

Temporal evolution of a post-caldera, mildly peralkaline magmatic system: Furnas volcano, São Miguel, Azores

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Abstract Furnas is one of three active central volcanoes on São Miguel Island, Azores, and is considered to be one of the most hazardous in the archipelago. In this study, the pre-eruptive magma plumbing system of the 10 young (<5 ka), intra-caldera, sub-Plinian, trachytic eruptions of the Upper Furnas Group (UFG) is investigated via whole rock major and trace element geochemistry, mineral chemistry, thermobarometry, and petrogenetic modelling. The main aim of this work is to elucidate the petrogenesis of the Furnas trachytes, constrain the P–T–fO₂ conditions under which they evolve, and investigate the temporal evolution of the magma plumbing system. Results indicate that the trachytes are derived predominantly from extended fractional crystallisation of alkali basalt parental magmas, at depths between ~3 and 4 km.

This is considered to take place in a density-stratified reservoir, with alkali basalt magmas at the base and hydrous trachytes forming an upper cap or cupola. The presence of this reservoir at shallow crustal depths beneath the caldera likely inhibits the ascent and subsequent eruption of mafic magmas, generating a compositional Daly Gap. Rare syenitic ejecta represent in situ crystallisation of trachytic magmas in the thermal boundary zone at the top of the reservoir. Trachytic enclaves within these syenites, in addition to banded pumices and ubiquitous clinopyroxene antecrysts in the UFG pumice falls, provide evidence for mingling/mixing processes within the magmatic system. Despite relatively uniform major element compositions, systematic trace element variations within individual eruptions highlight the importance of fractional crystallisation during late-stage evolution of the trachytes. This is facilitated by the accumulation of water and the development of mild peralkalinity, which contribute to low pre-eruptive melt viscosities and efficient crystal settling. Compositional zoning patterns between individual eruptions cannot be accounted for by periodic tapping of a single magma batch undergoing fractional crystallisation. Instead, up to four individual cycles are recognised, in which a zoned cap of eruptible trachytic magma, formed at the top of the reservoir, was erupted in one or more eruptions and was re-established via intermittent replenishment and subsequent fractional crystallisation.

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Introduction

Whilst peralkaline volcanic rocks are volumetrically less significant than their subalkaline counterparts, the petrogenetic processes that lead to their formation may be more complex (e.g. Mahood 1984; Mungall and Martin 1995; Scaillet and Macdonald 2001; White et al. 2009; Markl et al. 2010). Recent studies of peralkaline silicic magmatic systems have led to the general consensus that mafic magmas either act as a parental magma, linked to the peralkaline silicic magmas primarily by fractional crystallisation (e.g. Barberi et al. 1975; Self and Gunn 1976; Civetta et al. 1998; Peccerillo et al. 2007; Macdonald 2012), or alternatively act as a heat source generating crustal-derived partial melts which may then differentiate via fractional crystallisation (e.g. Bohron and Reid 1997; Trua et al. 1999; Avanzinelli et al. 2004). Both of these models can be affected by varying degrees of crustal assimilation and magma mixing (Macdonald et al. 2015).

Here, we investigate the petrogenetic processes that operate in the basalt-trachyte suite of Furnas volcano, São Miguel, Azores (Fig. 1). Furnas volcano is an example of an oceanic island volcanic centre that exhibits many of the petrogenetic problems associated with peralkaline magmatic systems, including a volumetric abundance of felsic products with a peralkaline affinity, an apparent Daly Gap, and a spatial concentration of felsic products within the volcano's caldera complex, with mafic products on the flanks (Booth et al. 1978; Moore 1991). Guest et al. (1999) suggested that, similar to neighbouring volcanoes Sete Cidades and Fogo (Água de Pau), the felsic magmas of Furnas were largely generated via fractional crystallisation (cf. Storey et al. 1989; Moore 1991; Widom et al. 1992) and that the observed geographical distribution of mafic and felsic eruption products could result from the presence of established trachytic magma reservoirs that intercept ascending mafic magmas (cf. Wolff 1987). However, the evolution of peralkaline trachyte has not been quantitatively demonstrated at Furnas, and constraints upon the plumbing system are currently limited to geophysical studies (e.g. Machado 1972; Camacho et al. 1997; Montesinos et al. 1999), which broadly point towards a magma reservoir at shallow depths in the crust (~5 km).

In this contribution, detailed petrography, mineral chemistry, thermobarometry, whole rock major and trace element geochemistry, as well as petrogenetic modelling, are applied to nine of the ten recent (<5 ka) post-caldera eruptions that together constitute the Upper Furnas Group (UFG). With this data set, this study aims to: (1) investigate the petrogenesis and differentiation of mildly peralkaline silicic magmas at Furnas volcano and the generation of a Daly Gap, (2) place constraints upon the P–T–fO₂ conditions of the magma

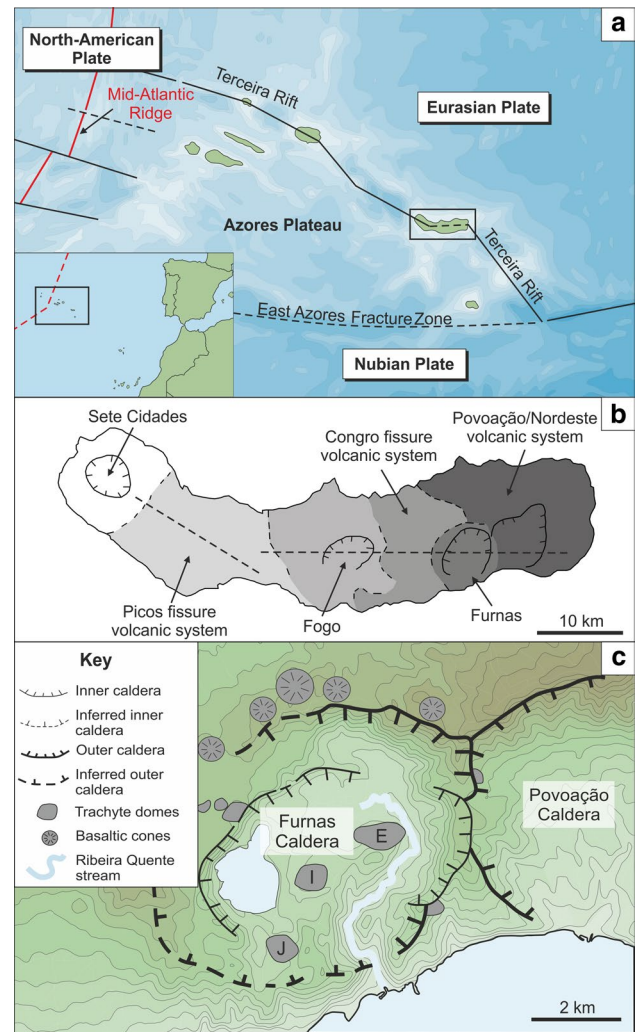


Fig. 1 Location maps of Furnas volcano, **a** summary map of the major tectonic features associated with the Azores triple junction, constructed after Searle (1980), Freire Luis et al. (1994), and Vogt and Jung (2004), **b** schematic map of São Miguel, highlighting the three active central volcanoes and fissure volcanic systems (adapted from Moore 1990), **c** summary map of the caldera complex of Furnas volcano and the neighbouring Povoação caldera. The lava domes associated with the Furnas E, I, and J eruptions are labelled. Modified after Guest et al. (1999) and Cole et al. (1999)

plumbing system beneath Furnas, and (3) evaluate the compositional evolution of a periodically tapped, metaluminous to mildly peralkaline magma plumbing system over a period of ~5000 years (Guest et al. 1999). A greater understanding of the pre-eruptive plumbing system of Furnas has implications for the processes that generate mildly peralkaline silicic magmas in such settings (e.g. Macdonald 2012), and the re-establishment and temporal evolution of post-caldera magmatic systems (e.g. Bachmann et al. 2012), as well as for the ongoing risk management of this highly active and hazardous volcano (e.g. Jones et al. 1999; Chester et al. 2002).

Geological background

The Azores archipelago is located in the central North Atlantic Ocean (lat. 37°N to 40°N, long. 25°W to 32°W), 1300 km west of the Portuguese mainland. The islands are situated in a unique geodynamic setting as a result of the triple junction where the North American, Eurasian, and Nubian plates meet (Fig. 1a). The nine islands themselves are the emergent portions of large volcanic centres that rise from the Azores Plateau, a bathymetric and gravitational anomaly associated with a morphologically complex area ($\sim 5.8 \times 10^6$ km²) of thickened oceanic crust that formed between 20 and 7 Ma (Kaula 1970; Ridley et al. 1974; Gente et al. 2003). The islands are divided into three groups based upon their geographical position. The central (Pico, Faial, São Jorge, Graciosa, Terceira) and eastern (Santa Maria and São Miguel) groups lie within a broadly triangular portion of the Azores Plateau, defined by three major tectonic features; the Mid-Atlantic Ridge in the west, the East Azores Fracture Zone (EAFZ) to the south, and the Terceira Rift to the north-east (Krause and Watkins 1970; Ridley et al. 1974; Vogt and Jung 2004; Luis and Miranda 2008). In contrast, the western group (Flores and Corvo) lie outside of this feature, on the western side of the Mid-Atlantic Ridge.

The island of São Miguel belongs to the eastern group and is the largest of the Azorean islands, comprising three active Quaternary trachytic central volcanoes [from west to east: Sete Cidades, Fogo (or Água de Pau), and Furnas] (Fig. 1b). Sete Cidades and Fogo are separated by the Picos basaltic fissure system (the Waist zone of Booth et al. 1978), whilst Furnas is separated from Fogo by the Congro basaltic fissure system (the Achada das Furnas complex of Booth et al. 1978). The easternmost part of the island is composed by the Pliocene–Pleistocene Povoação/Nordeste volcanic system, which is considered extinct (Fig. 1b). Unlike Sete Cidades and Fogo, Furnas lacks a distinctive constructive morphology and is instead made up of an approximately 7 × 6 km nested caldera complex (Fig. 1c) that is considered to have been formed by at least two caldera-forming eruptions dated at $\sim 30,000$ ¹⁴C y B.P. and $\sim 12,000$ ¹⁴C y B.P. (Duncan et al. 1999; Guest et al. 1999; cf. Sibrant et al. 2015).

Furnas eruptive history

Furnas is considered to be one of the most active and hazardous volcanoes in the entire Azores archipelago, having erupted at least ten times in the last 5000 years. During its subaerial existence, Furnas has exhibited effusive (dome-forming) to highly explosive (caldera-forming) eruptions of

felsic magmas (Cole et al. 1999; Guest et al. 1999, 2015). Approximately 1500 people inhabit Furnas caldera, with several hundreds more in the immediate vicinity. A stratigraphical study by Guest et al. (1999) revealed that more than 90 % of eruption products relating to Furnas are of pyroclastic nature, suggesting a volcanic history in which explosive activity is prevalent, certainly more so than the neighbouring Povoação/Nordeste volcanic system. Volcanism within the caldera complex has been exclusively trachytic in composition, with mafic products being limited to vents on the volcano's flanks. Although Guest et al. (1999) recognised a compositionally continuous sequence from basalt to trachyte, intermediate compositions were found to be subordinate in volume, indicating the presence of a compositional Daly Gap. The same study divided the stratigraphy of Furnas into three groups: the Upper (UFG), Middle (MFG), and Lower (LFG) Furnas groups. The LFG and MFG comprise numerous trachytic pyroclastic units, with minor trachytic domes and basaltic cones (Moore 1990). The boundary between the two groups is marked by the Povoação Ignimbrite Formation ($\sim 30,000$ ¹⁴C y B.P.), a voluminous, partially welded, metaluminous to mildly peralkaline ignimbrite that fills much of the neighbouring Povoação caldera and is believed to represent the first caldera-forming eruption of Furnas volcano (Schmincke and Weibel 1972; Booth et al. 1978; Duncan et al. 1999).

The UFG is made up of at least ten intra-caldera, sub-Plinian eruptions of trachytic pumice, named Furnas A to J (the latter is also known as Furnas AD 1630; Cole et al. 1995; Guest et al. 2015) in order of decreasing age (Booth et al. 1978) (Online Resource 1). Of these eruptions, the latter two are known to have been historic, occurring after settlement on the island. Three of these eruptions are known to have also produced lava domes in their final stages (Furnas E, I, and J) (Cole et al. 1999). The dominant lithologies of the UFG are inter-bedded pumice lapilli and ash beds that are inferred to be the result of complex transitions between magmatic and phreatomagmatic activity during eruption, though Furnas H appears to have comprised exclusively magmatic activity (see stratigraphic logs in Online Resource 2) (Cole et al. 1995; Guest et al. 1999). The base of the UFG is marked by a deposit from nearby Fogo volcano, named Fogo A (Walker and Croasdale 1971). Radiocarbon ages for Fogo A range from 4480 to 5380 ¹⁴C y B.P. (e.g. Moore and Rubin 1991; Wallenstein 1999; Pensa et al. 2015), but its age is generally reported as ~ 5 ka. Although the deposit is not genetically related to Furnas, it is well known and easily identified and, in the absence of well-exposed deposits relating to the second caldera-forming event at Furnas ($\sim 12,000$ ¹⁴C y B.P.), was chosen as the reference stratigraphic marker of the island that delimits the base of the UFG (Guest et al. 1999). Other

deposits from Fogo are also commonly included within the stratigraphical sequence of the UFG to act as chronological reference points (e.g. Cole et al. 1999).

Petrography

Pumice falls

The fall deposits of the UFG are dominated by highly vesicular (~85 %), aphyric to crystal poor (<5 % phenocryst content) pumice clasts. The dominant mineralogy shows little variation throughout the deposits, comprising alkali feldspar (~67 vol%), Fe–Ti oxides (~20 vol%), biotite (~10 vol%), clinopyroxene (~3 vol%), and apatite (accessory). Rarely, alkali feldspar microlites with high aspect ratios are observed. Additionally, a small number of pumice clasts from layer Lf of Furnas J (see Cole et al. 1995) exhibit pronounced banding, notably higher crystal contents, more abundant groundmass microlites, and rare inclusions of dense, crystalline material of similar mineralogy to the pumice, which range in size from ~0.5 to 6 mm.

