oxides; (i) detail of a REE oxide concentric rim.

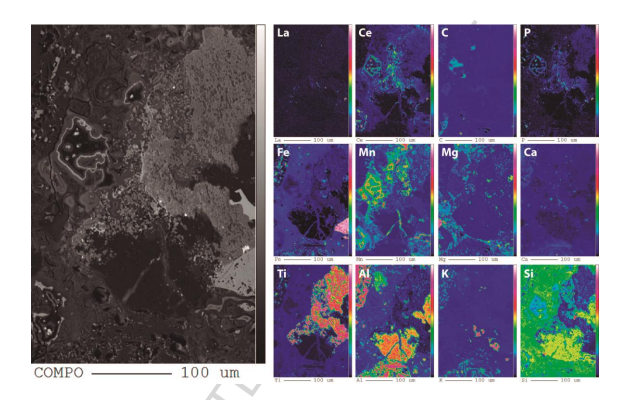


Figure 8. Wavelength-dispersive X-ray maps of representative elements for B-RH sample.

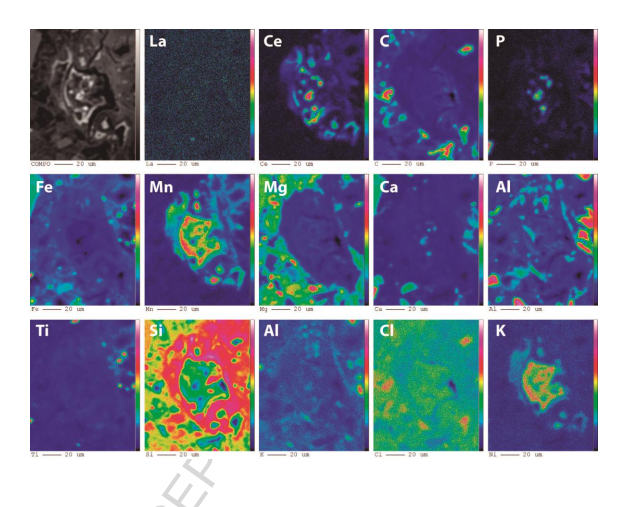


Figure 9. Wavelength-dispersive X-ray maps of representative elements for B-RH sample. Target area is a void selected from Fig. 7.

#### 4.3. Bulk Rock Geochemistry

#### 4.3.1. Major, Minor and trace elements

The bedrock geochemistry of the major, minor and trace elements from the three profiles (Tables 1A-B in the appendix) show a depletion in MgO,  $Fe_2O_3$  and CaO, an enrichment in silica (SiO<sub>2</sub>), alkalis (K<sub>2</sub>O and Na<sub>2</sub>O) and Al<sub>2</sub>O<sub>3</sub>, and low concentrations of MnO, TiO<sub>2</sub> and P<sub>2</sub>O<sub>5</sub> minor elements in relation to the first subaerial emissions (ultramafic, mafic and intermediate rocks). The concentration of trace elements is in agreement with this type of felsic rock, with low concentrations in compatible elements (Cr, Ni, Co, Sc, V) and an enrichment in incompatible elements (Rb, Sr, Ba, Nb, Zr and REE).

According to the TAS diagram (Fig. 3), the bedrocks are defined as trachytes, with slight differences: one is within the limit of the trachyte-benmoreite field (R-TR sample), another one is at the edge of the trachyte-phonolite field (R-PH sample) and the other is trachyte-rhyolite (R-RH sample). Consequently, these bedrocks are highly differentiated igneous rocks from the final stages of the fractional crystallisation.

#### 4.3.2. REE

The REE concentrations in the bulk rocks from the first magmatic cycle of Gran Canaria, are similar to the associated felsic rocks of the Miocene alkaline declining stage and range between 532 ±82 and 725 ±196 mg kg<sup>-1</sup> (Table 1b). Thus, the plutonic rocks (syenite stock) have a mean REE concentration of 532 ±82 mg kg<sup>-1</sup>; the subvolcanic rocks (trachyphonolites dykes and domes) have a mean REE concentration of 725 ±196 mg kg<sup>-1</sup>; the volcanic rocks (trachyphonolite and trachy-rhyolite lava flows and ignimbrites) have a mean REE concentration of 707 ±340 mg kg<sup>-1</sup>; the sedimentary rocks (sandstones and siltstone deposits) have a mean REE concentration of 681 ±443 mg kg<sup>-1</sup>; and the paleosols have a mean REE concentration of 684 ±273 mg kg<sup>-1</sup> (Table 1b). The mean REE concentration for alkaline declining stage is 672 ±296 mg kg<sup>-1</sup>. LREE/HREE ratio is 17 ±5.8 and the mean yttrium (Y) concentration is 57 ±30 mg kg<sup>-1</sup>. The Miocene erosional stage was characterised by a lower REE concentration of 399 ±8 mg kg<sup>-1</sup> from the Las Palmas Detritic Formation (siltstone). Chondrite-normalised plots showed a preferential concentration in LREE (Fig. 10).

The REE concentration of the second magmatic cycle (Roque Nublo Formation, with a magmatic total volume of 200 km<sup>3</sup>), which was part of the rejuvenation stage of the island, presents a mean of  $617 \pm 182 \text{ mg kg}^{-1}$  in the phonolitic domes;  $563 \pm 15 \text{ mg kg}^{-1}$  in trachyphonolitic lava flows and ignimbrites;  $500 \pm 41 \text{ mg kg}^{-1}$  in epiclastic sedimentary rocks (sandstones and siltstone deposits); and a paleosol with a concentration of 404 mg kg<sup>-1</sup>. These rocks, which are associated with the Pliocene magmatic chamber, show a REE concentration lower than the Miocene magmatic chamber.

