Mantle source heterogeneity, magma generation and magmatic evolution at Terceira Island (Azores archipelago): Constraints from elemental and isotopic (Sr, Nd, Hf, and Pb) data

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This work addresses the present-day (~100 ka) mantle heterogeneity in the Azores region through the study of two active volcanic systems from Terceira Island. Our study shows that mantle heterogeneities are detectable even when “coeval” volcanic systems (Santa Bárbara and Fissural) erupted less than 10 km away. These volcanic systems, respectively, reflect the influence of the Terceira and D. João de Castro plume end-members defined by Beier et al. (2008) for the Terceira Rift. Santa Bárbara magmas are interpreted to be the result of mixing between a HIMU-type component, carried to the upper mantle by the Azores plume, and the regional depleted MORB magmas/source. Fissural lavas are characterized by higher Ba/Nb and Nb/U ratios and less radiogenic 266Pb/204Pb, 143Nd/144Nd and 176Hf/177Hf, requiring the small contribution of delaminated sub-continental lithospheric mantle residing in the upper mantle. Published noble gas data on lavas from both volcanic systems also indicate the presence of a relatively undegassed component, which is interpreted as inherited from a lower mantle reservoir sampled by the ascending Azores plume. As inferred from trace and major elements, melting began in the garnet stability field, while magma extraction occurred within the spinel zone. The intra-volcanic system’s chemical heterogeneity is mainly explained by variable proportions of the above-mentioned local end-members and by crystal fractionation processes.

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1. Introduction

It has been demonstrated that the Earth’s mantle is heterogeneous (e.g., Gast et al., 1964) at several scales, from mm3 to 104 km3. This claim is supported by: 1) the study of mantle xenoliths (e.g., Bonadiman et al., 2005; Harte, 1987; Mattielli et al., 1999) and of outcropping mantle portions (e.g., Brunelli and Seyler, 2010; Reisberg and Zindler, 1986) preserving evidence for millimeter- to meter-scale heterogeneities caused by metasomatic processes; 2) the significant heterogeneity depicted by oceanic basalts (e.g., Hofmann, 2003); and 3) the existence of large mantle domains with peculiar characteristics, as shown by the Dupal anomaly (Dupré and Allègre, 1983; Hart, 1984).

The Azores islands, located on the Azores Plateau and close to the Mid-Atlantic Ridge, are characterized by significant elemental and isotopic variations, which are evident either from the study of a single island (Terceira: Dupré et al., 1982; S. Miguel: Beier et al., 2007; Haase and Beier, 2003; S. Jorge: Hildenbrand et al., 2008; Millet et al., 2009) or from regional studies (e.g., Beier et al., 2008, 2010; Moreira et al., 1999; White et al., 1979; Widom and Shirey, 1996). At Northeast the Azores Plateau is bordered by the Terceira Rift (Fig. 1A), which has been interpreted as an intra-oceanic spreading system (see Vogt and Jung, 2004 and references therein). Beier et al. (2008) identified the presence of several distinct compositional end-members along this structure, providing evidence for significant mantle source heterogeneity and focused magmatism with no significant melt or source interaction between different volcanic systems.

Although some authors have asserted that the Azores resulted from an anomalously volatile-enriched upper mantle domain (e.g., Bonatti, 1990), the conjunction of noble gas signatures with data from seismic tomography endorses the idea of a mantle plume influence on the Azores magmatism (Jean-Baptiste et al., 2009; Madureira et al., 2005; Moreira et al., 1999; Silveira et al., 2006, 2010). However, the...
relative role of a mantle plume and shallow level-reservoirs as the
cause for such variability has been the locus of an intense debate (e.g.,
Millet et al., 2009; Moreira et al., 1999; Widom et al., 1997).

Through a finite-frequency seismic tomography study, Yang et al.
(2006) demonstrated that the low-velocity anomaly is centered NE
of Terceira Island and is de
icted to the SE by asthenospheric
flow. This and the fact that some of the more unradiogenic He ratios for the
Azores islands were obtained in olivine phenocrysts from Terceira,
where 21Ne/22Necorr = 0.052 (Madureira et al., 2005), suggests that the
Azores mantle plume can be particularly well sampled by Terceira lavas.

In this study, elemental and isotope data (Sr, Nd, Pb, and Hf) were
obtained for Terceira’s ma
lavas. They were collected from the two
main active volcanic systems linked to the Santa Bárbara central
volcano and to the fissure zone, the latter being considered the
emerged record of the Terceira Rift. Based on available K/Ar and Ar/Ar
ages (Calvert et al., 2006; Féraud et al., 1980) the sampled lavas are
expected to be younger than 100 kyr. The obtained data will be used
to address the following issues: 1) to confirm and better constrain the
occurrence of intra-Terceira Island heterogeneity referred by Dupré
et al. (1982); 2) to address intra-volcanic system chemical variability;
3) to compare the Terceira isotopic signatures with those described by
Beier et al. (2008) for the Terceira Rift as a whole; 4) to constrain the
melting processes and the depth of magma extraction; and 5) to
discuss the origin of the main contributors to the Terceira source(s)
and their importance in the context of the Azores.

2. Geological background and sampling strategy

The Azores archipelago is located in the North Atlantic (37°–40° N)
near the RRR-type triple junction where the Eurasia, Nubia (Africa)

Fig. 1. A) Bathymetric chart of the Azores plateau and location of the Azores islands (adapted from Lourenço et al., 1998). Am, EA, Nu: America, Eurasia and Nubia lithospheric plates, respectively; EA: East Azores Fracture Zone; MAR: Mid-Atlantic Ridge; TR: Terceira Rift. Inset: location of the Azores Archipelago in the North Atlantic; B) Volcanic systems interpreted from the Terceira exposed lavas (adapted from Madeira, 2005; Nunes, 2000). CPC — Cinco Picos Caldera; GMC — Guilherme Moniz Caldera; SBC; Santa Barbara Caldera; PAV — Pico Alto Volcano; F — Fissure zone; AC — Algar do Carvão. The map shows the location of samples selected for geochemical analysis. The site of the Serreta submarine eruption (1998–2001) is indicated by the black dot offshore. Lavas grouped inside parentheses were sampled vertically in lava sequences forming the actual cliff. Radiogenic isotopic data (Sr, Nd, Pb, Hf) were obtained for underlined samples (see also Table 2). The Algar do Carvão lava flow has been dated at 2115 ± 115 a (Zbyszewski et al., 1971) and, more recently, at 1730 ± 70 a (Forjaz et al., 2004).
and America plates meet (Fig. 1A). The archipelago is composed of nine volcanic islands emerging from the Azores Plateau (400,000 km² of elevated oceanic crust, underlined by 2000 m isobaths; Lourenço et al., 1998) that extend to both sides of the Mid-Atlantic Ridge (MAR). To the east of the MAR, the northern boundary of the Azores Plateau is underlain by a 550 km long hyper-slow (~4 mm/a) rift system known as the Terceira Rift. The onset of the extensional regime in the Azores domain may have begun 20 Ma ago (Luís and Miranda, 2008) with focused spreading along the Terceira Rift occurring essentially in the last 1 Ma (Vogt and Jung, 2004). Its present-day extensional regime is confirmed by GPS and laser measurements on the islands (e.g., Miranda et al., 1998). The islands of S. Miguel, Terceira and Graciosa, as well as the submarine D. João de Castro Bank are all located along this rift and separated by deep volcanic zones that reach 3200 m bsl at the Hirondelle basin (Beier et al., 2008; Lourenço, 2007; Vogt and Jung, 2004). The Terceira Rift is considered the boundary between the Eurasia and Nubia lithospheric plates, which was previously located to the southwest as suggested by the elongation of S. Jorge Island, the Faial-Pico alignment and other elongated ridges with similar orientations (Vogt and Jung, 2004; see Fig. 1A).