Alkali feldspar is the most abundant phase and exists as phenocrysts (up to ~2 mm), microphenocrystic fragments, and, more rarely, groundmass microlites. Phenocrysts are commonly tabular and euhedral (Fig. 2a), but rare examples are rounded, with pronounced embayment structures and resorbed cores (Fig. 2b). Small crystal fragments with irregular, fractured forms are also present, which may be heavily resorbed. Such examples are particularly common in the banded pumices of Furnas J, described above. Where groundmass microlites are present, they are almost exclusively acicular and do not exceed 100 μm in length.

Clinopyroxene exists as microphenocrysts, typically <0.5 mm in length (Fig. 2c). Inclusions of acicular apatite are frequently observed, and Fe–Ti oxide inclusions are ubiquitous. More rarely, small (<50 μm) clinopyroxene crystals are seen with irregular crystal forms, either heavily fractured, or rounded and resorbed. Such examples are more abundant in the banded pumices of Furnas J.

Biotite occurs as small (<0.5 mm) microphenocrysts (Fig. 2d). Crystal forms are dominantly euhedral to subhedral and may show evidence for resorption. Fe–Ti oxides are generally very small (<0.5 mm), subhedral, and frequently associated with clinopyroxenes. More rarely, they may be found as inclusions within feldspars. They are ubiquitously associated with small, acicular apatite crystals, which may be partially or fully included within the Fe–Ti oxide host.

Lava domes

Samples collected from the two lava domes associated with Furnas I and J reveal mineral assemblages that are broadly

similar to those of the UFG pumice falls, including phenocrysts and microphenocrysts of alkali feldspar, clinopyroxene, Fe–Ti oxides, biotite, and apatite, in approximate order of decreasing abundance. However, due to several key mineralogical and textural differences, the two domes are treated separately.

The lava dome associated with Furnas J is dominated by alkali feldspar, which can be found as phenocrysts up to ~5 mm in size, and also as relatively coarse, acicular microlites in the groundmass (up to ~0.4 mm). Phenocrystic feldspars range from euhedral to subhedral, with rounded edges and uncommon embayments providing evidence for partial resorption. Feldspar microlites are frequently aligned, giving the groundmass a well-defined trachytic texture, which is also taken up by proximal phenocrysts. Rarely, feldspars are observed as glomerocrysts that reach sizes of up to ~2 mm and comprise a number of partially resorbed feldspar phenocrysts that meet at irregular and often lobate boundaries (Fig. 2e).

Clinopyroxene is found throughout the dome as phenocrysts (up to ~1 mm) and groundmass microlites (up to ~0.1 mm). As observed in the UFG pumice falls, it is characteristically green in thin section and ubiquitously associated with smaller Fe–Ti oxide crystals which are partially or totally included within the larger clinopyroxene crystal.

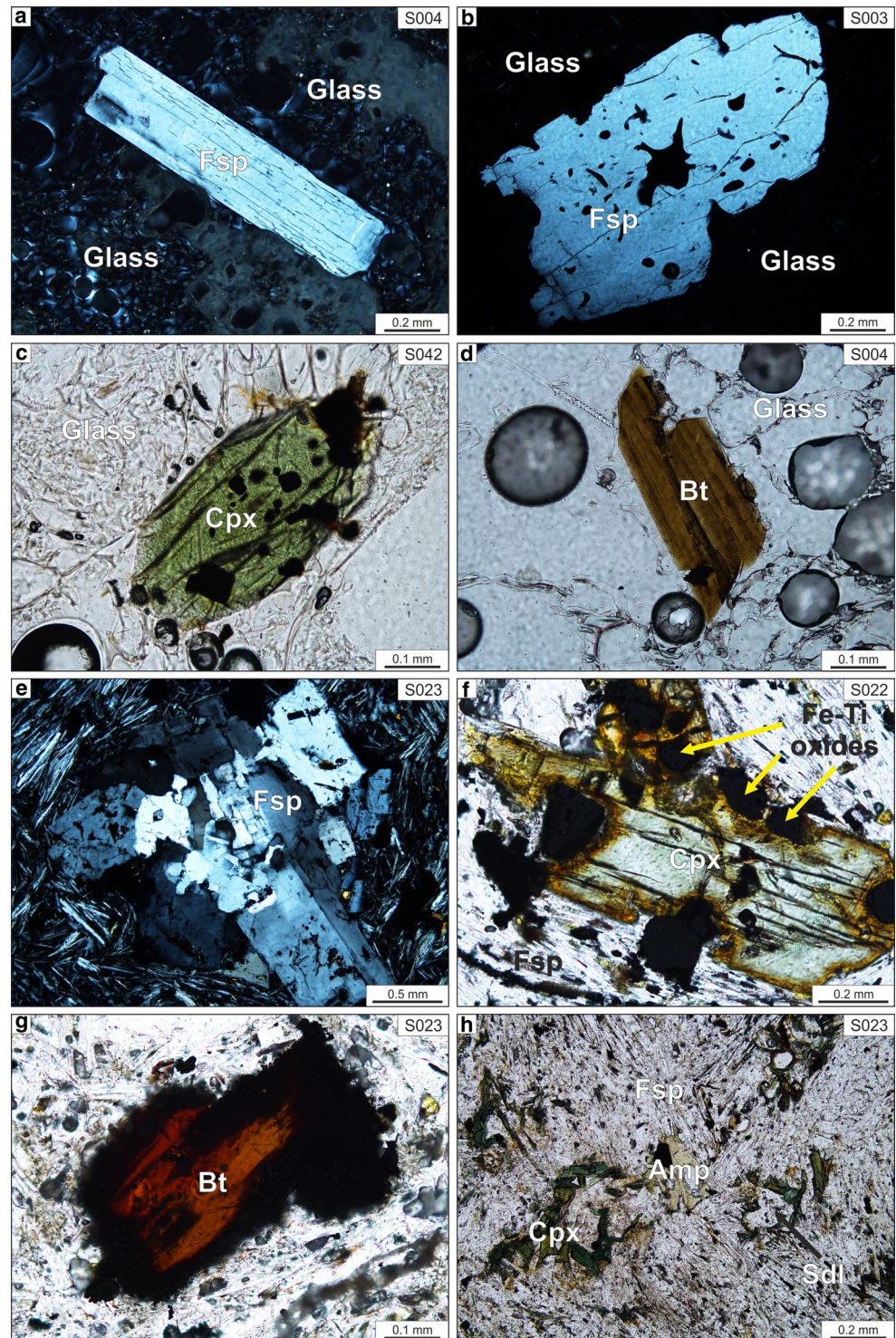
Phenocrystic and microphenocrystic clinopyroxenes display brown alteration rims that also extend along any fractures that cut through the crystal (Fig. 2f). In addition to Fe–Ti oxides, small apatite needles are also frequently observed as inclusions.

Fe–Ti oxides are found as uncommon microphenocrysts (generally between 0.25 and 0.5 mm in size) and abundant groundmass crystals (<50 μm). Microphenocrysts frequently exhibit exsolution lamellae of ilmenite. Such features are rare in the groundmass oxides. Like the UFG pumice falls, inclusions and partial inclusions of apatite are abundant.

Biotite is uncommon and is generally found as microphenocrysts, <0.2 mm in length. Examples are frequently well formed and often aligned with the trachytic texture in the groundmass. Rare examples of larger, phenocrystic biotite, up to 0.75 mm in length, are present in some samples. Unlike microphenocrystic biotite, these examples are ragged and exhibit prominent reaction rims (Fig. 2g), which contain abundant small Fe–Ti oxides, and also run along some fractures and cleavage planes.

In contrast to the UFG pumice falls, the Furnas J dome also contains sodalite, which can be found as well-formed, equant microphenocrysts that are typically 0.2 mm in size. Inclusions of feldspar and clinopyroxene are extremely common, and feldspar microlites frequently impinge upon the sodalite crystal faces. In some cases, sodalite appears rounded and partially embayed, indicating resorption.

Fig. 2 Representative photomicrographs for the UFG, **a** euhedral alkali feldspar phenocryst in vesicular glass (xpl), **b** highly resorbed and embayed alkali feldspar phenocryst (xpl), **c** clinopyroxene phenocryst with numerous Fe–Ti oxide and apatite inclusions (ppl), **d** subhedral biotite phenocryst in highly vesicular glass (ppl), **e** alkali feldspar glomerocryst with irregular internal boundaries, surrounded by a medium-grained, trachytic groundmass (Furnas I lava dome) (xpl), **f** clinopyroxene phenocryst with distinctive oxidation rim and numerous Fe–Ti oxide inclusions (Furnas J lava dome) (ppl), **g** large, reacted biotite phenocryst remnant (Furnas I lava dome) (ppl), **h** Na–Ca amphibole, aegirine-augite, and sodalite groundmass crystals, surrounded by alkali feldspar laths in the Furnas I lava dome (ppl). *Fsp* alkali feldspar, *Cpx* clinopyroxene, *Bt* biotite, *Amp* amphibole, *Sdl* sodalite. Sample numbers given in upper right of each image



The Furnas I lava dome rock is similar to that from the Furnas J dome, containing alkali feldspar phenocrysts surrounded by a coarse, alkali feldspar-based groundmass, as well as uncommon alkali feldspar glomerocrysts that show irregular and lobate interior grain boundaries. Clinopyroxene is found as anhedral phenocrysts (up to ~1 mm) in length and as small microlites in the groundmass. Fe–Ti

oxides are present as frequently exsolved microphenocrysts that are associated with the clinopyroxene phenocrysts, and as groundmass crystals. Biotite is observed in the Furnas I dome, as small subhedral microphenocrysts (<0.2 mm), or rare, ragged phenocrysts with an Fe–Ti oxide-rich reaction rim. Sodalite microphenocrysts are generally small (<0.2 mm) and euhedral, with abundant inclusions of

feldspar and clinopyroxene (Fig. 2h). The Furnas I dome also contains amphibole, found only as anhedral, interstitial crystals in the groundmass (Fig. 2h).

Syenite nodules

A small number of coarse-to-medium-grained syenitic clasts ejected as cognate xenoliths were sampled from the pyroclastic surge units associated with the Furnas J eruption. Individual specimens rarely exceed ~10 cm in size, and, in some instances, contain enclaves of darker, finer-grained rock. Boundaries between enclave material and host syenite range from sharp and well defined to irregular and diffuse, and, in some samples, both boundary types can be observed. Syenite mineral assemblages are dominated by cumulus alkali feldspar, which reaches sizes of up to 5 mm, and frequently displays perthitic textures. Intercumulus phases include clinopyroxene, amphibole, Ti-magnetite, ilmenite, biotite, pyrrhotite, and rare zircon (Online Resource 3). Mirolitic cavities are common in both the syenites and enclaves. Unlike reported syenitic ejecta from Terceira (Jeffery et al. 2016), Ascension (e.g. Harris 1983), and nearby Fogo volcano (Ridolfi et al. 2003), the Furnas syenites do not contain any of the Na–K–Ca–Ti–Zr silicate minerals that characterise agpaitic phase assemblages (e.g. aenigmatite, astrophyllite, eudialyte, cf. Marks et al. 2011). Instead, Ti-magnetite, ilmenite, and trace amounts of zircon indicate that the syenites are miaskitic.

Methods

Analytical techniques

Whole rock analyses

Whole rock major and trace element analyses for the UFG were undertaken at The Open University, UK, using an Applied Research Laboratories 8420+ dual goniometer wavelength-dispersive X-ray fluorescence (XRF) spectrometer. Additional analyses were produced using a Bruker AXS S4 Pioneer XRF spectrometer at the University of East Anglia, UK. The trace element concentrations of a small subset of samples were determined via inductively coupled plasma mass spectrometry (ICP-MS), using an Agilent 7500 s quadrupole mass spectrometer at The Open University, UK. Additional major and trace element analyses were undertaken at Bureau Veritas Mineral Laboratories, Canada, using inductively coupled plasma atomic emission spectroscopy (ICP-AES) for major element analysis and ICP-MS for trace element analysis.

The whole rock data set was supplemented by 117 quantitative analyses from various stratigraphic levels within the UFG, focussing on selected trace elements (Zr, Nb, Rb, Sr). This was achieved by X-ray fluorescence (XRF) at Keele University, UK, using a Niton XI3t energy-dispersive XRF spectrometer. Further details of analytical procedures and data quality are provided in Online Resources 4 and 5.

Mineral and glass analyses

Major element compositions of mineral phases and glass (both melt inclusions and groundmass glass) were analysed using a CAMECA SX 100 electron microprobe at The Open University, UK, and a JEOL JXA 8900 RL electron microprobe at the University of Göttingen, Germany. Full details of the analytical procedure and data quality are given in Online Resources 4 and 5.

Thermobarometry

In the presence of both Ti-magnetite and ilmenite, magmatic temperatures and oxygen fugacity (fO_2) estimates were calculated using the ILMAT program of Lepage (2003), applying the model of Andersen et al. (1993) with the calculation scheme of Stormer (1983) to determine values for X_{hem} , X_{ilm} , X_{mag} , X_{ulv} . The application of alternative calculation schemes (e.g. Carmichael 1967; Anderson 1968; Lindsley and Spencer 1982) led to variation of no more than 5 % in the calculated results. Equilibrium between mineral pairs was evaluated using the Mn–Mg partitioning test of Bacon and Hirschmann (1988). Pre-eruptive magmatic temperatures were also estimated using the alkali feldspar-melt thermometer of Putirka (2008). In addition to pre-eruptive temperatures, magmatic H_2O contents were also estimated, using the hygrometer of Mollo et al. (2015). Due to the absence of notable chemical zonation within Fe–Ti oxide and alkali feldspar crystals, both core and rim analyses were used in the calculations, which were based upon individual mineral (and glass) analyses rather than averages. Full details of the applied procedure are provided in Online Resource 4.