The REE concentration of the third magmatic cycle (Post-Roque Nublo Formation, with an erupted volcanic rock volume of 10 km<sup>3</sup>), is related to the rejuvenation stage of the island, and presents a value of 376 mg kg<sup>-1</sup> for the tephrite-phonolite lava flow and a value of  $523 \pm 132$  mg kg<sup>-1</sup> for the two red paleosols. In this case, the REE concentrations are also lower than those of the Miocene magmatic chamber and the erupted felsic rocks are negligible in relation to the subaerial, insular vulcanism.

#### 4.3.3. Element redistribution during soil formation

Geochemical losses and gains (% change of ratios) of the paleosol horizons with respect to the bedrock, differ between the three different profiles studied (Figs. 11-12, Tables C-D in the Appendix). Mg, Ca, P, Fe, and Ti were highly retained in the RH paleosol profile but just maintained in the rest and even reduced in B-PH. Elements such as Mn and Al increased in the same proportions in all profiles, except for a reduction in B-PH. Silica, Na and K suffered a moderate reduction. Some elements such as Cr, Cu, Cs, Sr, Hf, Pb, Th, Sb, Sc, Be, REE, U, Nb, V, Y and Zn were concentrated in B-RH, except Hf (Fig. 12) but Sn Ge, Ga, Ba, Ta, Zr, Rb and Ag were depleted. However TR and PH profiles lost Pb and Sc and gained in Cr, Hf, Zn; Rb and V. Overall, the only ones that increased in concentration throughout all of the profiles were REE, Cr, Nb, Th, Y and Zn. V-index values increased upwards in a similar manner for the three profiles, being twofold in PH and TR with respect to RH. Likewise, the loss on ignition (LOI) values increased upwards in the profile, from parent rock to B-horizons (Table A in the Appendix). The CIA index ranges from 48 to 52 in the bedrocks and 45 to 69 in the upper part of the profiles, with a generalised upwards increase, except for B-RH and BA-TR. The V index ranges from 0.4 to 1.7 in the bedrocks to 1.3 to 2.3 in the top horizons of the profiles, with a generalised increase upwards except for BA-TR (Table A in the Appendix).

REE distributions were significantly modified during weathering (Fig. 13 and Table D in the Appendix). An increase in REE concentrations was observed in all profiles. This increase was conspicuous for LREE in RH, whereas a slight decrease was also found for La in the TR profile, and for Pr, Nd, Sm, and Tb in the upper part of B-PH. Chondritenormalised plots showed a preferential concentration in LREE, both prominent in the PH and RH profiles (Fig. 14). None of the parent rocks show Ce anomalies but Eu differences were observed. Positive Ce anomalies were detected in the C and B-horizons of PH, Chorizon of RH and B-horizon of the TR profiles. An Eu negative anomaly remains in the PH profile and C-horizon in RH.

The Fe-Mn-P extractions performed for the trachy-rhyolite paleosol yielded a REE release efficiency of approximately 54% in Ce and about 17% for the rest of the REE when compared to the total amount of REE detected with total sample digestion (Fig. 15). The REE amount extracted from the poorly crystallised iron oxyhydroxides in soils (i.e. ferrihydrite and maghemite), ranged from 1.7 to 0.2% and from 1.4 to 6.3% respectively. For the more crystalline iron oxyhydroxides (i.e. goethite and hematite), REE extraction fluctuated from 92.6 to 98.4% of the total REE extraction.

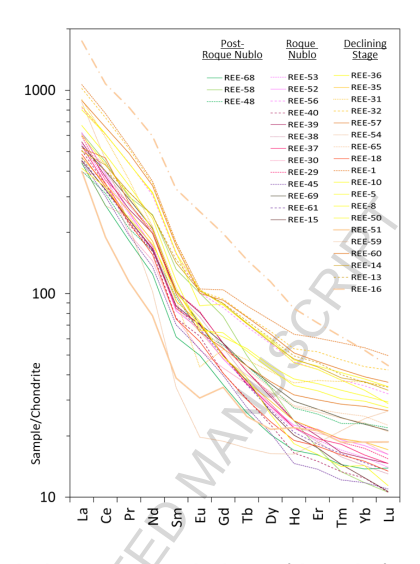


Figure 10. Chondrite-normalised REE distributions of the samples from the different stages on Gran Canaria (sample code and description in Tables 1a-b). Normalisation values after McDonough and Sun, 1995.



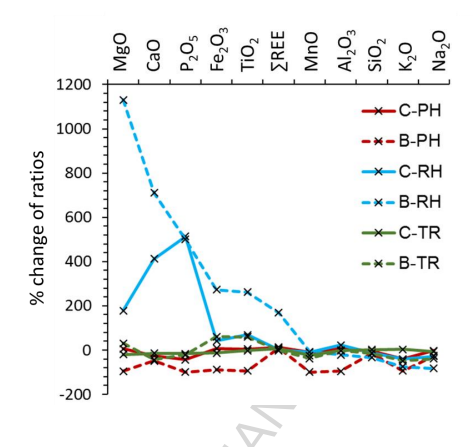


Figure 11. Geochemical plots of relative losses and gains of major oxides from the studied profiles (% change of ratios, equation [1]) in paleosol horizons with respect to bedrock. REE totals are included for comparison. PH: phonolite profile; RH: rhyolite profile; TR: trachyte profile; C, B, B' and BA refer to soil horizons.

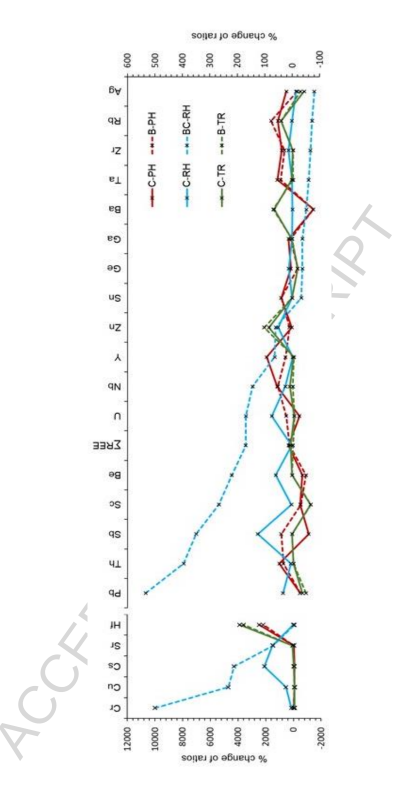


Figure 12. Geochemical plots of relative losses and gains of trace elements from the studied profiles (% change of ratios, equation [1]) in paleosol horizons with respect to bedrock. REE totals are included for comparison. PH: phonolite profile; RH: rhyolite profile; TR: trachyte profile; C, B, B' and BA refer to soil horizons.