The morphology and volcano-stratigraphy of Terceira Island indicate the existence of five main volcanic systems (Fig. 1B; Madeira, 2005; Nunes, 2000): Cinco Picos, Guillerme Moniz, Pico Alto, Santa Bárbara and one associated to the fissure zone (hereafter referred as the Fissural volcanic system). The latter connects the Santa Bárbara and Guillerme Moniz calderas and is believed to extend toward the SE coast of Terceira Island (Self, 1976). Cinco Picos comprises the oldest volcanic center (older than 370 ka; Calvert et al., 2006) and is currently dominated by the largest caldera of the Azores archipelago (~7 km diameter), which is partially filled with lavas erupted from the fissure zone. The Fissural, Santa Bárbara and Pico Alto systems are considered volcanically active, but for the latter only highly evolved lavas and pyroclastic deposits (pantellerites and comendites) outcrop at the surface (Gertisser et al., 2010; Mungall and Martin, 1995; Self and Gunn, 1976). The eruption of similar high-silica lavas also occurred during the last stage of Santa Bárbara’s evolution with the formation of ~80 lava domes (Madeira, 2005; Self and Gunn, 1976). For this volcano, Calvert et al. (2006) estimated the beginning of sub-aerial activity within the past 100 kyr.

For Terceira Island, Self (1976) reported the occurrence of 116 eruptions before the generation of a 23,000 year old ignimbrite. However, in historical times only three basaltic eruptions occurred restricted to the fissure zone. The last eruptions (1998–2001) took place offshore 10 km to the west of the island at the Serreta submarine ridge, where a previous eruption occurred in 1867 closer to the island (Gaspar et al., 2003). Our basaltic sample, PMT-Serr-2, was collected in April 1999 during one of the major submarine volcanic activity periods that brought lava balloons to the sea surface. This study is focused on mafic lavas from the Fissural and Santa Bárbara volcanic systems (See Fig. 1B for location) to assess the “present-day” Terceira mantle source(s) and magma formation processes.

3. Analytical procedure

Whole-rock analyses of major and trace elements were performed in Canada by Activation Laboratories using ICP-OES and ICP-MS, respectively. Alkaline dissolution with lithium metaborate/tetraborate followed by nitric acid was used for the analyses, except for Ni, Cu, Zn and Cd determination, for which multi-acid digestion (hydrofluoric followed by a mixture of nitric and perchloric acids) was carried out. Reproducibility was generally better than 5% for both major and trace elements and can be better evaluated at Supplementary Material (Table S1) where we also present analyses of standard reference materials (BIR-1,W-2a and DNC-1). More information on the analytical procedure can be found at http://www.actlabs.com.

The isotopic analyses of 12 representative samples from the Fissural and Santa Bárbara systems were performed at the Department of Earth and Environmental Sciences of the Université Libre de Bruxelles (Nd, Hf and Pb isotopes) and at Pacific Centre for Isotopic and Geochemical Research at the University of British Columbia (Sr isotopes). The samples were dried at 40 °C and crushed in an agate mortar under an extracting hood. The entire acid digestion process and the analyte separation on ion-exchange resin were performed in a class 100-laminar air flow cabinet. About 200 mg of powder was dissolved in closed Savillex® beakers using HF–HNO₃ (130 °C, 48 h) followed by evaporation, the addition of 6 N HCl, a second evaporation and final dissolution in 0.5 N HBr. Lead separation was performed using successive acid elution on an anionic resin (AG1-X8) column (for further details, see Weis et al., 2006), Sr, REE and Hf were isolated using a column loaded with 18.5 ml of Dowex AG50W-X8 100–200 mesh cation exchange resin. Nd was recovered by loading the REE fraction onto a column with HDEHP-coated Teflon. Hafnium was separated following the slightly modified two-column procedure described in Blichert-Toft et al. (1997), involving anion and cation exchange columns (see Mattielli et al., 2002; Weis et al., 2006). Measurements of the total procedural Pb, Nd, Sr and Hf blanks gave ca. 10 pg, 28 pg, 400 pg and 23 pg, respectively. The isotopic compositions of Pb, Nd and Hf were determined on a Nu plasma MC-ICP-MS (multiple collector inductively coupled plasma mass spectrometer). At the beginning of every analytical sessions, collected analyte (Pb or Nd or Hf) elution fractions were re-dissolved in 100 µl of concentrated HNO₃ evaporated and finally dissolved in 1.5 ml of HNO₃ 0.05 N. Tl was added to each Pb sample and standard to control instrumental mass fractionation. Solutions were prepared to obtain a Pb/Tl rate of 4 or 5 and a signal of 100 mV in the axial collector (200 Pb) as well as to reach the Pb and Tl concentrations of the standard (200 ppb of Pb and 50 ppb of Tl). Standards were systematically run between every two samples to monitor the instrumental mass bias during the analysis session, resulting in

\[ ^{143} \text{Nd}/^{144} \text{Nd} = 0.511960 \pm 0.000002 (2\sigma) \]

for the JMC475 Hf standard. All Pb measurements were automatically corrected according to the Tl mass fractionation and then by using the standard bracketing method with the recommended values of Galer and Abouchami (1998) and Abouchami et al. (2000). The Nd and Hf isotopic measurements were internally normalized to

\[ ^{143} \text{Nd}/^{144} \text{Nd} = 0.7219 \]

and

\[ ^{176} \text{Hf}/^{177} \text{Hf} = 0.282149 \]

(2\sigma, n = 31) for the JMC475 Hf standard.