Geochemical modelling

To investigate the petrogenesis of the UFG trachytes, a variety of approaches were applied, including Rayleigh fractionation and batch melting trace element modelling, least squares mass balance modelling, and thermodynamic modelling. The full details of the applied models are given in Online Resource 4.

Results

Major and trace element geochemistry

Whole rock major element data for the UFG units are reported in Online Resource 6. Notably, analyses for Furnas A and B have lower SiO_2 and Na_2O values, in addition to elevated LOI values, which are indicative of post-depositional alteration, and lead to misleadingly low calculated peralkalinity indices. For this reason, neither analysis is considered in the following discussion.

The UFG whole rock samples exhibit little major element variation, with SiO_2 values that cluster around 63 wt%, low MgO contents (<0.5 wt%), and high total alkali abundances (~13 wt%). Using the total alkali versus silica diagram of Le Bas et al. (1986), the UFG samples plot uniformly within the trachyte field (Fig. 3). The UFG pumice falls are mildly peralkaline, with peralkalinity indices (P.I.) that range from 1.02 to 1.13. The UFG lava domes range from metaluminous to mildly peralkaline, with a P.I. range of 0.95–1.05. All of the samples are silica undersaturated, with a range in normative nepheline of 0.15–6.10 wt%, correlating positively with P.I.. Due to the crystal poor nature of the UFG pumice falls, groundmass glass analyses are very similar in composition to whole rock and can be classified as trachyte (Fig. 3).

Trace element analyses for the UFG are presented in Online Resource 6. Chondrite-normalised REE patterns shown in Fig. 4a, and compared with a standard OIB (Sun and McDonough 1989), reveal a relatively uniform enrichment in LREEs relative to HREEs, with La_N/Yb_N ratios between 11 and 27. They also display a pronounced, negative Eu anomaly, with Eu/Eu^* values between 0.12 and 0.22. This anomaly is somewhat smaller in the lava domes (0.30–0.32), and in the first two eruptions of the UFG (Furnas A and B, 0.37 and 0.45, respectively), though the effects of post-depositional alteration cannot be ruled out in the latter.

Figure 4b presents a multi-element diagram, showing UFG data and a standard OIB, normalised against the primitive mantle composition of Sun and McDonough (1989). Notable troughs occur for Ba, Sr, Eu, Ti, and, in some cases, Pb. In particular, Sr and Ba reach concentrations which are lower than primitive mantle concentrations.

Mineral chemistry

All of the mineral chemical data discussed in the following section is provided in Online Resource 7.

Feldspar

The majority of UFG feldspar analyses lie at the anorthoclase-sanidine boundary, with compositions within Or_{23-48} ,

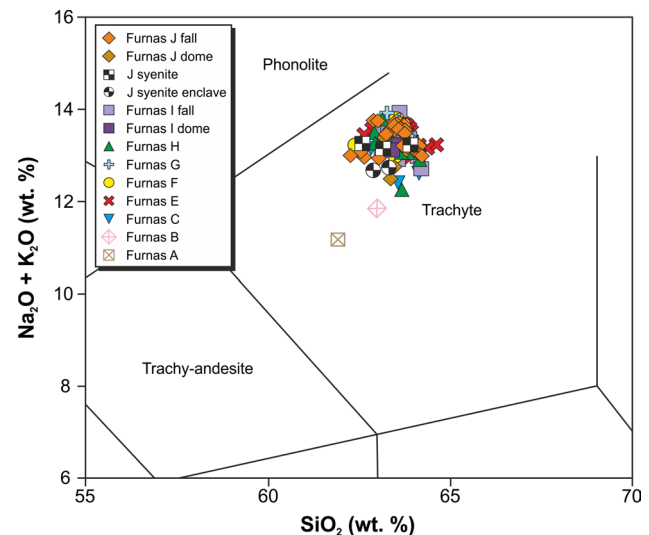


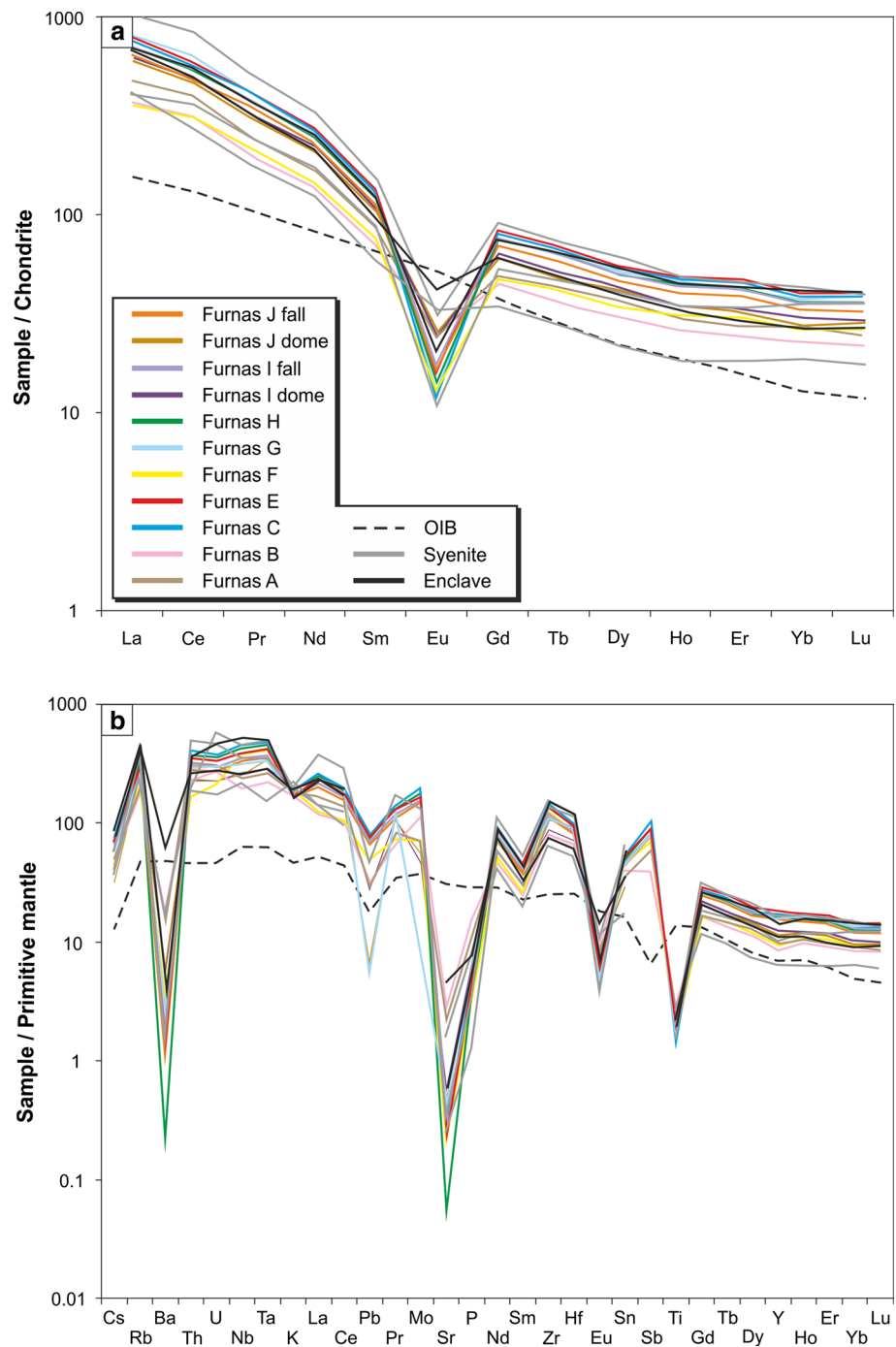
Fig. 3 UFG compositions plotted into the Total alkali–silica classification scheme of Le Bas et al. (1986). Whole rock analyses, groundmass glass and melt inclusions are plotted together and show uniform trachytic compositions

Ab_{48-69} , An_{0-7} (Fig. 5a). Feldspar phenocryst analyses from the lava domes of Furnas J and Furnas I exhibit a similar range to their respective pumice falls, though they extend to slightly more sodic compositions. Substituting elements such as Ba, Sr, and Fe are frequently below the detection limit. Feldspars from the Furnas J syenites are similar in composition to those of the other UFG units, with a range of Or_{26-57} , Ab_{43-72} , An_{0-5} , and no variation between syenite and enclave. Concentrations of Ba, Sr, and Fe are, like the UFG, frequently below detection limits.

Clinopyroxene

For classification, the ten component scheme of Marks et al. (2008) was applied. When plotted in the quadrilateral, three component (Wo-En-Fs) scheme of Morimoto et al. (1988) (Fig. 5b), UFG clinopyroxenes lie at the diopside-augite boundary. Analyses derived from clinopyroxene phenocrysts in the Furnas J and Furnas I lava domes do not show any significant compositional variation compared to their corresponding pumice falls. The analyses are best defined using Di, Hd, and Aeg, which exhibit ranges of Di_{45-70} , Hd_{12-37} , and Aeg_{2-18} (Fig. 5b). In Di–Hd–Aeg space, some Furnas samples exhibit a trend of enrichment in Hd, with little to no increase in the aegirine component. Other units show more homogenous compositions (e.g. Furnas H), though some comparatively Di-rich outliers are present. The Furnas J lava dome has a larger compositional range than the Furnas J pumice fall, extending to more Di-rich compositions. Similarly, the Furnas I lava dome clinopyroxenes show a somewhat greater range than the Furnas I pumice fall, though this

Fig. 4 Trace element variation diagrams for the UFG, **a** REE compositions of the UFG. All data are normalised relative to chondrite values and plotted against typical oceanic island basalt (OIB) compositions (Sun and McDonough 1989), **b** multi-element trace element variation diagram of the UFG. All data are normalised to primitive mantle values (Sun and McDonough 1989)



range extends to more Aeg-rich compositions. Two analysed groundmass crystals are outliers within this unit, with compositions of $\text{Di}_{33}, \text{Hd}_{37}, \text{Aeg}_{26}$, and $\text{Di}_{12}, \text{Hd}_7, \text{Aeg}_{58}$, and so may represent a continuation of the described trend, where Aeg enrichment begins after initial Hd enrichment.

The clinopyroxene analyses from the Furnas J syenite ejecta lie in stark contrast with the rest of the UFG units, with a compositional range from diopside, through aegirine-augite, to aegirine ($\text{Qd}_{0-93}, \text{Jd}_{0-8}, \text{Aeg}_{6-96}$). The Ti-aegirine component is generally present in small concentrations

(~6 mol%), but extends to values as high as 34 mol% in a small number of analyses. In Di-Hed-Aeg space, the data reveal a poorly defined trend of initial enrichment in hedenbergite, followed by aegirine enrichment.

Fe–Ti oxides

Quantitative analyses of both Ti-magnetite and ilmenite for the UFG are presented in the $\text{TiO}_2\text{–FeO–Fe}_2\text{O}_3$ ternary system in Fig. 5c. Analyses were stoichiometrically

corrected for ferric iron. Ti-magnetite shows a compositional range of Magnetite_{35–68}, Ulvöspinel_{22–63}, Spinel_{1–10}, with an average Al₂O₃ content of 0.9 wt%. MgO and MnO contents reach 1.3 and 2.7 wt%, respectively. The available ilmenite analysis from the Furnas J dome has a relatively MgO and MnO rich composition (Ilmenite₇₈, Pyrophanite₉, Geikielite₁₃). Ti-magnetite from the Furnas J syenite ejecta is similar to that of other UFG lithologies, with a range of Magnetite_{49–67}, Ulvöspinel_{32–50}, Spinel_{0–1}, whereas ilmenite from the syenites is notably enriched in MnO and shows essentially no MgO (Ilmenite_{53–75}, Pyrophanite_{24–46}, Geikielite_{0–1}).

Biotite

Biotite analyses from the UFG are Mg-rich, characterised by relatively constant Fe/(Fe + Mg) ratios (~0.41), and variable Si contents [5.44–5.71 atoms per formula unit (apfu)] (Fig. 5d). UFG analyses also exhibit high Ti contents (0.56–0.93 apfu). Occupation of the two hydroxyl sites is characterised by variably high F contents (0.32–1.63 apfu) and low Cl contents (<0.02 apfu). Biotite from the Furnas J syenite ejecta are very similar to those found in the pumice falls and lava domes, with variable Si contents (5.54–5.90 apfu), and high Ti (0.42–0.84 apfu). Similarly, the hydroxyl sites are dominated by F (0.75–1.12 apfu), with little Cl (<0.02 apfu). However, Fe/(Fe + Mg) ratios extend to larger values (0.44–0.72).

Amphibole

Following the nomenclature of Leake et al. (1997), all of the Furnas I dome analyses are Na–Ca amphiboles of magnesio-katophoritic to richteritic composition (Fig. 5e, f). Chlorine concentrations are generally close to the detection limit (<0.004 apfu). The role of F in the hydroxyl site was not quantified. Amphiboles in the Furnas J syenites exhibit a larger compositional range, which includes Na–Ca amphiboles of kataphoritic composition, and Na amphiboles of ferric-ferronyböitic and arfvedsonitic composition. Fluorine concentrations in the hydroxyl site range from 0.45 to 1.21 apfu and show no significant variation between Na–Ca and Na groups. Chlorine contents are extremely low (<0.006 apfu).

Discussion

In this section, the combined data set is used to provide insights into the pre-eruptive magmatic system of the young (<5 ka) volcanic activity of Furnas, exploring first the pre-eruptive P–T–fO₂ conditions of the UFG trachytes, and the petrogenetic processes that generate them. Ultimately,

this allows the construction of a conceptual model for the magma plumbing system. Second, the compositional zonation patterns observed throughout the UFG, both within and between individual eruptions, are used to place constraints upon the temporal evolution of the magma reservoir from which the UFG eruptions were fed.