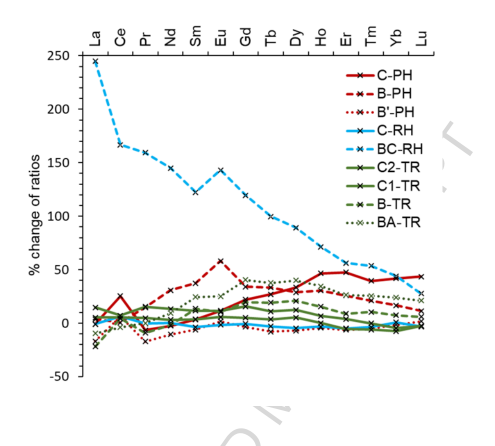


Figure 13. Geochemical plots of the relative losses and gains of rare-earth elements from the studied profiles (% change of ratios, equation [1]) in paleosol horizons with respect to bedrock. PH: phonolite profile; RH: rhyolite profile; TR: trachyte profile; C, B, B' and BA refer to soil horizons.



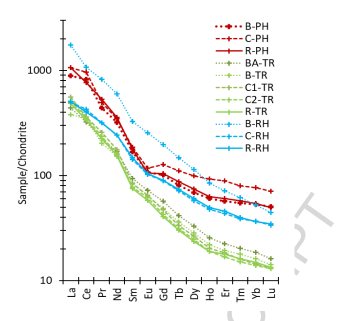


Figure 14. Chondrite-normalised REE distributions of the samples from the selected profiles (normalisation values after McDonough and Sun, 1995). Legend: PH: phonolite profile; RH: rhyolite profile; TR: trachyte profile; C, B, B', BA: soil horizons; R: bedrock.

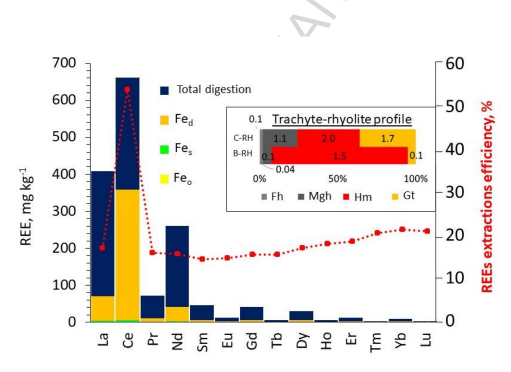


Figure 15. REE concentration of the Fe-Mn extractions (bar plots, left axis) and REE extraction efficiency (dotted red line, right axis) for C-RH and B-RH samples from the El Queso profile. The inset graph (upper right) shows the proportions of the different Fe oxides determined for each sample. Total digestion refers to the total amount of iron oxides, dithionite extract (Fe<sub>d</sub>) to hematite (Hm) plus goethite (Gt); sulphuric acid extract (Fe<sub>s</sub>) to maghemite (Mgh), and acid ammonium oxalate extract (Fe<sub>o</sub>) to ferrihydrite (Fh).

#### Appendix Tables

Table A. Major and rare-earth elements geochemistry of samples from the selected weathering profiles (major oxides in wt. %, REE in mg kg<sup>-1</sup>). Geochemical composition of the samples from the selected weathering profiles.

Profile			aga		El Queso trachy-rhyolite			Culata				
Lithology			honolit					DEE 40		trachyte		
Sample		C-PH	B-PH	REE-04 B'-PH	REE-13 R-RH	C-RH	B-RH		C2-TR	REE-21 C1-TR		
Horizon	R-PH 58	<u>С-РН</u> 56	<u>в-рн</u> 48		-	<u>C-RH</u> 59		<u>R-TR</u> 56	58	58	B-TR	BA-TR
SiO <sub>2</sub>			-	49 18	68		45				48	49
Al2O3	16	17	20		14	17	11	18	18	18 5	18	17
Fe <sub>2</sub> O <sub>3</sub>	6	6	6	7	4	6	15	6	5		9	10
MnO	0.5	0.4	0.3	0.4	0.2	0.2	0.3	0.1	0.1	0.2	0.1	0.1
MgO	1.2	1.2	1.2	1.6	0.5	1.5	6.5	0.9	0.7	0.8	1.2	1.3
CaO	1.2	0.9	3.0	2.1	0.5	2.8	4.4	2.3	2.1	1.8	1.3	1.5
Na2O	6.8	4.0	1.3	1.9	7.0	4.2	1.8	4.5	4.9	4.5	2.3	2.4
K2O	4.7	4.8	2.3	2.5	3.8	2.8	0.6	4.8	4.4	4.5	2.9	3.2
TiO <sub>2</sub>	1.0	1.1	1.0	1.2	0.8	1.3	2.8	1.5	1.6	1.5	2.5	3.0
P2O5	0.1	0.0	0.0	0.1	0.1	0.3	0.3	0.3	0.4	0.2	0.3	0.4
LOI	4	8	15	16	1	5	12	4	4	5	13	10
Total	99	99	99	100	100	100	99	99	99	99	99	99
La	253	251	210	263	120	119	414	115	121	132	90	104
Ce	472	592	503	475	247	263	659	206	217	221	215	198
Pr	49	46	41	57	29	29	76	21	22	24	19	21
Nd	163	159	146	213	111	111	272	71	73	81	70	78
Sm	27	27	25	36	22	21	49	11	12	12	13	14
Eu	6	7	6	9	6	6	14	3	3	4	4	4
Gd	21	25	20	28	18	18	39	8	9	9	10	11
Tb	3.2	4.0	2.9	4.2	2.7	2.6	5.3	1.1	1.1	1.2	1.3	1.5
Dy	18	24	17	24	15	14	28	6	6	7	7	8
Но	3.5	5.1	3.3	4.5	2.7	2.6	4.6	1.0	1.0	1.1	1.2	1.4
Er	9.7	14.3	9.1	12.2	7.4	7.0	11.5	2.9	2.7	3.0	3.1	3.6
Tm	1.4	2.0	1.4	1.7	1.0	1.0	1.5	0.4	0.4	0.4	0.4	0.5
Yb	8.7	12.4	8.6	10.2	5.9	6.0	8.5	2.4	2.2	2.3	2.6	3.0
Lu	1.2	1.8	1.2	1.4	0.9	0.8	1.1	0.3	0.3	0.3	0.4	0.4
∑REE	1036	1171	994	1139	588	601	1584	449	471	498	436	448
	-	13	-4	10	-	2	169	-	5	11	-3	0
Ce/Ce*	1.0	1.3	1.3	0.9	1.0	1.1	0.9	1.0	1.0	0.9	1.3	1.0
Eu/Eu*	0.8	0.8	0.8	0.9	0.9	0.9	1.0	1.1	1.1	1.0	1.0	1.0
LREE/HREE	22	17	22	19	16	17	25	31	33	33	26	23
, V-index	1.1	1.9	2.1	1.8	0.4	1.3	1.3	1.7	1.7	1.8	2.3	2.1
CIA-index	46	56	73	70	48	53	45	52	53	54	66	63