In our study, each session of analyses starts by a “batch run” of 10 analyses of the standard solutions in order to monitor the optimization (stability and sensitivity) of the machine. In addition, during this batch run, the reproducibility and accuracy of the values were checked. The quality of our analytical procedure (i.e. chemical preparation procedures and isotopic measurements), and especially the reproducibility of isotopic measurements, can be assessed by sample replicate and duplicate analyses, which is better than 49 ppm, 19 ppm, 25 ppm for

\[ ^{143} \text{Nd}/^{144} \text{Nd} \]

and

\[ ^{176} \text{Hf}/^{177} \text{Hf} \]

respectively. The Sr isotopic ratios were analyzed on a Finnigan Triton Thermo-Ionization Mass Spectrometer (TIMS). The NBS987 Sr standard was run after every 2 samples, giving a mean value of

\[ ^{87} \text{Sr}/^{86} \text{Sr} = 0.710592 \pm 0.0027 \]

for 6 measurements. The raw data were corrected for mass fractionation using

\[ ^{86} \text{Sr}/^{88} \text{Sr} = 0.1194 \]

In our study, each session of analyses starts by a “batch run” of 10 analyses of the standard solutions in order to monitor the optimization (stability and sensitivity) of the machine. In addition, during this batch run, the reproducibility and accuracy of the values were checked. The quality of our analytical procedure (i.e. chemical preparation procedures and isotopic measurements), and especially the reproducibility of isotopic measurements, can be assessed by sample replicate and duplicate analyses, which is better than 49 ppm, 19 ppm, 25 ppm for

\[ ^{143} \text{Nd}/^{144} \text{Nd} \]

and

\[ ^{176} \text{Hf}/^{177} \text{Hf} \]

respectively. For

\[ ^{206} \text{Pb}/^{204} \text{Pb} \]

and

\[ ^{208} \text{Pb}/^{204} \text{Pb} \]

the external reproducibility is ~180 ppm. Additional information on the analytical procedures, as well as on the usual accuracy and reproducibility obtained at the Université Libre de Bruxelles and the University of British Columbia, can be found in Weis et al. (2006).

Minerals from representative samples were analyzed with a JEOl JXCA 733X electron microprobe at Centro de Geologia da Universidade de Lisboa (CeGUL). The microprobe operated with a 5-µm beam diameter, an accelerating voltage of 15 kV (18 kV for opaque minerals) and a beam current of 25 nA. Natural mineral and pure metal (Cr, V and Ni) standards were used. Precision, as indicated by replicate determinations on an in-house standard, was better than 2% for the major elements.
4. Results

4.1. Petrography and chemical composition of the main mineral phases

The textures of mafic lavas from the Santa Bárbara and Fissural volcanic systems are predominantly porphyritic to seriate with major phenocryst phases (olivine, clinopyroxene and plagioclase) invariably surrounded by a microcrystalline matrix dominated by plagioclase and opaque minerals (Fig. 2). Opaque microphenocrysts are also present in most Santa Bárbara lavas. For the less evolved lavas (Mg#68), our petrographic observations indicate olivine as the liquidus phase, which was early joined by clinopyroxene and then plagioclase in the more evolved samples. The proportion of phenocrysts in Fissural lavas ranges between 1 and 15%, significantly less than the maximum of 25–30% (of the rock volume) observed for Santa Bárbara lavas.

Concerning Fissural lavas, plagioclase phenocrysts are nearly absent in rocks outcropping in the eastern area of Terceira Island and are considerably more abundant in lavas that erupted between the Santa Bárbara and Guilherme Moniz calderas, where more evolved lavas occur. This strongly suggests the presence of more mature magma chambers in the central part of the island.

In Santa Bárbara lavas, plagioclase is always the dominant phenocryst commonly forming glomeroporphyritic clusters. Olivine phenocrysts are poorly represented and often exhibit iddingsite along their rims and/or intra-granular fractures. The rarity of olivine is considered as a result of the evolved character of lavas (Mg#48) in this volcanic system. Biotite and/or kaersutite crystals can also be observed in some Santa Bárbara lavas occurring both as a matrix phase and as filling vesicles.

Representative chemical analysis of the major phenocryst phases in Fissural and Santa Bárbara lavas can be found in Supplementary Material (Table S2).

4.1.1. Olivine

Olivine phenocrysts with the highest forsterite (Fo) content were analyzed in Fissural lavas (88–90% Fo) sampled in the eastern area of Terceira Island, which also display higher measured NiO concentrations (0.3–0.4 wt.%; Supplementary Material, Table S2A). Forsterite ranges from 90% to 72% in phenocryst cores from these lavas and from 85% to 56% in olivine phenocrysts from the Santa Bárbara volcanic system.

For both volcanic systems there is a crude positive co-variation between the CaO concentration (0.15 to 0.38 wt.%) and %Fo in olivine phases (not shown). If the crystallization of the higher magnesium olivines occurred at deeper levels, this would be contrary to what is expected from the inverse correlation between pressure and the $D_{Ca^{2+}/Al}$ partition coefficient (Hirschmann and Ghiorso, 1994). According to petrographic observations, we explain such correlation as the result of CaO depletion stemming from coeval clinopyroxene crystallization (Libourel, 1999; Jurewicz and Watson, 1988). For some lavas, zoned compositional patterns display a large central plateau characterized by relatively high Fo contents, which is rimmed by a thin border marked by significantly lower MgO contents. This is interpreted as reflecting rapid MgO depletion in magmas as a consequence of sudden microlyte nucleation during mesostasis crystallization (e.g., Hildner et al., 2011).

4.1.2. Clinopyroxene

Clinopyroxene is a major phase in Terceira lavas, occurring as phenocrysts, microphenocrysts and matrix crystals. Ubiquitous phenocryst compositional zoning can be readily interpreted from crossed nicois observations and, in most cases, also from the darker brown color of the outer crystal rim. Terceira pyroxenes are dominated by quadrilateral components and, following the classification of Morimoto et al. (1988), can be considered as diopsides and augites. The highest Mg# of clinopyroxene cores is identical to that obtained for olivine, but their range is smaller than what was observed for olivine phenocrysts (Mg# range from 0.75 to 0.90 for Fissural lavas and from 0.73 to 0.80 for...
Santa Bárbara lavas; Supplementary Material, Table S2B). We consider the overlapping of the Mg# for the most magnesium pyroxenes and olivines as the result of equilibrium crystallization between those two phases given the similarity between their $K_{min}$ ratios (see Tappe, 2004). Fissural and Santa Bárbara clinopyroxene phenocrysts are distinguishable by higher Cr concentrations (up to 1.2 wt.% and Al/Ti ratios (4.57) of the former compared with those observed for Santa Bárbara pyroxenes (Cr$_2$O$_3$ up to 0.6 wt.%; Al/Ti up to 2.86), reflecting the less evolved character of the Fissural lavas.

4.1.3. Plagioclase

Plagioclase is the dominant phenocryst phase of Santa Bárbara lavas and is also important in the most evolved Fissural lavas sampled in the central area of Terceira Island. The analyzed phenocrysts display a wide range in composition from 40% to 85% An (Supplementary Material, Table S2C). Groundmass plagioclase laths extend the compositional range down to 30% An. Multiple and complex zoning surrounding resorbed cores is a common feature in Santa Bárbara plagioclase phenocrysts. Several profiles along Santa Bárbara plagioclase phenocrysts also display oscillatory zoning, which can be interpreted as a probable result of mixing between melt fractions with different evolutionary stages.