P–T–fO₂ constraints on the plumbing system

Thermobarometric models

Due to the scarcity of ilmenite in the Furnas samples, temperature and fO₂ estimates for the trachytes, based upon two-oxide equilibria, could only be achieved for the Furnas J lava dome and the syenite ejecta. Ti-magnetite analyses from the Furnas J pumice fall were also tested for equilibrium with Furnas J dome ilmenite, and a single additional pairing was found to be suitable for calculation. Estimates for the Furnas J lava dome indicate a temperature range of ~80 °C between 814 and 897 °C and fO₂ values that are slightly above the FMQ buffer (Fig. 6a). As the temperature decreases, the fO₂ values increase slightly, reaching approximately 0.5 log units above the FMQ curve at the lowest recorded temperatures. These values are representative of the final pre-eruptive magmatic temperatures and/or syn-eruptive conditions within the plumbing system, due to the rapid re-equilibration timescales of coexisting Fe–Ti oxides (e.g. Gardner et al. 1995; Venezky and Rutherford 1999; Pimentel et al. 2015). The pre-eruptive redox conditions of the UFG trachytes appear to be more reducing than those of nearby Sete Cidades, where a total range of ~FMQ + 1 to ~MH – 1 was reported by Beier et al. (2006). Interestingly, in the case of Sete Cidades volcano, both the caldera-forming and the post-caldera deposits (the latter being comparable to the UFG) record significantly more oxidising conditions than the UFG. This substantial variation may be linked to degassing of H₂ following the thermal disassociation of H₂O (cf. Mungall and Martin 1995) and would indicate a greater degree of magma degassing in the post-caldera magma system of Sete Cidades compared to that of Furnas. Equilibrium testing for the syenites indicates only four equilibrium pairs, yielding temperatures between 621 and 780 °C, and fO₂ values that become more reduced at lower temperatures, ranging from 0.5 to 2 units below FMQ (Online Resource 8).

Pre-eruptive temperature estimates for the trachytes were also calculated via alkali feldspar-melt thermometry (Putirka 2008), using an initial estimated pressure input of 100 MPa. The effect of pressure is, in this case, negligible, with a change in calculated temperature of <1 °C following a pressure increase of one order of magnitude. In each case, a range of up to 19 whole rock and groundmass glass compositions were used as potential equilibrium liquid

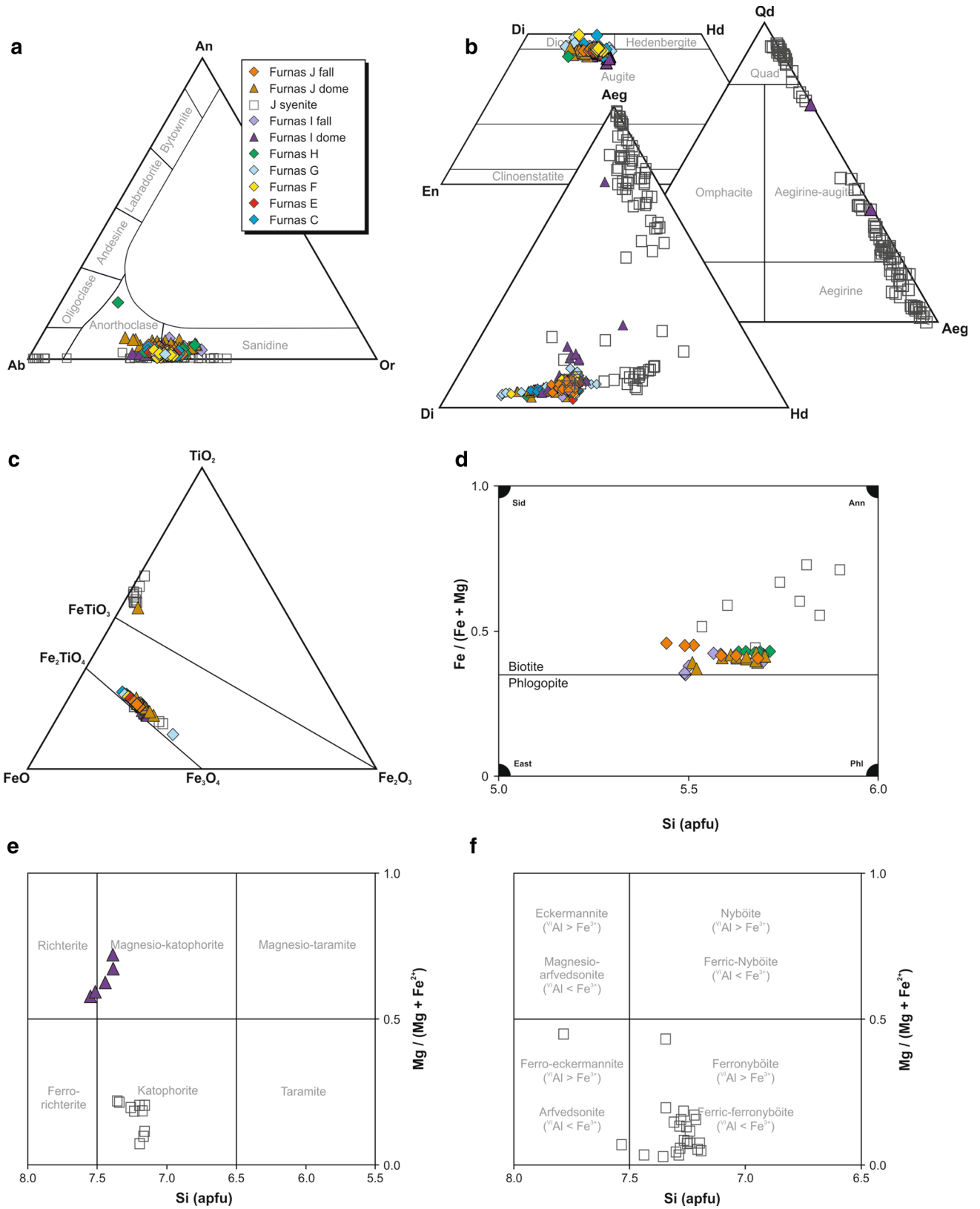


Fig. 5 Mineral compositions of the UFG, **a** alkali feldspar compositions plotted into the ternary An–Ab–Or system, **b** clinopyroxene compositions plotted into the pyroxene quadrilateral and, where relevant, the ternary Qd–Jd–Aeg system (Morimoto et al. 1988) and the ternary Di–Hd–Aeg system, **c** Fe–Ti oxide compositions plotted into the $\text{TiO}_2\text{--FeO--Fe}_2\text{O}_3$ ternary system, **d** biotite compositions for the Furnas H, I, and J pumice falls, and the Furnas J syenites, **e** Na–Ca amphibole compositions for the Furnas I lava dome and the Furnas J syenites, plotted in the scheme of Leake et al. (1997), **f** Na-amphibole compositions for the Furnas J syenites, plotted in the scheme of Leake et al. (1997)

compositions and the most suitable was identified using the $\text{KD}_{\text{Or-Ab}}$ equilibrium test presented by Mollo et al. (2015). In the majority of cases, all of the alkali feldspar analyses from a single eruption were found to be well equilibrated with a single groundmass glass or whole rock analysis from the same eruption. However, in the case of Furnas J, the most applicable equilibrium liquid for alkali feldspars from both the pumice fall and the lava dome was found to be a whole rock analysis of the lava dome (SM7-1) (Online Resource 8).

Calculated temperature estimates for the UFG are summarised in Fig. 6b. The UFG exhibits a temperature range of 855–965 °C, with some fluctuation between individual eruptions. For example, the pumice fall and lava dome of Furnas J record temperatures in excess of 915 °C, whilst the majority of earlier UFG eruptions yield somewhat lower temperatures that are generally below 900 °C. Furnas F represents a notable exception to this, with a narrow range of hotter temperature estimates between 922 and 935 °C. Estimates derived from syenite ejecta are similar to those of the UFG falls and domes, but with greater scatter, ranging from 892 to 969 °C. The limited available estimates for the enclaves yield temperatures that are not distinguishable from the host syenites. The comparatively wide range of temperature estimates observed in the syenites may reflect the temporal variation in feldspar nucleation that inevitably results from in situ crystallisation.

Estimates of pre-eruptive $\text{H}_2\text{O}_{\text{melt}}$ of the trachytes were determined for each unit of the UFG via alkali feldspar-melt equilibria (Mollo et al. 2015). In each case, equilibrium liquid compositions were carried over from alkali feldspar-melt thermometry. Due to the dependency of calculated $\text{H}_2\text{O}_{\text{melt}}$ values upon the input temperature, the average temperature estimate derived from alkali feldspar-melt thermometry was used as an input for each of the UFG units. Calculated $\text{H}_2\text{O}_{\text{melt}}$ estimates for the UFG are summarised in Fig. 6b. Results indicate that the UFG was hydrous, with average $\text{H}_2\text{O}_{\text{melt}}$ values around 5 wt%, and a total range from 2.4 to 6.1 wt%. This range is consistent with the measured water contents of trachytic glass inclusions within alkali feldspars sampled from Furnas J (Jeffery 2016). The negative correlation between

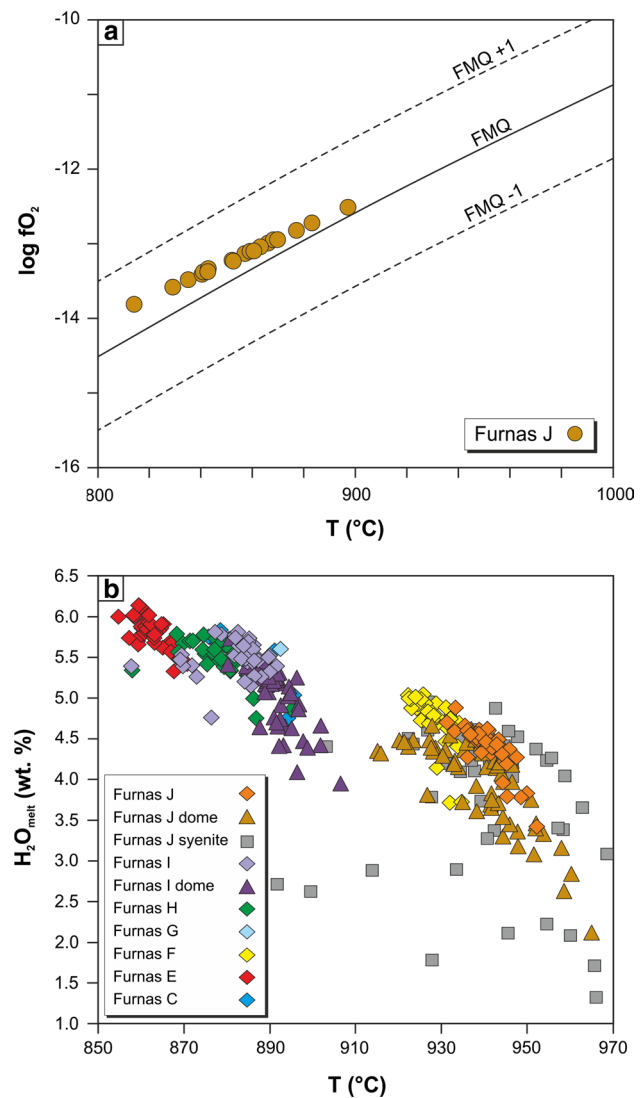


Fig. 6 Summarised results of thermobarometry and hygrometry, **a** $T\text{--}f_{\text{O}_2}$ estimates for Furnas J derived from two-oxide models. FMQ buffer reaction curve calculated at 100 MPa, **b** $T\text{--}\text{H}_2\text{O}_{\text{melt}}$ estimates produced via alkali feldspar-melt models

temperature and $\text{H}_2\text{O}_{\text{melt}}$ indicates that the UFG trachytes were H_2O undersaturated prior to eruption. There is some evidence for variation in $\text{H}_2\text{O}_{\text{melt}}$ within individual eruptions. In the case of Furnas J and Furnas I, the late-stage lava domes both yield $\text{H}_2\text{O}_{\text{melt}}$ estimates that extend to somewhat lower values than the preceding pumice falls. This is most obvious for Furnas J, where the dome analyses return a range that extends as low as 2.4 wt%, which is >1 wt% lower than the lowest estimate for the Furnas J pumice fall. Syenite ejecta and their enclaves yield a range of estimates that are comparable to those of Furnas J and Furnas F, but show substantial scatter, ranging from 1.3 to 5.3 wt%.