Oxides formulae and LOI values are shown in wt% (1 wt% equals 10,000 mg kg-1), and the REE in mg kg<sup>-1</sup>. REE vs.R (%) is the REE concentration related to REE concentration in the bedrock (R), in percentage.  $Ce/Ce^* = CeN/(LaN \times PrN)^{0.5}$ ;  $Eu/Eu^* = EuN/(SmN \times GdN)^{0.5}$ ; V index (Vogt's Residual Index)= (Al<sub>2</sub>O<sub>3</sub>+K<sub>2</sub>O)/(MgO+CaO+Na<sub>2</sub>O). CIA index (Chemical Index of Alteration) = 100\*(Al<sub>2</sub>O<sub>3</sub>)/(Al<sub>2</sub>O<sub>3</sub>+K<sub>2</sub>O+CaO+Na<sub>2</sub>O).

Table B. Minor and trace element geochemistry of samples from the selected weathering profiles (all values in mg kg<sup>-1</sup>, except Au in ppb). Geochemical composition of the samples from the studied weathering profiles.

Profile Lithology			aga honolit			E <b>l Queso</b> chy-rhyo		<b>Culata</b> trachyte				
Sample				REE-04		REE-14		RFF-18			REE-22	RFF-23
Horizon	R-PH	C-PH	B-PH	B'-PH	R-RH	C-RH	B-RH	R-TR	C2-TR	C1-TR	B-TR	BA-TR
Ag	3.1	4.6	3.8	2.9	2.1	1.8	<0.5	1.6	1.2	1.4	2.3	
As	16	15	< 5	< 5	7	5	4	< 1	1	2	5	< 5
Au	< 1	2	< 1	< 1	7	< 1	< 1	< 1	3	< 1	< 1	< 1
Ва	23	23	98	116	616	608	300	1690	1612	1702	1000	1123
Ве	12	15	24	13	5	8	16	4	4	4	4	4
Bi	< 0.1	0.1	< 0.4	< 0.4	< 0.1	< 0.1	0.3	< 0.1	< 0.1	< 0.1	< 0.4	< 0.4
Br	7.0	< 0.5	< 0.5	2.1	< 0.5	2.1	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5
Cd	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5
Со	< 0.1	< 0.1	2.0	4.0	< 0.1	6.6	70.2	9.8	6.6	10.6	12	22
Cr	31	23	30	40	12	29	1160	28	45	29	130	130
Cs	2.6	1.1	3.0	2.3	0.1	2.2	4.4	0.3	0.4	0.9	0.9	1.2
Cu	3	2	10	10	2	13	96	12	10	9	40	80
Ga	39	42	37	31	32	32	20	28	29	28	28	27
Ge	2	2	2	1	2	2	1	1	1	1	1	1
Hf	1.2	1.8	23.6	18.4	0.9	0.8	6.6	0.3	0.3	0.3	14.2	13.0
Ho	3.5	5.1	3.3	4.5	2.7	2.6	4.6	1.0	1.0	1.1	1.2	1.4
In	0.2	0.2	0.2	0.2	0.1	0.1	0.1	< 0.1	< 0.1	< 0.1	< 0.2	< 0.2
Мо	4	< 2	< 2	< 2	< 2	4	< 2	< 2	< 2	< 2	< 2	2
Nb	410	416	269	267	121	151	295	119	136	129	122	124
Ni	11	7	< 20	< 20	3	14	763	21	15	17	40	50
Pb	15	15	21	21	6	8	38	< 5	< 5	5	8	8
Rb	122	105	69	68	80	81	22	76	80	72	54	68
S	32	8	n.d.	n.d.	59	39	9	6	5	4	n.d.	n.d.
Sb	0.7	0.2	0.5	< 0.5	0.4	0.9	1.8	0.2	0.2	0.2	< 0.5	< 0.5
Sc	3	3	4	6	8	8	28	4	4	4	13	13
Se	< 0.5	5.8	< 0.5	< 0.5	< 0.5	4.3	< 0.5	< 0.5	< 0.5	< 0.5	n.d.	n.d.
Sn	7	7	5	5	6	6	4	2	2	2	2	2
Sr	71	67	104	98	26	418	410	1464	1483	1472	869	928
Та	24	26	17	16	9	9	4	7	7	7	7	7
Th	27	31	21	20	12	13	61	12	12	12	13	12
TI	0.2	0.1	0.2	0.2	<0.1	0.1	0.1	0.1	0.1	0.1	<0.1	<0.1
U	7	4	6	7	2	4	6	3	3	3	4	4
V	51	42	36	43	16	58	221	127	121	121	253	254
W	< 1	< 1	2	2	< 1	< 1	< 1	< 1	< 1	< 1	2	2
Y	102	158	82	123	74	73	121	29	29	30	31	39
Zn	319	295	290	330	169	256	270	121	104	117	60	90
Zr	1555	1676	1213	966	991	1124	330	739	749	757	769	725

Table C. The change of percentage ratio values (equation [1]) from B and C horizons with regards to the bedrock of the major and trace elements, including the  $\sum$ REE. These values are also shown as graphs in Figs. 11 and 12. The elements and oxides are ordered as in Figs. 11 and 12.