4.1.4. Oxides

Opaque minerals can be found as inclusions in phenocryst phases, as (micro)phenocrysts in Santa Bárbara lavas and as crystals from the microcrystalline matrix. Inclusions and (micro)phenocrysts are represented by minerals from the spinel group, mainly chromite and titanomagnetite (Supplementary Material, Table S2D). Ilmenite is the dominant oxide in the microcrystalline matrix both in Fissural and in Santa Bárbara lavas but can also occur as (micro)phenocrysts in Santa Bárbara lavas or as exsolution lamellae in titanomagnetite phenocrysts.

4.2. Whole-rock chemistry

Whole-rock data were obtained from unaltered samples, usually with less than 10% of phenocrysts to be representative of magmatic liquid compositions (Table 1). However, some Santa Bárbara lavas display higher volume proportions of plagioclase phenocrysts (up to 35%), which suggests plagioclase accumulation. This interpretation is strengthened by a positive correlation between the Eu/Eu* and Sr/Nd ratios (not shown), and, accordingly, those samples were omitted in this discussion and did not contribute to our conclusions.

4.2.1. Major elements

Similarly to other Azores Islands and OIB, the elemental data obtained for the Terceira Fissural and Santa Bárbara lavas are distinct from Atlantic MORB. Most primitive lavas (here considered to have MgO > 8 wt.%) present higher TiO$_2$ and P$_2$O$_5$ contents than typical N-MORB (2.3–3.0 and 0.35–0.86 wt.% versus 1.19–1.77 and 0.10–0.16 wt.%, respectively; e.g., Sridhar and Ray, 2003) and Na$_2$O/K$_2$O ratios (2.8 to 4.3) similar to the values reported for other Atlantic islands (e.g., Davies et al., 1989; Schminke, 1973). This more SiO$_2$-undersaturated tendency of Terceira lavas partially explains the common occurrence of ilmenite, a mineral phase whose crystallization is strongly inhibited by the significant Ti incorporation in clinopyroxene in SiO$_2$-undersaturated rocks (e.g., Toplis and Carroll, 1995). The high ilmenite abundance allows the distinction of Terceira lavas even from moderate alkaline lavas such as those of Madeira Island (see Mata and Munhá, 2004).

4.2.2. Trace elements

Terceira most primitive lavas (MgO > 8 wt.%) are characterized by fractionated REE patterns with primitive mantle normalized La/Yb and Tb/Yb ratios ranging from 7.4 to 11.5 and 2.2 to 2.7, respectively. Variable La/Yb and Tb/Yb ratios for basaltic samples are suggestive of differences in the degree of partial melting and in the amount of residual garnet on magma genesis (e.g., George and Rogers, 2002). For Terceira Island, the presence of garnet in the mantle source is also suggested by the low concentrations of HREE (Yb: 1.87–2.67 ppm; Lu: 0.25–0.37 ppm) in the most primitive lavas when compared with the mean values obtained for N-MORB (3.05 ppm and 2.37 ppm respectively; Sun and McDonough, 1989). Moreover, the presence of residual garnet is supported by the $^{230}$Th excesses up to 26% over $^{232}$U determined by Turner et al. (1997) for Terceira lavas.

Looking at primitive mantle normalized diagrams, it is possible to recognize for both systems the enrichment of Nb and Ta relative to LREE and HREE and the presence of K negative anomalies (Fig. 4). However, Fissural lavas are almost ubiquitously characterized by a positive Ba anomaly that is never observed in Santa Bárbara lavas, suggesting that magmas from these two volcanic systems correspond to distinct magmatic series. This assertion is also strengthened by the different incompatible trace elements and isotopic ratios found for Fissural and Santa Bárbara lavas (see Fig. 5 and Section 4.3). Santa Bárbara lavas present trace element patterns similar to those described for HIMU-type basalts, while Fissural samples, with higher Ba/Nb, Ba/Th and Rh/Th ratios, are more akin to EM I-type magmas (see insets in Fig. 4 and Weaver, 1991).

The submarine lavas erupted in 1999 along the Serreta ridge, at the prolongation of the fissure zone, represents a special case since they display trace element characteristics similar to those erupted from the Santa Bárbara central volcano (Fig. 5). This suggests that the Santa Bárbara mantle source is presently being sampled by eruptive events at the fissure zone.

4.3. Isotopic data

The Sr, Nd, Pb and Hf isotopic data obtained for some Fissural and Santa Bárbara lavas are summarized in Table 2. Considering the very young age of the studied lavas (~0.1 Ma; Calvert et al., 2006) and the very long half-life of the radioactive parent isotopes (~0.703×10$^{10}$ y), the analyzed values are equal to the calculated initial isotopic ratios.

In the Sr–Nd isotope diagram, Terceira samples are plotted in the mantle array in the vicinity of the HIMU and DMM components (Fig. 6A). They display a restricted range of isotopic ratios ($^{87}$Sr/$^{86}$Sr; $^{142}$Nd/$^{144}$Nd: 0.512908–0.512980; $^{143}$Nd/$^{144}$Nd: +3.5 to +6.7), showing evidence for the time-integrated depletion of Rb and Nd relative to the more compatible Sr and Sm, respectively.

In Pb isotope diagrams (Fig. 6B and C), the studied Terceira samples plot slightly beneath the NHRL. In Sr–Nd isotope diagrams, it is evident that Fissural and Santa Bárbara lavas display distinct Pb isotope signatures (see also Dupré et al., 1982) with the latter being more radiogenic ($^{207}$Pb/$^{204}$Pb from 19.52 to 19.99; $^{206}$Pb/$^{204}$Pb from 15.614 to 15.620; $^{208}$Pb/$^{204}$Pb from 39.226 to 39.266) than Fissural lavas ($^{207}$Pb/$^{204}$Pb from 19.339 to 19.486; $^{206}$Pb/$^{204}$Pb from 15.529 to 15.617; $^{208}$Pb/$^{204}$Pb from 38.872 to 39.204). Moreover, looking at the fissure zone, there is a tendency for a
Table 1

Whole-rock elemental analyses of representative samples. LOI: loss on ignition. Mg#: atomic 100 × [Mg/(Mg + Fe2+)] using FeO calculated from Fe2O3 according to the procedure of Midlenost (1989). Trace element concentration was determined by ICP-MS. Bold and underlined Pb concentration was determined by TD-MS. See text for further analytical details. Lithotype determined according Le Bas et al. (1986).

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(continued on next page)
westward increase in Pb isotopic ratios. The most unradiogenic sample was collected in the eastern part of the island (PMT-15 near Porto Judeu, see Fig. 1B for location), and more radiogenic values were obtained for the 1761 historic lava (PMT-27) and the Serreta sample, which are more similar to those values obtained for Santa Bárbara lavas (see insets of Fig. 6A and C).