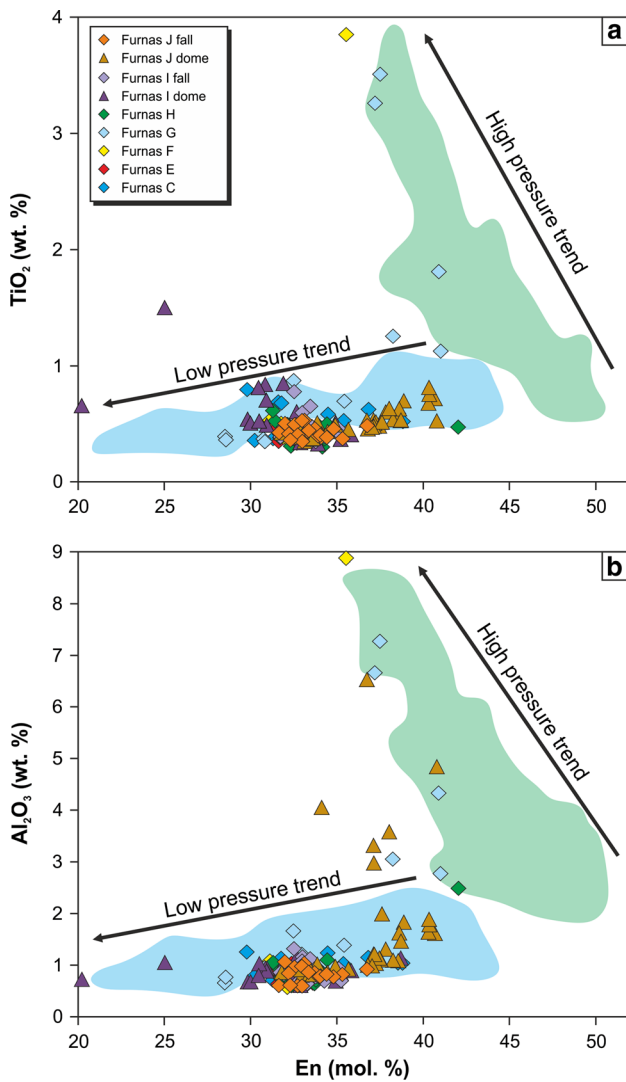


Fig. 7 Variations in TiO_2 and Al_2O_3 content of clinopyroxene, plotted with reference to the high- and low-pressure trends identified at nearby Sete Cidades (Beier et al. 2006)

Equilibrium testing of clinopyroxene indicates that none of the reported UFG clinopyroxenes were in equilibrium with their host trachytes, precluding quantitative estimation of pre-eruptive conditions. Beier et al. (2006) noted a similar lack of crystal-melt equilibrium in trachytes from Sete Cidades. Application of hypothetical liquid compositions derived from Rhyolite-MELTS modelling indicates that the most suitable equilibrium liquid compositions are broadly trachyandesitic. Despite the lack of equilibrium, clinopyroxene chemistry may still allow qualitative insights into the magma system. Beier et al. (2006) observed two trends in clinopyroxene Al_2O_3 and TiO_2 content at Sete Cidades: (1) high Al_2O_3 and TiO_2 in the mafic rocks, and (2) low Al_2O_3 and TiO_2 in the trachytes. These trends were attributed to high- and low-pressure crystallisation conditions,

respectively. With the exception of a small amount of clinopyroxenes from a number of the UFG units, the UFG clinopyroxene population adheres to the low Al_2O_3 and TiO_2 trend (Fig. 7). On this basis, the bulk of clinopyroxene crystallisation may be constrained to shallow crystal conditions. The small number of high-pressure clinopyroxenes may reflect an earlier high-pressure fractionation step, similar to that of Sete Cidades, which is poorly represented in the UFG trachytes.

Constraints from H_2O solubility

Further constraints can be placed upon the minimum depth of the magma reservoir through the calculation of saturation pressures of the melt. To achieve this, the solubility models of Di Matteo et al. (2004) and Papale et al. (2006) were applied. The former applies a polynomial fit of experimental data to parameterise H_2O solubility and yields an estimated minimum pressure of $122 \text{ MPa} \pm 15 \text{ MPa}$, assuming water saturation at 5.7 wt%. The second model was applied using a representative anhydrous MET whole rock composition, and an input temperature of $908 \text{ }^\circ\text{C}$ (average alkali feldspar-melt equilibria-derived T of UFG), and produces a minimum pressure of 156 MPa.

Petrogenesis of the UFG trachytes

To investigate the roles of fractional crystallisation and partial melting of crustal rocks in the generation of the UFG trachytes, we employ Rayleigh fractionation and batch melting models, using sample 98SM33 of Elliott et al. (2007) as the parental magma, or a partially melted crustal gabbro, respectively (see Online Resource 4 for full details). Applied mineral-melt partition coefficients are given in Table 1. Modelling of compatible trace elements such as Sr, Ba, and Ni indicate that batch melting models of gabbroic crust with variable mineral assemblages are unable to reproduce the low concentrations observed in the Furnas rocks (Fig. 8a, b, c). Instead, Rayleigh fractionation curves provide a better fit and suggest that the compositional variability of the UFG trachytes [least evolved trachytes (or LETs) to most evolved trachytes (or METs)] is largely controlled by ~75–85 % fractionation of an alkali basaltic parent. Modelled fractionation curves for incompatible elements such as Nb and Zr indicate a similar range of fractionation values (~70–85 %) (Fig. 8d). This is consistent with nearby Sete Cidades (Beier et al. 2006) and Fogo (e.g. Storey 1981; Widom et al. 1992), as well as other notable peralkaline centres such as Terceira (e.g. Mungall and Martin 1995) and Pantelleria (e.g. Neave et al. 2012), where fractional crystallisation plays a dominant role in magma evolution.

To further investigate the role of fractional crystallisation, major element mass balance models were applied

Table 1 Partition coefficients used for trace element modelling

| | Pl | Cpx | Ol | Mt | Bt | Ap | Alk fsp | Ilm |
|------------------------------|--------|--------|--------|-------|--------|--------|---------|--------|
| Step 1 RFC and batch melting | | | | | | | | |
| Nb | 0.0100 | 0.100 | 0.0100 | 0.900 | 0.0880 | | | |
| Zr | 0.0100 | 0.120 | 0.0600 | 0.700 | 0.0170 | | | |
| Sr | 10.0 | 0.700 | 0.0200 | | 0.700 | 1.20 | | |
| Ba | 0.300 | 0.0400 | 0.0300 | | 3.68 | 0.0500 | | |
| Ni | 0.270 | 2.50 | 34.0 | 48.0 | 23.9 | | | |
| Step 2 RFC | | | | | | | | |
| Nb | | | | 0.180 | 0.740 | | 0.00400 | 11.6 |
| Zr | | | | 0.250 | 0.100 | | 0.0300 | 0.0740 |
| Sr | | | | 0.000 | 0.490 | 2.00 | 2.30 | 0.740 |
| Ba | | | | 0.100 | 10.0 | 8.00 | 5.90 | |
| Ni | | | | | 89.6 | 0.450 | 0.905 | 6.20 |

All coefficients sourced from the GERM database (Nielsen 2006); see Online Resource 4 for details

Pl plagioclase, Cpx clinopyroxene, Ol olivine, Mt Ti-magnetite, Bt Biotite, Ap apatite, Alk fsp alkali feldspar, Ilm ilmenite

using the mineral compositions reported in Table 2. The results indicate that evolution from parental basalt to the LET (step 1) can be adequately explained ($\sum r^2 = 0.023$) by 87.5 % fractional crystallisation of a phase assemblage including clinopyroxene (45 %), plagioclase (24 %), Ti-magnetite (12 %), olivine (7 %), biotite (10 %), and apatite (2 %) (Table 3). The second evolutionary step (LET to MET) can be achieved by a further 38 % fractional crystallisation of an assemblage dominated by alkali feldspar (94 %), with subsidiary Ti-magnetite (1 %), ilmenite (1 %), biotite (3 %), and apatite (1 %) ($\sum r^2 = 0.018$). Together, steps 1 and 2 indicate that the differentiation of parental basalt to the most evolved UFG trachyte can be accounted for by a total of 92 % fractional crystallisation of an assemblage that is dominated by clinopyroxene and plagioclase during basalt to trachyte evolution, and alkali feldspar during continued evolution within the trachytes. The predicted late-stage prevalence of alkali feldspar fractionation is in agreement with the mineral assemblages of the trachytes, where alkali feldspar constitutes the dominant phenocryst phase. Predicted mafic-to-intermediate assemblages are broadly similar to available descriptions of mafic-to-intermediate rocks from across São Miguel (e.g. Moore 1991; Beier et al. 2006; Jeffery unpublished data).

The results of 150 Rhyolite-MELTS models indicate that a liquid of similar composition to the UFG trachytes (with respect to MgO, SiO₂, Al₂O₃, total alkalis, peralkalinity index, and H₂O_{melt}) can be produced via fractional crystallisation of a hydrous (1.5 wt%) alkali basalt parent under low pressure (150 MPa), at redox conditions close to the FMQ buffer reaction curve (Fig. 9). These conditions are consistent with those predicted via thermobarometrical models. This model predicts olivine as the liquidus phase (1265 °C), followed by clinopyroxene and Ti-magnetite

(1115 °C), with clinopyroxene crystallisation ceasing at 905 °C. This is followed by apatite (1065 °C) plagioclase feldspar (985 °C), biotite (900 °C), and ilmenite (795 °C). The model reaches compositions similar to the METs of the UFG at ~815 °C, at which point the melt has undergone ~86 % fractionation of olivine (~10 %), clinopyroxene (~42 %), feldspar (~22 %), Ti-magnetite (~10 %), apatite (~1 %), and biotite (~1 %). Higher pressure (300 and 500 MPa) models fail to reach the silica and total alkali values exhibited by the UFG, peaking at approximately trachyandesitic compositions. Lower pressure (50 MPa) models underestimate liquid Al₂O₃ contents due to the earlier crystallisation of feldspar, leading to higher peralkalinity indices than observed in the UFG. Polybaric models also perform well, generating liquid compositions similar to the UFG when initial water content and redox conditions are set to 1.5 wt% and the FMQ buffer, respectively. However, these models yield silica and total alkali contents that are slightly lower than the UFG.

In addition to the described petrogenetic models, geochemical characteristics of the UFG highlight the role of fractional crystallisation. The observed depletion of Ti and P indicate the fractionation of Fe–Ti oxides and apatite, respectively, whilst the extreme depletion of Sr and Ba, and the large negative Eu anomaly that is typical of Furnas trachytes, reflects extensive fractionation of both plagioclase and alkali feldspars. Based upon the presented evidence, it is suggested that the trachytes of the UFG are primarily derived from between ~70 and ~90 % low-pressure fractional crystallisation of an alkali basaltic parental magma under redox conditions close to the FMQ buffer, although a minor contribution from crustal assimilation or partial melting of crustal lithologies cannot be ruled out entirely (e.g. Widom and Farquhar 2003; Snyder et al.

Fig. 8 Results of trace element modelling. Calculated Rayleigh fractionation curves are labelled with RFC, and batch melting curves are labelled with BM. *Ticked* intervals represent 10 % fractionation or partial melting. Analyses derived from ED-XRF are given with *filled symbols*, whilst those produced via WD-XRF or ICP-MS are shown with *open symbols*. Parental compositions are shown with a *black circle*

2004; Genske et al. 2013, 2014; Larrea et al. 2014). These elevated fractionation values are compatible with estimates from other peralkaline systems such as Pantelleria (White et al. 2009; Neave et al. 2012), Sete Cidades (Renzulli and Santi 2000; Beier et al. 2006), Fogo (Widom et al. 1992) and the East African Rift (Scaillet and Macdonald 2003; Rooney et al. 2012). The LETs are likely to have evolved via at least 70 % fractionation from alkali basalt, through trachybasalt, basaltic trachyandesite, and trachyandesite, fractionating an assemblage that is initially dominated by clinopyroxene with subsidiary olivine and with feldspar becoming more significant at intermediate compositions. The METs are then produced by further fractional crystallisation of an assemblage dominated by alkali feldspar, with small amounts of biotite, Ti-magnetite, and apatite. The crystallisation of ilmenite is not predicted until 795 °C, which may suggest that the UFG magmas were predominantly stored at temperatures in excess of this value.

Origin of syenitic cognate xenoliths

Syenitic ejecta from the Furnas J eruption are considered to represent the final stages of crystallisation (<800 °C), characterised by an assemblage that, whilst still dominated by alkali feldspar, also includes a range of clinopyroxene compositions (diopside, augite, hedenbergite, aegirine-augite, aegirine), sodic amphiboles, Ti-magnetite, and ilmenite. Amphiboles and clinopyroxenes of broadly similar composition were observed as interstitial phases in the Furnas I lava dome, suggesting that, like ilmenite, they only form at temperatures that are lower than those at which the UFG magmas were stored.

Based upon their geochemical and petrographical characteristics, the syenites are considered to represent in situ crystallisation at the thermal boundary zone of a magma reservoir, rather than cumulates from which significant melt volumes were extracted (cf. Widom et al. 1993). For example, evolution from LET to MET has been shown to be controlled by the fractionation of alkali feldspar, biotite, Ti-magnetite, and apatite. Neither Na-amphibole nor Na-clinopyroxene is predicted in thermodynamic models, and both significantly decrease the fit of mass balance models when included, suggesting that the syenites cannot represent a cumulate from which the trachytes are derived.