_	From Fig. 10	C-PH	B-PH	C-RH	BC-RH	C-TR	B-TR
	MgO	5	-95	178	1131	-21	30
	CaO	-25	-49	413	710	-15	-43
	$P_2O_5$	-43	-99	513	499	-17	-21
	Fe <sub>2</sub> O <sub>3</sub>	7	-88	42	273	-13	61
	TiO <sub>2</sub>	5	-94	68	261	-3	60
	∑REE	13	10	2	169	6	-3
	– MnO	-10	-100	-9	-8	-25	-38
	$AI_2O_3$	6	-96	23	-22	0	-2
	SiO <sub>2</sub>	-3	-17	-14	-34	2	-15
	Na <sub>2</sub> O	-41	-94	-41	-75	3	-48
	K <sub>2</sub> O	-3	-26	-28	-84	-7	-39
_	From Fig. 11	C-PH	B-PH	C-RH	BC-RH	C-TR	B-TR
	Cr	-25	3	153	9987	-71	-79
	Cu	-80	-70	550	4700	-76	-70
	Cs	-63	-13	2100	4300	-28	-67
	Sr	-36	-32	1508	1477	70	68
	Hf	2490	2195	5	-68	3900	3597
	V	17	42	263	1281	-52	-50
	Pb	-29	-29	33	533	-38	-50
	Th	47	32	5	395	-5	-5
	Sb	-60	40	125	350	0	0
	Sc	-30	-32	3	267	-67	-68
	Be	-38	-50	60	220	0	0
	U	-26	22	74	169	-8	-8
	Nb	55	52	25	144	9	-2
	Y	93	24	-1	64	-5	-6
	Zn	2	10	51	60	84	102
	Sn	40	40	0	-33	0	0
	Ge	5	-20	13	-38	-20	-20
	Ga	14	5	0	-38	2	0
	Ва	-77	-77	-1	-51	66	69
	Та	54	42	0	-60	2	-3
	Zr	38	28	13	-67	-2	-4
	Rb	52	77	1	-73	41	41
_	Ag	21	-18	-14	-81	-43	-30

Table D. The change of percentage ratio values (equation [1]) from B and C horizons with regards to the REE in the bedrock. These values are also shown as graphs in Fig. 13. The elements and oxides are ordered as in Fig. 13.

From Fig. 12	C-PH	B-PH	B'-PH	C-RH	BC-RH	C2-TR	C1-TR	B-TR	BA-TR
La	-1	4	-17	-1	245	5	15	-22	-10
Ce	25	1	7	6	167	5	7	4	-4
Pr	-7	15	-17	0	160	5	15	-9	0
Nd	-2	31	-10	0	145	3	13	-2	9
Sm	3	37	-6	-4	122	4	12	14	24
Eu	12	58	2	-2	143	6	11	10	25
Gd	22	34	-3	-1	120	5	16	20	41
Tb	27	33	-8	-3	100	4	11	19	38
Dy	33	29	-7	-5	89	5	12	21	40
Но	47	30	-4	-3	71	0	7	15	35
Er	47	26	-6	-5	56	-5	4	9	26
Tm	39	21	-5	-4	54	-6	0	11	26
Yb	42	17	-2	1	44	-7	-5	7	24
Lu	43	11	2	-3	28	-3	-2	6	21

#### 5. Discussion

#### 5.1. REE distribution

The redistribution of REE in weathered environments and soils may be controlled by different groups of phases, for instance, the aluminium-phosphate-sulphate (APS) minerals included in the alunite-jarosite group (Scott, 1987; Stoffregen and Alpers, 1987; Dill et al., 1995). However, all members of this group have been identified in the weathering profiles during the present work. A second mineral group that could also adsorb REE during weathering processes are clay minerals, as has been demonstrated by experimental studies (Coppin et al., 2002). Kaolinite may play an important role in removing REE from aqueous solutions by sorption (Coppin et al., 2002; Karakaya, 2009). Conversely, Galán et al. (2007) found a lack of correlation between the total concentration of REE and kaolinite content in the clay-sized fraction. In the same way, relationships between kaolinite and REE were not found on our Gran Canaria profiles. We reported the existence of secondary illite, smectite and kaolinite generated by weathering but they present a clear lack of REE, as shown by the relation of Ce and Si compositional maps (Figs. 8 and 9). This suggests that adsorption onto the surfaces of clay minerals is probably not a prevailing mechanism of REE retention in the Gran Canaria profiles studied. Analogous results were observed in the laterite profiles developed on the Neogene basalts from the northern coast of Hainan Island, South China (Ma et al., 2007), where kaolin-group minerals did not present significant REE content.

The REE also present a high affinity for P-compounds (Tyler, 2004) and these REEbearing minerals are very typical in weathered environments (Oelkers and Poitrasson, 2002; Marques et al., 2004; Santana et al., 2015). Indeed, on the basis of the mineralogical, textural and geochemical studies carried out in the present work, we can support such an affinity in the weathering profiles, where phosphates control the mineral distribution of REE (Fig. 6). Thus, we report monazite-(Ce) as a main REE phosphate (Fig. 7). This occurrence has been described in other paleosols and weathering profiles worldwide, such as the Las Mercedes bauxite deposit in the Dominican Republic (Torró et al., 2017), Greece bauxites (Eliopoulos et al., 2014) or the Quyang bauxites, China (Wang et al., 2010).