This work presents the first Hf isotopic data for Terceira Island. 176Hf/177Hf ratios range between 0.283002 and 0.283099 (εHf: +8.1 to +11.6) with some samples displaying more depleted signatures than those reported for S. Miguel (εHf: −0.2 to +8.7) and Pico (εHf: +7.2 to +9.5) (see Beier et al., 2007; Elliott et al., 2007). Terceira samples tend to be less radiogenic than MORB lavas and plot slightly below the mantle array (Fig. 6D).

5. Discussion

Distinct elemental and isotopic signatures were presented for Fissural and Santa Bárbara lavas. However, before assessing the role of heterogeneities in the Terceira mantle source, it is important to evaluate the geochemical changes imprinted on primary magmas during their ascent to the surface.

5.1. Fractional crystallization

The distribution of mafic Terceira lavas in the TAS diagram (Fig. 3) allows for the recognition of two main groups: one formed by lavas from the Fissural volcanic system, plotting in the basaltic and hawaiitic compositional fields, and another formed by the lavas from the Santa Bárbara volcanic system, which display a much wider dispersion (from basalts to benmoreites). This compositional range is even larger when we consider the high volume of comenditic lava domes and coulées formed.

Table 1 (continued)

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Fig. 3. Terceira lavas plotted on a TAS diagram (Le Bas et al., 1986). The black line represents the alkaline and sub-alkaline divider from Irvine and Baragar (1971). With the exception of sample PMT-65 (K-Trachybasalt), all samples plotted in S1, S2 and S3 are classified as hawaiites, mugearites and benmoreites, respectively, given that (Na2O − 2.0 ≥ K2O).

This work presents the first Hf isotopic data for Terceira Island. 176Hf/177Hf ratios range between 0.283002 and 0.283099 (εHf: +8.1 to +11.6) with some samples displaying more depleted signatures than those reported for S. Miguel (εHf: −0.2 to +8.7) and Pico (εHf: +7.2 to +9.5) (see Beier et al., 2007; Elliott et al., 2007). Terceira samples tend to be less radiogenic than MORB lavas and plot slightly below the mantle array (Fig. 6D).
during the latest eruption events of the Santa Bárbara system (Self, 1976). This difference indicates the existence of a more mature magma chamber system feeding the Santa Bárbara volcano, which is also in accordance with the mixing of melt fractions at different evolutionary stages as interpreted from plagioclase phenocrysts (see Section 4.1.3). Despite the more primitive character shown by Fissural lavas, the presence of samples that can be considered primary magmatic liquids is scarce. Following the criteria of Frey et al. (1978), only the faintly porphyritic lavas sampled in the eastern area (PMT-8, PMT-14 and PMT-15, see Fig. 1B for location) represent Terceira primary-like magma compositions (Mg#: 71.8–72.8; Ni: 267–283 ppm). Their MgO contents (MgO: 11.34–11.83 wt.%) are also in agreement with the proposal of Beier et al. (2008) for the composition of primary magmas erupted along the Terceira Rift (MgO ~12.5 wt.%).

Based on distinct isotopic signatures and incompatible element ratios we demonstrated that the Fissural and Santa Bárbara volcanic systems are not comagmatic. Accordingly, we consider the apparent single liquid line of descent shown in Fig. 7 as representing the evolution of a series of broadly similar, but not cogenetic, magma batches, evolving by similar processes in their travel to the surface.

Least-squares mass balance calculations considering pairs of less to more evolved magmas point to an amount of fractional crystallization between 50% and 65% to generate the observed range for each magmatic series. For the Fissural series, the lowest sum of squared residuals (0.246) was achieved by fractionating olivine (~25.5%), clinopyroxene (~53.7%), plagioclase (~14.8%) and Cr-spinel (~6.0%). Plagioclase and titanomagnetite become dominant over olivine and Cr-spinel when considering the most evolved samples. Concerning Santa Bárbara, the lowest sum of squared residuals (1.251) implies the fractionation of plagioclase (~42.9%), clinopyroxene (~40.4%), titanomagnetite (~15.4%) and minor olivine (~1.3%). The somewhat high value of the sum of squared residuals calculated for the Santa Bárbara series results from the large compositional range considered (from basalts to benmoreites) and from the onset of crystallization of accessory phases (e.g. apatite) during the late stages of magma evolution toward benmoreitic compositions.

The validity of these models can be tested by calculating the variation in the concentration of trace elements (Ni, Sc and Sr) compatible with the fractionated mineral phases. Lines plotted in Fig. 8 (from A to C) were constructed considering the proportion of the minerals assessed from the mass balance calculations. These diagrams show a good fit between the model and the data. The same conclusion can be achieved from Fig. 8D, which emphasizes the role of the most abundant phenocrysts on major

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**Fig. 4.** Spiderdiagrams and REE profiles obtained for the Fissural and Santa Bárbara lavas considered in this work. Normalization values are from McDonough and Sun (1995). Lines with circles are for basalts, lozenges for hawaiites, squares for mugearites and triangles for benmoreites. The Serreta submarine sample is represented by a hatched line with white filled circles. Inset diagrams: typical patterns of lavas with strong HIMU and EM I signatures (from Weaver, 1991).

**Fig. 5.** Diagram (Zr/La vs. Rb/Th) showing the distinct elemental signatures of the Fissural and Santa Bárbara lavas. The Serreta submarine sample (white circle) erupted from the Fissural system is characterized by relatively high Zr/La and low Rb/Th ratios, more typical of the Santa Bárbara volcanic system. Data previously published for other Serreta lava balloons (Gaspar et al., 2003) are also presented.
element variation. The important role of clinopyroxene fractionation in both magmatic series is also expressed by the decrease of the CaO/Al2O3 ratio along with magma evolution (Fig. 7). Moreover, the negative correlation between the CaO/Al2O3 ratio and the alkalinity index [(Na2O+K2O) − 0.37⁎(SiO2−39); Albarède et al., 1997; not shown] demonstrates the influence of clinopyroxene fractionation on the increase of alkalinity of the Fissural basalts with differentiation. This is demonstrated by the (Na2O+K2O)/SiO2 ratio calculated for the pyroxene-dominated fractionating assemblage, which is significantly lower than that observed for the least evolved samples (0.014 versus 0.076, respectively). For the Santa Bárbara series, we note that the removal of Fe–Ti oxides can account for the compositional gap observed in the TAS diagram between mugearites and benmoreites (Fig. 3), given that their fractionation would trigger a sudden silica increase in the residual liquids (see also Mungall and Martin, 1995).