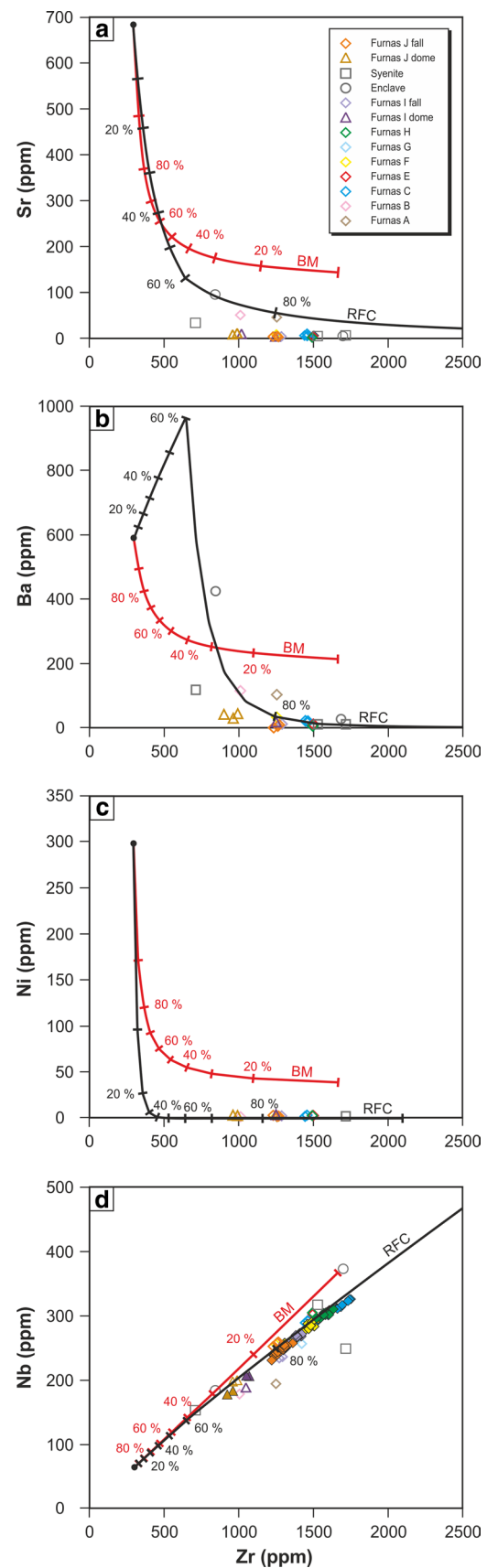


Table 2 Mineral compositions used for major element mass balance modelling

| Mineral | Pl | Alk fsp | Ol | Cpx | Mt-1 | Mt-2 | Ilm | Bt | Ap |
|--------------------------------|-------|---------|-------|-------|-------|-------|-------|-------|-------|
| SiO ₂ | 54.71 | 65.15 | 40.44 | 52.26 | | | | 36.70 | |
| TiO ₂ | | 0.09 | | 0.84 | 14.65 | 20.47 | 48.97 | 6.51 | |
| Al ₂ O ₃ | 29.28 | 19.00 | | 2.21 | 0.99 | 4.39 | 0.03 | 13.30 | |
| Fe ₂ O ₃ | | | | 3.37 | 40.01 | 26.61 | | 6.85 | |
| FeO | 0.82 | 0.37 | 11.75 | 0.38 | 39.25 | 42.36 | 42.26 | 9.01 | |
| MnO | | | 0.22 | 0.12 | 2.63 | 0.52 | 3.93 | 0.41 | |
| MgO | | 0.01 | 47.32 | 16.71 | 1.77 | 4.99 | 3.31 | 13.91 | |
| CaO | 10.30 | 0.54 | 0.28 | 22.37 | | | | | 55.70 |
| Na ₂ O | 4.83 | 6.87 | | 0.28 | | | | 0.97 | |
| K ₂ O | 0.43 | 6.69 | | | | | | 8.71 | |
| P ₂ O ₅ | | | | | | | | | 41.82 |
| H ₂ O | | | | | | | | | 0.59 |

Mineral compositions for alkali feldspar (Alk fsp), Ti-magnetite (Mt-1), and ilmenite (Ilm) were taken from the data set of this study. Compositions for plagioclase (Pl), olivine (Ol), clinopyroxene (Cpx), Ti-magnetite (Mt-2) and biotite (Bt) were taken from Beier et al. (2006)

Similarly, the syenites and their enclaves show a wide range of trace element compositions that encompass those of the UFG, rendering many of the UFG eruptions less evolved than the syenites. If the syenite nodules did represent residual cumulates, they should be uniformly less evolved than the UFG trachytes. Furthermore, the syenites exhibit Eu anomalies that are comparable to, or greater than, those of the UFG trachytes. In the UFG rocks, a negative correlation is observed between Eu/Eu* and differentiation indices such as Zr and Nb, suggesting that the accumulation of feldspar could only result in a less negative, or even positive, Eu anomaly. As such, it is suggested that the syenites represent near-complete, in situ crystallisation of unerupted trachytic magmas similar to those of the UFG. Their trace element variability is ultimately derived from the varying degrees of fractionation that also yield trace element diversity in the UFG trachytes, though hydrothermal alteration and the presence of accessory phases such as zircon also play a role.

The role of magma mixing/mingling processes

In addition to FC processes, a number of features of the UFG eruption products provide evidence for the role of replenishment and magma mingling between two variably evolved trachytes in the magma storage zone. For example, enclaves hosted in syenite ejecta display chilled margins, rounded, lobate forms, as well as large alkali feldspar phenocrysts which exhibit similar compositions to host syenite feldspar and pronounced disequilibrium textures at their rims, suggesting that they are xenocrysts derived from the host syenite. These features are all indicative of liquid emplacement, suggesting that the enclaves represent

trachyte-syenite mingling in the highly crystalline mush zone at the margins of a magma reservoir.

Additionally, the presence of banded pumices in the uppermost pumice fall deposit of Furnas J (Lf) may provide direct evidence for mingling between two variably evolved trachytes during the later stages of the eruption, as has been described for the C11 explosive eruption (~1000 ¹⁴C y B.P.) of Caldeira volcano, Faial (Pimentel et al. 2015). Grey bands within the predominantly white pumices of Lf exhibit markedly lower vesicularity and thus may have been volatile poor (Cole et al. 1995). The same study presented evidence for variable viscosity of the two types based upon deformation characteristics, as well as geochemical analyses for each type, which, on the basis of Zr content, suggests that the dense grey pumice may be marginally less evolved. However, it should be noted that both analyses are ~250–300 ppm lower in Zr than the Lf analyses of this study and are instead comparable to the Furnas J lava dome analyses. This observation implies that there may be a relatively sharp geochemical transition from high Zr (~1200 ppm) to low Zr (~900 ppm) within the Lf unit. Additionally, the analyses of Cole et al. (1995) exhibit Ba contents an order of magnitude greater than typical UFG concentrations, which may indicate local resorption of alkali feldspar. A single analysis within the data set of this study reveals an anomalous enrichment of Sr (95 compared to <10 ppm), which may reflect the same process.

A conceptual model for the UFG magma plumbing system

Based upon the range of pressure estimates derived from H₂O solubility models and thermodynamic models

Table 3 Results of major element mass balance modelling

| Sample | Liquid composition | | | Fractionating phases | |
|-------------------------------------|--------------------|-------|--------|----------------------|------|
| | Basalt | | LET | | |
| | 98SM33 | | S022 | | |
| | Obs | Calc | Obs | | |
| <i>Step 1: Alkali basalt to LET</i> | | | | | |
| SiO ₂ | 46.22 | 46.17 | 63.20 | Cpx | 45.1 |
| TiO ₂ | 3.41 | 3.29 | 0.57 | Plg | 23.9 |
| Al ₂ O ₃ | 10.89 | 10.93 | 17.83 | Mt | 12.3 |
| FeO ^T | 11.44 | 11.48 | 3.34 | Bt | 10.2 |
| MnO | 0.17 | 0.19 | 0.22 | Ol | 7.1 |
| MgO | 11.53 | 11.53 | 0.38 | Ap | 1.6 |
| CaO | 12.01 | 12.04 | 0.88 | | |
| Na ₂ O | 2.12 | 2.13 | 7.37 | | |
| K ₂ O | 1.59 | 1.65 | 6.03 | | |
| P ₂ O ₅ | 0.63 | 0.6 | 0.09 | | |
| Σr ² | 0.023 | | | | |
| F | 0.125 | | | | |
| Sample | Liquid composition | | | Fractionating phases | |
| | LET | | MET | | |
| | S022 | | SM 1-2 | | |
| | Obs | Calc | Obs | | |
| <i>Step 2: LET to MET</i> | | | | | |
| SiO ₂ | 63.20 | 63.28 | 63.16 | Alk fsp | 94.1 |
| TiO ₂ | 0.57 | 0.57 | 0.42 | Bt | 3.2 |
| Al ₂ O ₃ | 17.83 | 17.86 | 17.40 | Mt | 1.4 |
| FeO ^T | 3.34 | 3.44 | 4.21 | Ap | 0.7 |
| MnO | 0.22 | 0.22 | 0.30 | Ilm | 0.6 |
| MgO | 0.38 | 0.4 | 0.32 | | |
| CaO | 0.88 | 0.83 | 0.77 | | |
| Na ₂ O | 7.37 | 7.31 | 7.75 | | |
| K ₂ O | 6.03 | 5.94 | 5.49 | | |
| P ₂ O ₅ | 0.09 | 0.16 | 0.06 | | |
| Σr ² | 0.018 | | | | |
| F | 0.618 | | | | |

Ap apatite, Plg plagioclase, Alk fsp alkali feldspar, Cpx clinopyroxene, Bt biotite, Ol olivine, Mt magnetite, Ilm ilmenite. LET least evolved trachyte, MET most evolved trachyte

(~122–156 MPa), the magma storage region of the UFG eruptions can be constrained to shallow crustal depths between 3 and 4 km (based upon an average crustal density of 2800 kg/m³). This range compares favourably with previous estimates of magma storage beneath Furnas volcano (Machado 1972; Sigmundsson et al. 1995; Blanco et al. 1997; Camacho et al. 1997; Montesinos et al. 1999) and also the shallow magma reservoir of Sete Cidades (Beier et al. 2006). The results of thermodynamic modelling, as well as the predominantly low Al₂O₃ and TiO₂ contents of clinopyroxene, suggest that much of the

compositional range from basalt to MET is present at this depth (Fig. 10).

Based upon the decreasing melt densities predicted by Rhyolite-MELTS from basalt to trachyte (~2580 to ~2230 kg/m³), it is suggested that the reservoir beneath Furnas is compositionally zoned, with dense mafic magmas at the base, and low-density, hydrous silicic magmas at the roof. Trachytic magmas at the roof may be further enriched in volatiles via upward migration of a vapour phase, further stabilising the density stratification. This configuration may account for the apparent Daly Gap observed at Furnas

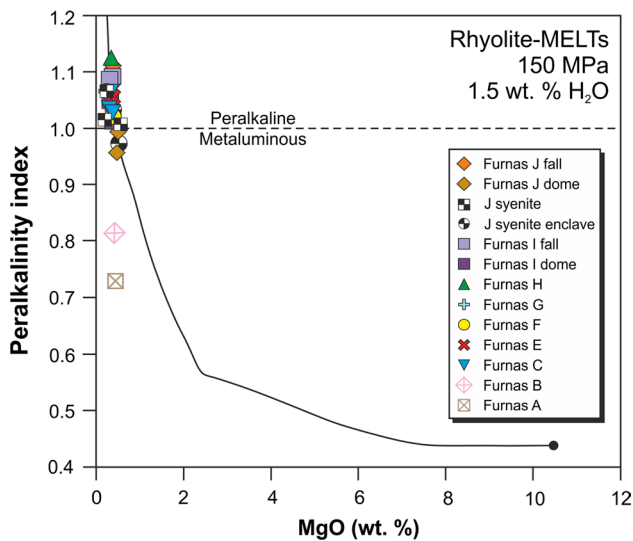


Fig. 9 Results of Rhyolite-MELTS fractional crystallisation modelling from an alkali basalt (black circle). Modelled curved produced at 150 MPa, with an initial water content of 1.5 wt%

(Guest et al. 1999), as ascending mafic magmas may stall in an established shallow crustal magma reservoir and differentiate, ultimately contributing relatively small volumes of residual trachytic liquid to the silicic cap of the reservoir rather than erupting themselves (e.g. Mahood 1984; White et al. 2009; Neave et al. 2012). The likely presence of older, frozen reservoirs at this level is envisaged to further inhibit rapid magma ascent (e.g. Zanon and Pimentel 2015). In contrast, basalts that ascend outside the confines of the caldera and avoid the shallow system reached the surface and were erupted as the monogenetic cones and fissure eruptions that are typical of the volcano’s flanks and the neighbouring Congro fissure volcanic system (Zanon 2015).

The generation of crystal poor silicic magmas, such as those of the UFG, is commonly accounted for through the application of models which involve the extraction of interstitial melt from large crystal mush bodies, citing processes such as compaction, microsettling, hindered settling, and gas-driven filter pressing (e.g. Sisson and Bacon 1999; Bachmann and Bergantz 2004; Hildreth 2004; Pistone

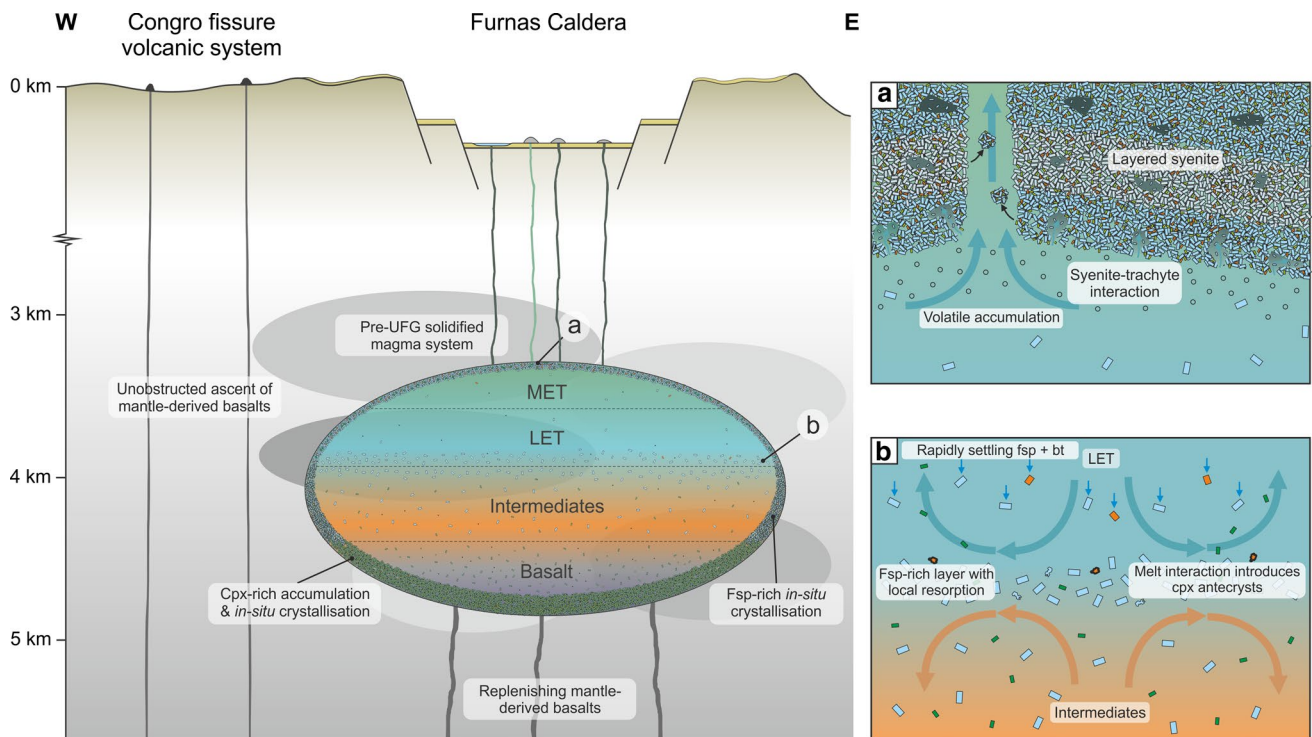


Fig. 10 A schematic model for the structure of the shallow plumbing system of Furnas volcano. A density-stratified magma reservoir with a trachytic cap is envisaged, at depths of approximately 3 to 4 km, **a** syenite ejecta derived from an upper solidification front may be sampled during eruption. The solidified remnants of pre-UFG magma systems may provide an additional source of syenitic xenoliths, **b**