REE phosphate grains (monazite-(Ce)) show evidence for an authigenic origin: subhedral to anhedral crystals, irregular grain morphologies, rounded grain edges, oval to elongated

grain shapes and partial dissolution at its borders (Fig. 7). We interpret that this REE phosphate is inherited from the bedrock and was not formed during weathering. However, weathering processes were able to generate the alteration of primary monazite-Ce and the corresponding mobilisation of REE, forming secondary phosphates during incipient stages of alteration (Zapata and Roy, 2004). In Gran Canaria soils, these secondary minerals are represented by rhabdophane-(Ce) which develops concentrically around primary crystals of monazite-(Ce), defining foamy aggregates that generate "popcorn" textures (Fig. 7f). This REE mineral association is similar to those reported in other localities worldwide (Braun et al., 1993; Taunton et al., 2000; Harlavan and Erel, 2002, Reinhardt et al., 2018). Lower alteration indexes (V and CIA-index) observed in the RH paleosol profile are evidence of slight weathering (Table 2a), coupled with the highest REE concentrations. In terms of weathering, CIA-indexes were lower in Gran Canaria profiles than in the other well-developed soils studied (Condie et al., 1995; Panahi et al., 2000; Galán et al., 2007; Bao et al., 2008; Foley and Ayuso, 2013). As we previously suggested, Gran Canaria paleosols may not be strongly weathered. The occurrence of other secondary REE oxides developing concentric rims over inherited monazite-Ce and secondary rhabdophane-Ce (Fig.7 g-i) would be associated with significantly advanced stages of weathering (Table 4).

The existence of quartz in the upper layers of the paleosols (Fig. 5) has been related to an allochthonous provenance and can be used as a tracer of the aeolian inputs (Fernandez-Caldas et al., 1982; Mizota and Matsuhisa, 1995; Menéndez et al., 2007). In general, atmospheric deposition is a minor source of REE in soils (Wang et al., 2004) and it was confirmed on Gran Canaria in a previous study, in which current aeolian dust deposits were analysed and showed lower REE concentrations than in the paleosol profiles ( $\sum REE = 226 \text{ mg kg}^{-1}$ ; Menéndez et al., 2019). Overall, the allochthonous components scarcely contribute to the REE bulk geochemistry of the upper soil horizons (Condie et al., 1995).

In soils, significant REE enrichment seems to be mostly associated with Mn oxides. There is numerous evidence of Ce enrichment in relation to Fe-Mn oxide formation (Ma et al., 2007; Tripathi and Rajamani, 2007; Feng et al., 2011). Frequently, the nano-sized REE oxides are below the spatial resolution of the spectroscopy tools, thus being difficult to observe (Yusoff et al., 2013). The association between LREE and Mn-Fe oxides suggests that Ce oxidation may have been performed through oxide-assisted scavenging of dissolved Ce(III) (Ohta and Kawabe, 2001; Ma et al., 2007; Pourret and Davranche, 2013;

Yusoff et al., 2013). Since the incorporation of REE in the Fe–Mn-oxide structure is limited, sorption is one of the main mechanisms for REE scavenging onto Fe–Mn-oxides. Hence, REE contents are generally higher in amorphous Fe and Mn-oxides than in crystalline ones (Compton et al., 2003; Laveuf and Cornu, 2009). The REE tend to be higher in amorphous phases due to a possible expulsion during ageing and goethite crystallisation (Compton et al., 2003; Yusoff et al., 2013) or because the amorphous phase typically has a higher surface area for adsorbing ions. In the Gran Canaria profiles, the proportions of the poorly crystallised forms, ferrihydrite and maghemite, are significantly lower (8 to 24% of the total Fe concentration measured in C-RH and B-RH, respectively) than those of the crystallised phases of goethite and hematite (Fig. 15). Therefore, the bulk of the Fe-extractions of REE were mainly obtained from goethite and hematite forms (93 to 98% of the total REE measured in the Fe-extractions).

The presence of Fe oxides in paleosols suggests oxidising conditions. Thus, Ce is likely to be in the tetravalent state and, as a result, paleosols display a positive Ce anomaly in the trachyte and trachy-phonolite profiles (Table 2a and Fig. 15). According to Yusoff et al. (2013), the highest Ce(IV) concentrations are contained in cerianite and aluminophosphates, localised in the pores in Mn-rich domains. In the C-horizon, Ce(III) and Ce(IV) are mainly distributed in REE-bearing minerals of the rhabdophane group, found in pores and cracks (Janots et al., 2006). In soils, Ce(III) oxidation into Ce(IV) is typically associated with the dissolution/precipitation processes that operate during weathering (Mongelli, 1997; Ma et al., 2007; Sanematsu et al., 2013; Yusoff et al., 2013).

Iron can co-precipitate with elements such as Al, Ti, Mn, Ni, Zn and Cr, among others, in the form of poorly crystalline phases, such as ferrihydrite (Cornell and Schwertmann, 2003, and references therein). In paleosols and very old soils, ferrihydrite has generally been transformed into hematite and/or goethite incorporating those elements. In addition, Gálvez et al. (1999) observed how phosphorus can be incorporated (or occluded) in the structure of the crystalline phases formed from the precursor ferrihydrite. Thus, in the studied profiles, the REE-bearing phosphates (rhabdophane- (Ce) and LREE oxides) might have co-precipitated with Mn and Fe oxides.

#### 5.2. REE and weathering processes

The present study has revealed significant differences in REE concentrations when comparing the parent rocks (499 to 1036 mg kg<sup>-1</sup>) with the corresponding weathering

profiles (-42 to 0% change in the upper horizons and 2 to 169% in the lower ones). Even so, all lithologies present similar magnitudes in Ce (0.9 for negative anomalies and 1.1 to 1.3 for positive ones) and Eu anomalies (0.8 to 0.9 for negative anomalies and 1.1 for positive ones), as well as in LREE/HREE fractionation (from 16 to 33; Table 2a).