5.2. Magma contamination

The different incompatible trace element compositions of Fissural and Santa Bárbara magmas can be strengthened by comparing Santa Bárbara and Fissural lavas with similar MgO content (not shown; see Table 1). Santa Bárbara/Fissural normalized element ratios higher than unity were obtained for Th, U, Zr, Hf, and to a lesser extent K and Rb, showing their enrichment in Santa Bárbara lavas when compared to

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Fig. 6. Isotope systematics of the studied Terceira lavas. NHRL: Northern Hemisphere Reference Line (Hart, 1984). In the εNd–εHf isotope diagram the mantle array is the one defined by Andres et al. (2002). Fields for mantle components are based on data compiled by Stracke et al. (2003) for Atlantic MORB (DMM), Saint Helena (HIMU), Samoa and Society (EM II) and Walvis ridge and Pitcairn (EM I). The previously published Azores data is from the compilation made by Stracke et al. (2003), complemented with data from Elliott et al. (2007) and Beier et al. (2008).
those from the Fissural volcanic system. Conversely, calculated ratios for Ba and REE are lower than unity as the result of their relative depletion in Santa Bárbara magmas. These elements are not expected to be significantly fractionated by the mineral assemblages that were assumed to control the fractional crystallization process previously described. Thus, such differences can be related to source heterogeneity or, alternatively, to magma contamination. It is important to evaluate the latter possibility because most subaerial Fissural historical lavas were erupted in the vicinity of hyperalkaline trachytic lavas (pantellerites) belonging to the Pico Alto volcanic system (see Fig. 1B). However, trace element patterns of Fissural lavas reveal a lack of negative Ba, Sr and Ti anomalies that would be expected as the result of the assimilation of those pantelleritic lavas (Fig. 9). Also, the absence of positive Sr and Ti anomalies precludes the assimilation of cumulates complementary to the pantellerites. Moreover there is no significant correlation ($R^2 < 0.5$) between the Ba anomaly and an evolution index (MgO) as would be expected from an assimilation and crystal fractionation (ACF) process.

5.3. Melting constraints

At Terceira Island, primary magmas are only recognized for the Fissural volcanic system, which erupted lavas with MgO ~11 to 12 wt. % in equilibrium with liquidus olivine (Fo90) at Porto Martins and Porto Judeu areas (see Fig. 1B for location). These olivines display low-NiO contents (<0.35 wt.%), precluding the possibility of a pyroxenite-type source for their host lavas (Sobolev et al., 2005).

Experimental work on the melting of dry peridotites (e.g., Hirose and Kushiro, 1993, 1998) has shown that SiO$_2$ and MgO magma contents are independent of source composition, reflecting essentially the intensive parameters prevailing during magmatic extraction. Using the MgO and SiO$_2$ contents of the primary magmas and the equations proposed by Albarèrede (1992), we estimate that magmatic extraction occurred at temperatures of 1350 ± 40 °C and pressures around 1.7 ± 0.27 GPa.

The estimated temperature of magmatic segregation shows that Terceira primary magmas were hotter than the expected temperature of the asthenosphere (~1280 °C, e.g., McKenzie and Bickle, 1988), which indicates a local temperature excess that is compatible with a thermally driven mantle plume (see also Putirka, 2008).

The depths estimated for magma extraction (from ~50 to ~55 km, considering an average crustal thickness of 12 km, an oceanic crust density of 3.0 g/cm$^3$ and a mantle density of 3.3 g/cm$^3$) correspond to the spinel stability field, contrary to what would be anticipated from the REE fractionated patterns (Fig. 4; see also Fig. 10) or from the $^{230}$Th excess over $^{238}$U that were reported for Terceira lavas (Turner et al., 1997), both of which suggest melting in the presence of residual garnet. This contradiction has been described elsewhere and was explained by Eggins (1992) as the result of the onset of melting in the garnet stability zone (depths greater than 75 km) and later re-equilibration and segregation at lesser depths within the spinel stability zone (see also Francis, 1995). A similar explanation was also given by Beier et al. (2008) for Terceira and Graciosa lavas. Moreover, these authors stressed that the age of the crust at the base of Terceira is compatible with a lithosphere thickness of ~60 km, which would support the migration of the melting front of a mantle diapir up to the spinel stability zone. From the data distribution in the diagram of Fig. 10, a significant range of melting degrees of a garnet-bearing source is required to explain part of the trace element variability. This percept is
Fig. 8. A, B and C) Compatible trace element variation plotted against Th, taken as an index of magma evolution. Hatched lines represent evolutionary trends calculated using the Rayleigh’s law equation and the mineral proportions based on the least squares major element modeling (see text for details). Ni, Sc, Sr and Th partition coefficients are from Ewart and Griffin (1994), Lemarchand et al. (1987), Bindeman et al. (1998), McKenzie and O’Nions (1991) for plagioclase, from Duke (1976), Lemarchand et al. (1987), McKenzie and O’Nions (1991) for olivine and clinopyroxene, and from Nielsen (1992), Lemarchand et al. (1987), Ewart and Griffin (1994) for chromite and titanomagnetite. Vectors show the effect of instantaneous removal (10%) of olivine, clinopyroxene, plagioclase and magnetite. D) Variation diagram also showing Terceira lavas and phenocrysts composition. The dominant role of clinopyroxene + olivine for the Fissural and of clinopyroxene + plagioclase fractionation for the Santa Bárbara magmatic series is clear from the comparison of the liquid lines of descent and the composition of the main phenocrysts phases.

Fig. 9. Spider diagram comparing the Fissural lavas (that occur in the central Terceira Island) with the Pico Alto trachytes. Normalization values are from McDonough and Sun (1995). Fissural lavas are plotted with the same symbols as in Fig. 4.

Fig. 10. Primitive mantle normalized Ce/Yb ratios plotted against (Ce)N. Melting trends for garnet (Gt) and spinel (Sp) lherzolites were determined assuming an enriched mantle source with 4 ppm and 0.5 ppm of Ce and Yb, respectively. The modeling was performed considering the batch melting of a source with 60% ol + 25% opx + 10% cpx + 5% gt and 60% ol + 25% opx + 10% cpx + 5% sp. The Ce and Yb partition coefficients were taken from McKenzie and O’Nions (1991). Note that the least evolved samples (lower Ce contents) plot close to the melting trend of the garnet lherzolite. White circle: Serreta sample. The dotted lines represent two sub-parallel trends formed by the Santa Bárbara samples, emphasizing the role of variable degrees of partial melting to the variability of the Terceira lavas.
strengthened by the two sub-parallel trends formed by lavas from the same volcanic system (Santa Bárbara).

The higher degree of partial melting interpreted for the relatively primitive Serreta lava is interpreted as reflecting the presence of a thinner lithosphere according to the age decrease of the oceanic crust toward the MAR.

5.4. Source implications

As mentioned above, the significant differences in incompatible trace element ratios and isotopic signatures emerging from the comparison between the two studied volcanic systems (Fissural and Santa Bárbara) cannot be explained by fractional crystallization, assimilation and/or distinct degree of partial melting. From this perspective, these differences are believed to reflect the heterogeneity of the mantle domain(s) sampled by Terceira magmas.