interaction between individually convecting layers introduces an antecrystic clinopyroxene population to the cap, whilst rapidly settling feldspars form a Sr- and Ba-rich feldspar layer. Settling biotite phenocrysts originating from the upper most evolved trachytes (METs) are heavily resorbed by hotter underlying least evolved trachytes (LETs) and intermediates

et al. 2015). Such models are attractive primarily because of the difficulties associated with two-phase flow in relatively cool, high viscosity ($\sim 10^{4.5}$ Pa s) melts (Scaillet et al. 1998). However, their application to peralkaline magmatic systems is complicated by the apparent absence of erupted crystal-rich melts (equivalent to the ‘monotonous intermediates’ associated with non-peralkaline systems, Macdonald 2012) and also by the tendency of peralkaline melts to have lower viscosities than their metaluminous and peraluminous equivalents (e.g. Giordano et al. 2006; Neave et al. 2012). The efficiency of two-phase flow in the UFG trachytes was investigated through the estimation of crystal free melt viscosities. Application of the model of Giordano et al. (2008) indicates that the trachytes, at a temperature of 908 °C and H_2O_{melt} values from 2.4 to 5.7 wt%, have viscosities ranging from $10^{2.7}$ to $10^{4.1}$ Pa s. The UFG trachytes therefore had pre-eruptive viscosities that were up to two orders of magnitude lower than typical silicic melts (Scaillet et al. 1998), which has implications for the dominant differentiation mechanism.

The efficiency of crystal settling was evaluated through the calculation of Stokes’ settling velocities for alkali feldspar crystals (determined to be the most important crystallising phase during the later stages of magma evolution). This was supplemented with the hindered settling equation, which allows the determination of settling rates in polydispersed suspensions (Bachmann and Bergantz 2004):

$$U_{\text{hs}} = U_{\text{Stokes}} \times f(c)$$

where U_{hs} is the hindered settling velocity, U_{Stokes} is Stokes settling velocity, and $f(c)$ is a correction factor calculated as:

$$f(c) = \frac{(1 - c)^2}{(1 + c^{1/3}) \left[\frac{5c}{3(1-c)} \right]}$$

where c is equal to the crystal fraction. Crystal sizes were set to 2 mm, in accordance with petrographic observations. Results indicate that even for the highest viscosity estimate for the UFG, settling rates range from 0.51 m/a^{-1} (hindered settling, 50 % crystallinity) to 1.76 m/a^{-1} (unhindered). Settling rates for the lowest viscosity trachytes reach values as high as 13.7 m/a^{-1} (hindered settling, 50 % crystallinity) and 47 m/a^{-1} (unhindered). Assuming a volume of melt that is typical for the UFG eruptions (0.1 km^3 DRE; Booth et al. 1978), and hypothetical magma reservoir aspect ratios of 1:1:1 and 8:1:1, alkali feldspar crystals would need to travel maximum distances of ~ 470 – 235 m to be removed from the volume of erupted melt. Available radiocarbon ages (and associated errors) for the UFG indicate that repose times do not exceed 918 years, but are likely to be ~ 190 years (Furnas I to J) or less (e.g. Furnas H to I, Guest et al. 1999). Over such short timescales (190 years), alkali

feldspar crystals are capable of settling $\sim 100 \text{ m}$ at the very least (highest viscosity), but are capable of settling far greater distances (up to 9600 m using the lowest calculated viscosity). It is therefore suggested that, unlike the rhyolitic systems to which crystal mush models may apply, crystal settling was a viable and efficient mechanism of differentiation in the UFG trachytes and may account not only for the nearly aphyric nature of the erupted trachytes, but also for the rapid rates of differentiation implied by the wide range in their degree of evolution.

Syenitic ejecta are considered to represent a solidification front at the volatile-rich cap of the reservoir, in which trachytic magmas near the margins of the reservoir solidify via in situ crystallisation. Phase assemblages including aegirine, Na-amphibole, and ilmenite are consistent with near-complete in situ crystallisation, whilst the presence of miarolitic cavities and pyrrhotite indicates the accumulation of volatile phases. Crystallisation temperatures estimated from Ti-magnetite-ilmenite pairs are at least 70 °C lower than estimates for the UFG pumice falls. High temperature estimates derived from alkali feldspar-melt equilibria (892 to 969 °C) are similar to those of Furnas J and likely reflect the relatively early formation of feldspar compared to later ilmenite, aegirine, and amphibole. Enclaves within the syenites are considered to represent interaction between the upper thermal boundary layer and the crystal poor liquids in the underlying crystal poor cupola. Small volumes of crystal poor trachytes are inferred to have infiltrated the overlying crystal mush and been quenched. In Zr–Nb space, the majority of syenites and enclaves appear to lie along the same liquid line of descent as the UFG, implying a cogenetic association. Although Zr–Nb relationships could be uncoupled by the crystallisation of Zr-bearing phases, the observed rarity of zircon, as well as its restriction to pore spaces, imply that late-stage zircon was retained rather than fractionated. The substantial range in the degree of evolution observed in their whole rock chemistry (~ 700 – 1700 ppm Zr) is likely to be a temporal effect, where the composition of a given syenite clast or enclave is related to the dominant melt composition in the roof zone of the reservoir, a feature which is shown to vary considerably. Additionally, it seems probable that this cupola would be zoned to some extent, exhibiting the same degree of composition variation as the UFG trachytes themselves. We therefore invoke an ‘onion-skin’ arrangement, similar to that of Widom et al. (1993), where random sampling of various layers of the roof zone yields a range of compositions. It should also be noted that a single syenite analysis appears to deviate from the UFG liquid line of descent in Zr–Nb space, suggesting that it may represent a fragment of an alternative liquid line of descent. It thus cannot be ruled out that some syenitic ejecta may instead represent the ‘fossilised’

remnants of a pre-UFG magmatic system, entrained pre- or syn-eruptively within the erupted trachyte.

Attempts to employ clinopyroxene thermobarometry revealed that the clinopyroxenes are not in equilibrium with the host trachytes (see above), instead being equilibrated with a broadly trachyandesitic composition. This is consistent with thermodynamic models, which suggest that clinopyroxene becomes saturated in the melt at 1115 °C and crystallises until 905 °C. Settling rates of clinopyroxene through the UFG trachytes are approximately three times faster than those of alkali feldspar, suggesting that the ubiquitous presence within the erupted trachytes requires a mechanism that is either continuous within the magma plumbing system, or occurs immediately prior to each eruption. Two possibilities for an origin are considered: (1) they are xenocrysts derived from the incorporation and assimilation of mafic wall rock material or (2) they are antecrystic relics derived from liquid–liquid interaction between the erupted trachytes and less evolved magmas. There is no petrographic evidence, such as crustal or cumulate xenoliths, to support the former hypothesis, and trace element models do not indicate partial melting as a significant petrogenetic process in the UFG. As such, the second hypothesis is favoured and, given the small but consistent presence of clinopyroxene throughout the UFG, implies the episodic interaction between trachytic and less evolved, clinopyroxene-bearing magmas, most likely at an interface between strata with contrasting densities.

Development of zoned magma bodies

Despite the apparent major element homogeneity and low eruptive volumes (generally $\sim 0.1 \text{ km}^3$ DRE, Booth et al. 1978) of the intra-caldera UFG trachytes, trace element compositions indicate substantial variation. For example, both the Furnas J and Furnas I eruptions began with an initial pumice fall, followed by a late-stage lava dome; in both cases, the lava dome is, according to a variety of differentiation indices (Zr, Nb, REE, Eu/Eu*), less evolved than its respective pumice fall. Such phenomena are generally attributed to the development of a zoned magma reservoir prior to eruption (e.g. Hildreth 1981) and suggest that the uppermost, eruptible region of the Furnas magma reservoir is able to develop rapidly and maintain compositional zonation over the timescales of the UFG. A variety of processes have been suggested to play a role in the development of zonation (e.g. Hildreth and Wilson 2007); however, due to the described low magma viscosities, it seems likely that zoning patterns in the UFG are the result of the efficient fractionation of alkali feldspar crystals, driving the melt towards MET compositions. The upwards increase in volatile content and associated decrease in viscosity may lead

to a geochemical gradient via variably efficient fractionation (e.g. Furnas J, L1 to Lf).

The preservation of zonation in the UFG magmas until eruption indicates the absence of efficient convective cells able to rehomogenise the reservoir, which may be accounted for by a strong density gradient linked to the $\sim 4 \text{ wt\%}$ range in estimated melt water contents (cf. Hildreth and Wilson 2007). Alternatively, compositional zonation might be stabilised through the generation of multiple, individually convecting layers (e.g. Huppert and Sparks 1984), though the applicability of such a model in a low volume of magma ($< 0.3 \text{ km}^3$) may be restricted. If the zoning patterns of the UFG magmas are considered to have been largely unaltered by conduit processes during ascent, then the potentially sharp compositional contact between the pumice falls and lava domes of Furnas I and J may provide evidence for individual strata, reflecting a compositional discontinuity that was present in the magma reservoir between hydrous MET, and underlying, volatile-poor LET. This would also indicate that either a deeper part of the reservoir was tapped than in the majority of previous eruptions or that the cap had not yet regenerated the comparatively large volumes of MET associated with older eruptions (e.g. Furnas C, $\sim 0.3 \text{ km}^3$ DRE).

The zoning patterns observed in the Furnas C (and possibly F) deposits imply a more complex regime in which less evolved melts are capable of erupting prior to more evolved melts. The observed zoning patterns require more complex reservoir geometries, such as sequential eruption of multiple magma bodies, each acting as an independent evolving system (e.g. Cooper et al. 2012; Zanon et al. 2013; Pimentel et al. 2015). Alternatively, this complex zonation might originate from an irregular chamber geometry which allows lower regions to be tapped first.

Temporal evolution of the UFG magma reservoir

In addition to trace element zonation within individual eruptions, the UFG as a whole exhibits higher-order variations which provide insights into the temporal evolution of the previously described eruptible cap of the shallow crustal reservoir. The complex trace element profiles of the UFG (Fig. 11) are clearly inconsistent with the tapping of a single large magma batch, emplaced prior to the Furnas A eruption, and allowed to fractionate for ~ 5000 years (maximum age of the UFG). Instead, the observed variability suggests a cyclical regime, in which a zoned, silicic cap forms at the top of the reservoir, and is partially erupted in one or more individual eruption events, before being re-established via continued fractionation and potentially mafic replenishment (Fig. 12). This is consistent with neighbouring Fogo volcano, where a similar regime of zoned cap formation and tapping over the same time

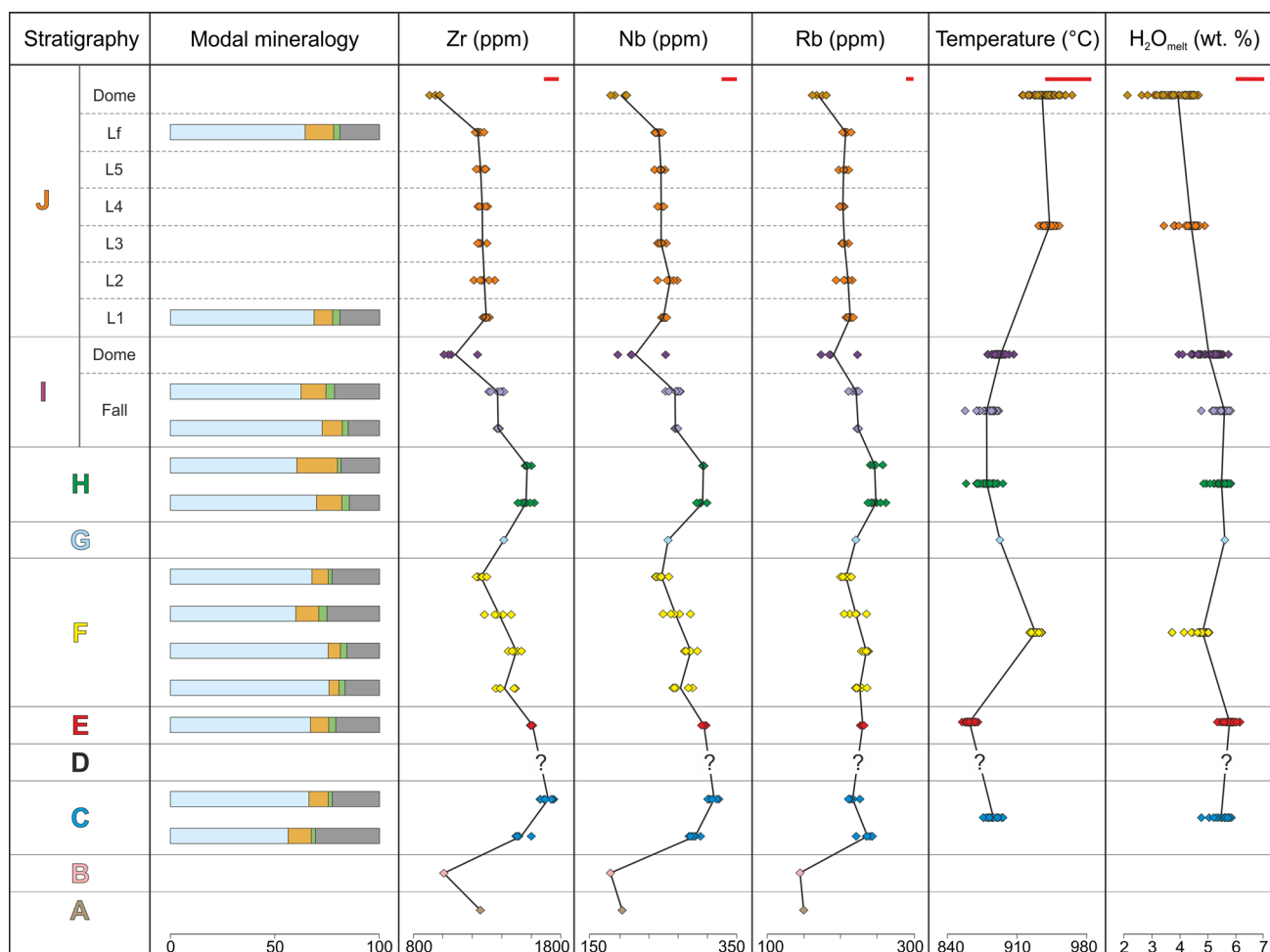


Fig. 11 Variations in trace element concentrations, temperature estimates, and H_2O_{melt} estimates with stratigraphic height. Relevant errors are shown with red lines at the top of each column. Modal