The increase of REE concentration during the weathering of the felsic volcanic rocks from Gran Canaria agrees with the results from other studies (Condie et al., 1995; Galán et al., 2007; Bao and Zhao, 2008; Yusoff et al., 2013; Calagari et al., 2015; Santana et al., 2015; Yanots et al., 2015). In contrast, depletion in REE during intense weathering of the parent rock has been reported by Beyala et al. (2009). There seems to be a preferential fixation of REE minerals in the early, rather than the more intense, weathering stages (Bao and Zhao, 2008). Primary mineral weathering occurs in all types of soils (Bockheim and Gennadiyev, 2000), thus dissolving accessory and essential phases (Nesbitt, 1979; Harlavan and Erel, 2002; Laveuf and Cornu, 2009). Moreover, an increase in REE concentration with depth has been described, a clear indication of leaching from the surface layers and deposition downward in the accumulation layers (Panahi et al., 2000; Ma et al., 2007; Chang et al., 2016). A variety of factors have been identified as influencing the retention of mobile REE, including preferential Mn (Ohta and Kawabe, 2001) and organic complexation (Taunton et al., 2000) and REE uptake by plant roots, linked to the absorption of Fe (Brioschi et al., 2013).

The REE mobilisation under supergene conditions involves the sequential release of REE and their redistribution along the weathering profile (Nesbitt, 1979). Europium and Cerium present two oxidation states at ground-surface conditions (Henderson, 1984; Laveuf and Cornu, 2009). The Ce<sup>+4</sup> fixation on secondary minerals (Bao and Zhao, 2008; Laveuf and Cornu, 2009) might be the reason for the noticeable efficiency found in the dithionite extraction for Ce (54% of the Ce obtained by total digestion; Fig. 15). This result suggests that about 54% of this Ce could be absorbed by the Fe-Mn and LREE oxides (Figs. 4 to 7). Otherwise, it is necessary to remark that in the volcanic rocks, the REE are preferentially and randomly hosted within the vitreous mesostasis, thus hindering the REE-bearing mineral formation (Wilson, 1989). In addition, part of the Ce in the profiles still remains in crypto-crystallised form.

Under reducing conditions, Eu can be mobilised by phosphate and carbonate complexation (Wood, 1990). Negative Eu anomalies are, in most cases, related to the

presence of feldspar or feldspar-derived alteration products, such as clay minerals (Galán et al., 2007).

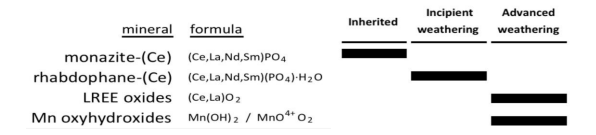
Evidence for REE mobilisation in weathering profiles includes fractionation of LREE to HREE. However, this fractionation was primarily inherited from the felsic volcanic rocks (Fig. 14). The same pattern has been identified in rhyolitic dykes by Benaouda et al. (2017), where volcanism-related hydrothermal processes are held responsible for this fractionation. There is no evidence of general hydrothermalism in the Gran Canaria rocks and initial REE fractionation should be mainly inherited from magmatic REE fractionation. This fractionation continues through the weathering process, being enhanced in the upper part of the profiles (Table 2b) and absorbed by amorphous Mn and Fe oxides (Laveuf and Cornu, 2009; Beyala et al., 2009).

In the studied profiles, we consider immobile elements to be those that become concentrated during the mineral transformation processes, despite their lack of mobility. Most of the elements are mobilised and transferred downwards through the weathering profile, as noticed in previous works (Ma et al., 2007). Nevertheless, TiO<sub>2</sub> and Zr are universally used for quantifying mass losses of major and trace elements in soils and other residual weathering products (Nesbitt, 1979; Brimhall and Dietrich, 1987). However, Bern et al. (2017) showed that Ti can be predominantly transported in the form of colloids. Other studies consider Th as an immobile element in tropical settings (e.g. Braun et al., 1993, 1998), but neither Th, TiO<sub>2</sub> nor Zr behave as immobile elements in the Gran Canaria profiles (Fig. 12). Physical weathering produces a loss of mass fragments, which leads to a preferential lixiviation of finer grain sizes. These processes subsequently involve a physical REE fractionation that is difficult to quantify. A positive percentage change should be detected for those elements but this trend depends on the profile (Figs. 11 to 13). Hence, none of the analysed elements appear to behave as if they were immobile (all positive values) except REE, Cr, Nb, Th, V, Y and Zn. Higher efficiency in their redistribution within the weathering profile probably implies that they were hosting the secondary minerals from the edaphic process. Secondary minerals (i.e. iron and manganese oxides) play an effective role in the transportation and concentration of a group of trace elements such as Co, Cr, and V, during weathering processes (Pokrovsky et al., 2006; Calagari et al., 2015). Rubidium is released during the break down of feldspar during weathering, thus explaining its depletion in RH but not in PH and TR paleosols.

The estimated weathering indexes (CIA and V-index, Table 2a) showed lower values in the RH profile, in concordance with REE decreasing with increasing degrees of weathering (Taunton et al., 2000, Chang et al., 2016), and in agreement with other studies of kaolinised profiles (Mongelli, 1993; Galán et al., 2007; Bao and Zhao, 2008; Yusoff et al., 2013). The CIA values of the upper parts of the profiles are lower than those obtained in other weathering profiles studied, for instance: CIA=70 in Condie et al. (1995), CIA=79 in Panahi et al. (2000), CIA=91 in Galán et al. (2007), CIA=98 in Bao and Zhao (2008), and CIA=99 in Foley and Ayuso (2013). It is worth pointing out that our profiles are paleosols and are, therefore, always truncated. In any case, the CIA value of the most preserved profile (BA-horizon, TR) only reaches 63. Thus, we suggest that, despite truncation, the hereby reported weathering paleosol profiles from Gran Canaria were not fully developed.