In their study on Terceira Rift magmatism, Beier et al. (2008) showed the existence of significant heterogeneity at the scale of that structure, whereas smaller scale heterogeneity is evidenced by the existence of distinct intra-island magmatic signatures (e.g., Dupré et al., 1982; Haase and Beier, 2003; Millet et al., 2009). This study, which confirms the existence of mantle variability at the scale of a small island (398 km²), evidences that such heterogeneities are even detectable when two "coeval" volcanic systems erupted less than 10 km away.

The linear arrays observed in the $^{207}\text{Pb}/^{204}\text{Pb}$ versus $^{206}\text{Pb}/^{204}\text{Pb}$ space can be interpreted as secondary isochrones or, considering that the reference isotope is the same in both ratios, as binary mixing lines. Our data for the Fissural volcanic system define an array (see inset of Fig. 6) with a slope of 0.1621 ($R^2 = 0.965$), which, if interpreted as an isochron, corresponds to an age of circa 2.47 Ga. Accordingly, this would be the age of the U/Pb increase responsible for the positioning of the Terceira samples well to the right of the Geochron (e.g., Chase, 1981).

A linear array is also observed for the same magmatic series in the $^{208}\text{Pb}/^{204}\text{Pb}$ versus $^{206}\text{Pb}/^{204}\text{Pb}$ diagram, which can be used to decipher the meaning of both Pb linear arrays (e.g., Abouchami et al., 2000). If these arrays have an age meaning, a broad positive correlation should exist between the time-integrated Th/U ratios inferred from Pb isotopic ratios ($K_{\text{Th}}$, Galer and O’Nions, 1985) and the elemental Th/U measured in the samples. However, there is not a correlation between measured Th/U ratio and $K_{\text{Th}}$ (not shown) leading to the interpretation that the observed linear arrays represent the binary mixing of mantle components with different time-integrated compositions.

Santa Bárbara lavas plot close to the Terceira end-member defined by Beier et al. (2008) and clearly to the right of the Geochron, presenting $^{206}\text{Pb}/^{204}\text{Pb}$ ratios up to 19.992 (Fig. 11; Table 2). This implies that their sources were affected sometime in the past by a significant U/Pb increase. They also depict trace element patterns (Fig. 4) similar to those characterizing typical HIMU-type basalts (Weaver, 1991). However, the $^{208}\text{Pb}/^{204}\text{Pb}$ ratios are significantly lower than those attributed to the more extreme HIMU signatures (e.g., Mangaia: 21.864; Woodhead, 1996). This implies the mixing of a high-$\mu$ end-member with a component characterized by significantly less radiogenic lead isotope signatures.

Millet et al. (2009) showed that the Azores lavas form two distinct trends on the $\Delta 8/4$ versus $^{206}\text{Pb}/^{204}\text{Pb}$ diagram, converging in the North Azores Fracture Zone MORB field (Fig. 12). This suggests that the unradiogenic-Pb component required by the isotopic signature of Santa Bárbara lavas is related to the mantle domain feeding the Mid-Atlantic Ridge. This is also endorsed by studies involving He and Ne, which argue for mixing between a low $^{21}\text{Ne}/^{22}\text{Ne}$ and $^{4}\text{He}/^{3}\text{He}$ component that is required for Terceira Island and a radiogenic He and nucleogenic Ne end-member characterizing the unradiogenic-Pb component (Madureira et al., 2005; Moreira et al., 1999).

For Santa Bárbara, the proposed mixing between a HIMU-type end-member and a MORB-type one is supported by the $^{143}\text{Nd}/^{144}\text{Nd}$ versus $^{206}\text{Pb}/^{204}\text{Pb}$ diagram (Fig. 11). Conversely, Fissural lavas form a trend toward lower $^{143}\text{Nd}/^{144}\text{Nd}$, $^{206}\text{Pb}/^{204}\text{Pb}$ and also $^{176}\text{Hf}/^{177}\text{Hf}$, $^{207}\text{Pb}/^{204}\text{Pb}$ and $^{208}\text{Pb}/^{204}\text{Pb}$ ratios (see Fig. 6; Table 2). These require the involvement of an end-member, characterized by a relatively lower time-integrated Sm/Nd, Lu/Hf, U/Pb and Th/Pb ratios as compared with the Terceira end-member sampled by Santa Bárbara lavas. Such more unradiogenic-Hf and -Nd end-member is, also, clearly distinct from the enriched end-member described at the eastern extremity of the Terceira Rift at S. Miguel Island (Fig. 11), for which several distinct origins have been proposed (cf. Beier et al., 2007; Elliott et al., 2007; Widom et al., 1997). Their isotopic signatures are more akin to those depicted by the end-member described by Beier et al. (2008) for D. João de Castro Bank (Fig. 11; Table 2), whose presence was not detected in Terceira Island by these authors.

The affinities of Fissural lavas with the D. João de Castro Bank end-member are reinforced by the relatively high Ba/Nb ratios (~10).
characterizing lavas from the two volcanic systems (Table 1; Beier et al., 2008). The high Ba/Nb ratios reflect either the mantle source composition or, alternatively, Ba–Nb fractionation during partial melting processes. These elements can only be fractionated in the presence of residual phlogopite, ilmenite or rutile (Bodinier et al., 1996; LaTourrette et al., 1995; Ryerson and Watson, 1987), contributing to decrease (phlogopite) or increase (ilmenite or rutile) of the Ba/Nb ratio. However, residual ilmenite or rutile should be reflected by a negative Ti anomaly that is not observed in Fissural lavas. These also display (Ce/Nb) sub-continental lithospheric mantle (SCLM) is considered to be highly heterogeneous processes generating amphibole-rich domains (Lundstrom et al., 2003), which given amphibole/melt partition coefficients generating amphibole-rich domains (Lundstrom et al., 2003), which given amphibole/melt partition coefficients would also be characterized by low U/Pb ratios (e.g., LaTourrette et al., 1995).

Owing to their long-lived isolation from the convective asthenospheric mantle, SCLM is considered to be highly heterogeneous presenting compositions reflecting the age and tectonothermal events of the overlying crust (e.g., Griffin et al., 1999). However, there is growing evidence for the positioning of the SCLM below the HFNd mantle array (e.g., Geldmacher et al., 2008; Griffin et al., 2000; Tappe et al., 2007) conferring plausibility to the hypothesis of a small contribution of SCLM to the Fissural magmas, as has been proposed based on other grounds, for the Azores (Millet et al., 2009; Moreira et al., 1999; O’Reilly et al., 2000; Widom and Shirey, 1996).

From the above discussion, we can conclude that the isotopic variability of the studied Terceira lavas reflects the mixing of several end-members. Santa Bárbara lavas reveal the mixing of an HIMU-like end-member that is believed to reflect the contribution of the mantle plume to the Terceira basalts, with the upper mantle domain feeding the Mid-Atlantic Ridge. Fissural lavas require an additional component with some characteristics that are similar to the SCLM.