Lastly, the results of Nesbitt (1979) and Duddy (1980) are particularly significant in showing that REE could just be redistributed, rather than lost, within a weathering profile. Net losses and/or gains of specific REE are not observed in the TH profile nor the PH profile, or in other studies, such as that by Condie et al. (1995). Further studies of the RH profile should be conducted in order to understand its substantial (threefold) increase in REE content.

Table 4. Sequence of crystallisation for the REE-bearing mineral phases, depicting relationships between REE inherited phases (monazite-Ce), minerals generated during early stages of weathering (rhabdophane)-Ce, and LREE oxides associated with advanced stages of weathering. Note the close linkage between LREE oxides and Mnoxyhydroxides.



# **5.3. REE mineralisation and environmental and socio-economic limitations for its exploitation on Gran Canaria**

The occurrence of weathering and the corresponding formation of soils generated a significant enrichment of REE in these types of secondary formation, compared to the original primary rock (Angelica and da Costa, 1991; Foley and Ayuso, 2015). Therefore, secondary deposits should be considered as the most suitable targets when considering the potential exploration and corresponding exploitation of REE resources.

Gran Canaria is a limited territory (1,560 km<sup>2</sup>) with a population of 843,158 inhabitants, whose one major source of income is tourism (4.5 million tourists per year; ISTAC, 2017). The natural heritage (geodiversity and biodiversity) is valued and highly protected, mainly over the centre and south-western half of the island (49.4% of the island surface; source: Canary Government). In fact, both the local population and most of the services and facilities are concentrated in the north-eastern and eastern parts of the island and, hence, the proportion of protected areas here is much lower. Thus, most of the protected areas are concentrated in the south-west. The tourist industry is focussed along the southern and south-western coasts, where the Miocene mafic (shield phase) and felsic (decline phase) magmatic rocks from the first magmatic cycle mainly crop out. The few occurrences of Plio-quaternary magmatic rocks from the Roque Nublo and Post Roque Nublo Formations also crop out here, the latter being of minor interest to REE sources. Therefore, most of the key outcrops appear in protected areas, where extractive operations are prohibited and mining is not viable.

Our geochemical exploration study confirmed that the felsic magmatic rocks (phonolites, trachytes, rhyolites and syenites) and the associated paleosols of Gran Canaria can be considered to be a type of low-grade and bulk tonnage ore deposit for oceanic, intraplate volcanic islands, similar to that described by Jowitt et al. (2017) for highly fractionated rhyolites. The total volume of the felsic rocks in the alkaline declining stage (emitted between 14.1 and 7.3 Ma) is approximately 1,000 km<sup>-3</sup> (Schmincke, 1976, 1982; Schmincke and Sumita, 2010) and, assuming that these rocks have a density of 2,700 kg m<sup>-3</sup>, this yields 2,700 MT. Taking into account the mean 672 mg kg<sup>-1</sup> REE grade in this alkaline stage, the total tonnage of REE on Gran Canaria would be about 1,814 MT of potential REE resources. Accordingly, intraplate oceanic islands with alkaline felsic rocks

(lava flows, ignimbrites and plutonic stocks) can be considered as new potential, nonconventional rare-earth element resources.

Nevertheless, the gross amount of about 1.814 MT of potential REE resources must be recalculated downwards due to the limitations of the area occupied by the protected natural spaces, the dispersed rural population and the numerous coastal tourist infrastructure. For all these reasons, it would be necessary to carry out a detailed geochemical exploration in outcrops of felsic rocks and paleosols that are free of these environmental and socio-economic conditions.

The results obtained within this study, comprising the geochemical exploration of REE on this hotspot-derived volcanic island, can serve as a guide for future mineral prospects in these oceanic, intraplate environments. The felsic magmatic rocks related to the insular alkaline declining stage, which appear after the shield-building stages in all hotspot archipelagos, are the most suitable materials in which to carry out mineral exploration for REE. However, since islands are small geological features, mostly constrained by both environmental and socioeconomic issues, the potential exploitation of these resources might be restricted.

#### 6. Conclusions

Bulk major and trace geochemistry has been analysed in a set of felsic magmatic rocks and related paleosol profiles from different magmatic episodes on the island of Gran Canaria. A substantial mobilisation of REE, among other elements, has taken place during the course of the soil-forming processes. Our key findings can be summarised as follows:

(i) REE concentrations are significantly relevant in these types of differentiated magmatic materials, in terms of the estimated volume erupted during the alkaline declining stage of the island of Gran Canaria (1,000 km<sup>3</sup> over 7 Ma), which averages a REE sum of 672  $\pm$ 296 mg kg<sup>-1</sup>, an yttrium content of 57  $\pm$ 30 mg kg<sup>-1</sup>, and a LREE/HREE ratio of 17  $\pm$ 6;

(ii) REE concentrations vary from 2 to 169% in the lower horizons and are associated with phosphates (such as primary monazite-(Ce)), and with secondary soil weathering products linked to the co-precipitation of Fe-Mn oxides (such as rhabdophane-(Ce) and LREE oxides), but not to clay minerals, such as smectite, kaolinite or illite;

(iii) High LREE/HREE fractionation and slightly negative Eu anomalies were mainly inherited from the parent rocks, whereas positive Ce anomalies were enhanced by soil development;

(iv) REE were found to be evenly redistributed in newly-formed minerals (i.e. rhabdophane-(Ce) and LREE oxides), particularly in the case of cerium, mainly linked to crystalline oxides, such as goethite and hematite, rather than to amorphous phases.

As a corollary, on the basis of our results, this research supports the potential of felsic materials derived from the declining stage of an oceanic intraplate volcanic island, such as Gran Canaria, as being a type of low-grade and bulk tonnage REE ore deposit. However, environmental and socioeconomic factors may drastically limit the extension of island-based, REE-enriched deposits in terms of long-term exploitation.

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#### **Disclosure statement**

No potential conflict of interest was reported by the authors.

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