HIMU signatures have been considered to stem from recycling and processing at the subduction zones of altered oceanic crust, resulting in the addition to the deep mantle of material characterized by low Rb/Sr and high U/Pb ratios, which evolve over time to high $^{206}\text{Pb}/^{204}\text{Pb}$ and low $^{87}\text{Sr}/^{86}\text{Sr}$ ratios (e.g., Chauvel et al., 1992; Hofmann and White, 1982; Stracke et al., 2003; Weaver, 1991). These oceanic materials are believed to penetrate the Earth mantle as part of the downgoing slabs, which, as shown either by tomographic (e.g., Maruyama et al., 2007) or by density constraints (e.g., Hirose et al., 2005), can reach the lower levels of the mantle. The most Pb-radiogenic Santa Bárbara sample ($^{206}\text{Pb}/^{204}\text{Pb}=19.992$) presents a $^{87}\text{Sr}/^{86}\text{Sr}$ (0.70357) ratio slightly higher than the samples from HIMU type-localities, where this ratio is usually below 0.703 (e.g., Jackson and Dasgupta, 2008). For the Azores, this was also reported by Millet et al. (2009) in his study of S. Jorge Island suggesting that such relatively high $^{87}\text{Sr}/^{86}\text{Sr}$ ratios are characteristic of the HIMU-type end-member at the scale of the Azores archipelago.

At least a fourth contributor is required to explain the He and Ne isotopic characteristics described by Moreira et al. (1999) and Madureira et al. (2005) for Terceira Lavas. Indeed, the analyzed noble gases trapped in melt inclusions occurring in olivine phenocrysts (R/Ra up to 11.4; $^{21}\text{Ne}/^{22}\text{Ne}_{\text{corr}}=0.052$) are clearly distinct from those assigned to mantle components similar to the dominant end-members in the Terceira sources (DMM as imaged by MORB; R/Ra = 8 ± 1; Zindler and Hart, 1986; $^{21}\text{Ne}/^{22}\text{Ne}_{\text{corr}}=0.075$, Sarda et al., 2000; HIMU: R/Ra −5 to 7, Hanyu and Kaneoka, 1997; Moreira and Kurz, 2001; Parai et al., 2009; $^{21}\text{Ne}/^{22}\text{Ne}_{\text{corr}}=0.050$ to 0.066, Parai et al., 2009). Those Terceira signatures, requiring the long term isolation of a low (U + Th)/He and (U + Th)/$^{206}\text{Ne}$ reservoir (Class and Goldstein, 2005; Jackson et al., 2010; Porcelli and Elliott, 2008) have been considered to reflect the incorporation of lower mantle material in the Azores mantle domain (Madureira et al., 2005; Moreira et al., 1999). Therefore, there is a contradiction resulting from the apparent decoupling between the lithophile and Pb isotopic data, pointing to the existence of recycled materials, and He and Ne isotopic signatures, showing evidence for the contribution of lower mantle material. This has been explained elsewhere by assuming that the noble gas signatures of the deep mantle reservoir are virtually “unaffectted” by the addition of crustal materials, which were almost completely stripped of He and Ne by degassing at mid-ocean ridges and, later, at subduction zones. In opposition, their lithophile, Os and Pb isotope signatures are strongly constrained by the recycled crustal material (e.g., Brandon et al., 2007; Jackson et al., 2009; Mata et al., 2010).

The heterogeneity of mantle plumes is thought to reflect either the role of entrainment of the surrounding mantle (e.g., Hauri et al., 1994) or the sampling of a heterogeneous thermal boundary layer (TBL) from which the plume rises (e.g., Farnetani et al., 2002). As shown above, the Azores mantle plume incorporates at least ancient recycled oceanic crust and a lower mantle contribution. Admitting that the extension associated with the Terceira Rift can only be regarded as a mechanism that facilitates the melting and/or magma extraction from the plume head imaged by seismic tomography (Yang et al., 2006) regional-scale studies can be used to infer information about the plume structure. Indeed, numerical simulations have shown that the spatial distribution of heterogeneities sampled by a volcanic province can provide insights on the spatial fields of the plume and the TBL structures. A recent study of the Azores, also including the Flores and Corvo Islands located to the west of the MAR, suggests the existence of a chemical axisymmetric pattern, which, if reflecting a plume concentric zoning (Beier et al., 2010), would imply a layered TBL (see Farnetani and Hofmann, 2009, 2010).

Tomographic studies indicate that the Azores plume is centered NE of Terceira Island (Yang et al., 2006). However, a comparison of our results with those obtained by Millet et al. (2009) shows that S. Jorge
lavas are significantly more Pb-radiogenic ($^{206}\text{Pb}/^{204}\text{Pb}$ up to ~20.5) than those from Terceira ($^{206}\text{Pb}/^{204}\text{Pb}$ up to 19.992). This suggests that S. Jorge lavas carry to the surface a stronger plume signal than those at Terceira, where the HIMU signatures are more diluted by a stronger contribution of the local depleted upper mantle (see Fig. 11). We interpret this to reflect a larger degree of melting at Terceira as a result of the extensional regime characterizing the Terceira Rift.

6. Concluding remarks

Through the study of the volcanic systems (Santa Bárbara and Fissural) erupting mafic lavas at Terceira Island (Azores), we demonstrated the existence of present-day (less than 100 ka) small-scale lateral mantle source heterogeneity (Fig. 14). The western Terceira lavas, represented by the Santa Bárbara volcanic system and the Serreta ridge, sample the Terceira end-member defined by Beier et al. (2008) for the Terceira Rift. This end-member is here interpreted as the result of mixing between plume-derived ancient recycled oceanic crust (HIMU-type component) and the regional depleted MORB magmas/source. The D. João de Castro Bank end-member (Beier et al., 2008), not previously reported for Terceira Island, contributes to the Fissural volcanic system. It is suggested that the chemistry of these lavas reflects the interaction of the Azores plume with SCLM, the latter probably residing in the upper mantle as delaminated megaliths. Published noble gas data (Madureira et al., 2005; Moreira et al., 1999) on lavas from both volcanic systems also argue for the presence of a relatively undegassed component, which is interpreted to be inherited from the lower mantle reservoir sampled by the Azores plume.

The intra-volcanic system’s chemical variability is mainly explained by variable proportions of the above-mentioned local end-members and by crystal fractionation processes. Magmatic evolution processes were more important for the Santa Bárbara volcanic system where magmas reach comenditic compositions. Fissural lavas range from basalts to hawaiites, with the latter occurring preferentially in the central part of the island where the crust is presumably thicker. As indicated by trace elements the partial mantle melting began in the garnet stability zone, although major element compositions suggest that magma segregation occurred within the spinel zone. A higher degree of partial melting was invoked for the Serreta submarine lava, which is interpreted to reflect the presence of a thinner lithosphere toward the MAR (Fig. 14).

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Fig. 14. Cartoon illustrating the mantle heterogeneity beneath Terceira Island sampled by Fissural and Santa Bárbara mantle sources. The white dashed line represents depths of magma extraction, which is believed to be constrained by the thickness of the lithosphere.