mineralogy: alkali feldspar = blue, biotite = orange, clinopyroxene = green, Fe–Ti oxides = grey

period has been suggested (Widom et al. 1992). At Furnas, four periods are recognised, during which a single, zoned magma cap was erupted sequentially in up to three individual eruptive events, reforming after each period. Due to the relatively large errors and consequent degree of overlap in the radiocarbon ages of the UFG, the repose periods between individual eruptions cannot be determined accurately. However, it is suggested that the length of the repose time is likely to be linked to the temporal model of this study, with the longest repose times being associated with periods during which the zoned, eruptible cap was being reconstructed.

The first zoned magma body of the UFG is the least constrained, but is inferred to have been tapped twice during the Furnas A and B eruptions, during a period of up to ~3000 years (Online Resource 1), followed by a lengthy re-establishment phase, during which the reservoir cap evolved via 30–40 % fractionation prior to the Furnas C

eruption. Available radiocarbon ages suggest that the Furnas B to C repose period may have been up to 918 years.

The Furnas C, E, and F eruptions together indicate a trend of progressively less evolved trachytes, suggesting that they represent sequential, and probably relatively rapid, tapping of a single zoned magma body that formed following Furnas B. This indicates that, despite the relatively substantial volume of the Furnas C eruption compared to the other UFG eruptions (0.3 km³ DRE), the system was tapped a further three times prior to re-establishment, which may be linked to the repose time between eruptions, or the overall rate of trachyte replenishment in the cap. The repose periods between these eruptions are likely to have been comparatively short, limiting the effects of continued differentiation between eruptions. This period was followed by a period of fractionation, which culminated in the Furnas H eruption, and equated to ~10–15 % fractionation. During this period, the reservoir was tapped

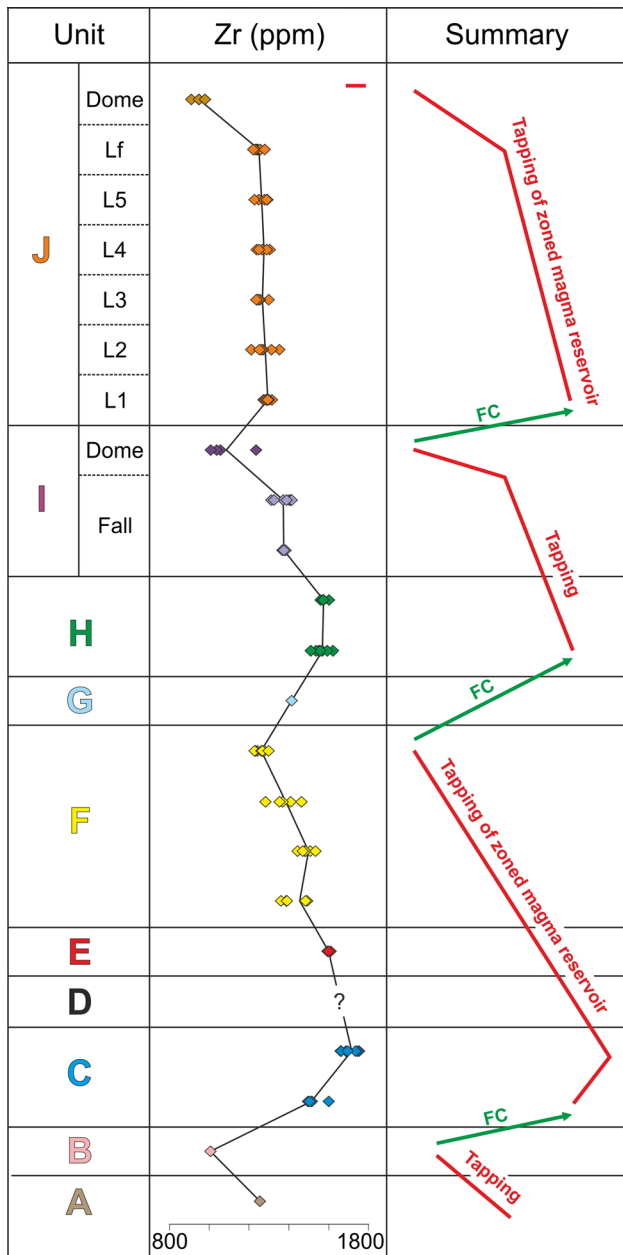


Fig. 12 A schematic model for cyclic replenishment and eruption phases throughout the UFG, based upon trace element compositions. See text for discussion. FC fractional crystallisation

once by the Furnas G eruption, but this did not inhibit continued fractionation prior to Furnas H.

The third zoned magma body is considered to have formed prior to Furnas H and been sequentially tapped by the Furnas H and I eruptions. This is consistent with the comparatively short repose time between Furnas H and I (~20 years, Booth et al. 1978; Guest et al. 1999). Together, these eruptions define a trend of decreasing evolution that reverses the effect of fractional crystallisation by up to ~35 %. This was followed by a repose period of

~190 years, during which the system evolved by up to 20 % to produce the fourth zoned magma body, which was subsequently erupted in the Furnas J eruption. Based upon the similarity between the trace element profiles of the Furnas I and J eruptions, it is possible that the shallow magmatic system of Furnas has entered a new repose period whilst the eruptible cap reforms, though it should be noted that the current repose time of 385 years is approximately twice that of the I to J repose time (~190 years).

Implications for silicic volcanism on São Miguel

The magma system of Furnas shares a number of features with neighbouring Fogo and Sete Cidades volcanoes. For example, a prominent, subcaldera, shallow crustal (~3–4 km) magma reservoir has been identified at each volcano, in which mafic-to-intermediate magmas differentiate primarily via fractional crystallisation to form a cap of variably evolved trachyte (e.g. Storey 1981; Widom et al. 1992; Renzulli and Santi 2000; Beier et al. 2006). Evidence for magma mingling and mixing, such as phenocryst textures (Beier et al. 2006), hybrid magmas (Storey et al. 1989), and banded pumices (Widom et al. 1992; this study), is present at all three centres. However, felsic eruptive products are somewhat less dominant at Sete Cidades (~75 vol%; Moore 1991) than at Furnas and Fogo (≥90 vol%; Moore 1991), and the Daly Gap observed at both Furnas and Fogo is absent (Beier et al. 2006). One solution is that the upper, trachytic caps of the magma plumbing systems of both Furnas and Fogo are more developed, possibly due to periods of reduced basaltic input to the lower part of the systems (cf. Macdonald 2012). They would therefore form a more significant barrier against the ascent and eruption of basaltic magmas. In contrast, the magma plumbing system of Sete Cidades may be somewhat less developed and so unable to entirely prevent the ascent of basaltic magmas.

Conclusions

The compositional characteristics of the young (<5 ka), intra-caldera trachytes of Furnas volcano can be adequately accounted for by extended fractional crystallisation from an alkali basalt parental magma at minimum depths of ~3–4 km. Fractionation is initially dominated by clinopyroxene, which is eventually replaced by alkali feldspar in the latter stages of differentiation. Despite their major element homogeneity, trace element compositions highlight the prominent role of continued fractional crystallisation within the trachytes, revealing up to 50 % fractionation from the least evolved trachytes (LETs) to the most evolved trachytes (METs). This process is enhanced by the accumulation of water and the development of peralkalinity.

The magma plumbing system is shown to comprise a single prominent zone in which alkali basalts evolve through to trachytic residual liquids. Due to density variations, this reservoir is envisaged to be stratified, with high-density mafic magmas at the base and relatively cool, hydrous trachytes forming an upper cap. Rare syenitic ejecta are considered to represent in situ, near-complete crystallisation of the trachytes in the roof zone of the reservoir, which is sampled randomly during eruption. Mingling and mixing within the trachytes is recorded by trachytic enclaves within the syenite ejecta and by the presence of banded pumices in Furnas J. Further evidence for mixing can be seen in the ubiquitous presence of small quantities of clinopyroxene antecrysts within all of the UFG eruptions, which are likely to be introduced to the trachytes during interaction with underlying, hotter, intermediate magmas. The presence of such a shallow reservoir may account for the Daly Gap at Furnas, as ascending mafic magmas are intercepted before reaching the surface, instead ponding in the lower reservoir and fractionating towards felsic compositions.

Despite the low volumes of the UFG eruptions ($\leq 0.3 \text{ km}^3$ DRE), variations in trace element concentrations within individual eruptions indicate zonation of the upper cap of the reservoir prior to eruption, with the most evolved melts typically being the first to erupt. Evidence exists for the presence of both chemical gradients and relatively sharp compositional stratification, which, in the case of the latter, are considered to represent sharp compositional discontinuities in the pre-eruptive magma reservoir, providing direct evidence for the presence of density stratification. Less distinct gradients likely result from the accumulation of volatiles, as well as the associated increase in crystal settling rates. Overall, trace element profiles of the UFG are not compatible with continued fractionation and tapping of a single magma batch and are instead indicative of a cyclic regime in which a compositionally stratified trachytic cap has formed and been sequentially erupted in up to three individual eruptions, before being re-established. This study suggests that there have been up to four such cycles of eruption and re-establishment within the UFG, with the 385-year repose time since the most recent eruption most likely representing a re-establishment phase.

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References

- Andersen DJ, Lindsley DH, Davidson PM (1993) QUILF: a Pascal program to assess equilibria among Fe–Mg–Mn–Ti oxides, pyroxenes, olivine and quartz. *Comput Geosci* 19:1333–1350
- Anderson AT (1968) Oxidation of the La Blanche Lake titaniferous magnetite deposit, Québec. *J Geol* 76:528–547
- Avanzinelli R, Bindi L, Menchetti S, Conticelli S (2004) Crystallisation and genesis of peralkaline magmas from Pantelleria volcano, Italy: an integrated petrological and crystal-chemical study. *Lithos* 73:41–69
- Bachmann O, Bergantz GW (2004) On the origin of crystal-poor rhyolites: extracted from batholithic crystal mushes. *J Petrol* 45:1565–1582
- Bachmann O, Deering CD, Ruprecht JS, Huber C, Skopelitis A, Schnyder C (2012) Evolution of silicic magmas in the Kos-Nisyros volcanic center, Greece: a petrological cycle associated with caldera collapse. *Contrib Mineral Petrol* 163:151–166
- Bacon CR, Hirschmann MM (1988) Mg/Mn partitioning as a test for equilibrium between coexisting Fe–Ti oxides. *Am Mineral* 73:57–61
- Barberi F, Ferrara G, Santacroce R, Treuil M, Varet J (1975) A transitional basalt-pantellerite sequence of fractional crystallisation, the Boina Centre (Afar Rift, Ethiopia). *J Petrol* 16:22–56
- Beier C, Haase KM, Hansteen TH (2006) Magma evolution of the Sete Cidades volcano, São Miguel, Azores. *J Petrol* 47:1375–1411
- Blanco I, García A, Torta JM (1997) Magnetic study of the Furnas caldera (Azores). *Ann Geofis* 40:341–359
- Bohrson WA, Reid MR (1997) Genesis of silicic peralkaline volcanic rocks in an oceanic island setting by crustal melting and open-system processes: Socorro Island, Mexico. *J Petrol* 38:1137–1166
- Booth B, Croasdale R, Walker GPL (1978) A quantitative study of five thousand years of volcanism on São Miguel, Azores. *Philos Trans R Soc A* 288:271–319
- Camacho AG, Montesinos FG, Vieira R (1997) A three-dimensional gravity inversion applied to São Miguel Island (Azores). *J Geophys Res* 102:7717–7730
- Carmichael ISE (1967) The iron-titanium oxides of silicic volcanic rocks and their associated ferromagnesian silicates. *Contrib Mineral Petrol* 14:36–64
- Chester DK, Dibben JL, Duncan AM (2002) Volcanic hazard assessment in western Europe. *J Volcanol Geotherm Res* 115:411–435
- Civetta L, D'antonio M, Orsi G, Tilton GR (1998) The geochemistry of volcanic rocks from Pantelleria Island, Sicily Channel: petrogenesis and characteristics of the mantle source region. *J Petrol* 39:1453–1491
- Cole PD, Queiroz G, Wallenstein N, Gaspar JL, Duncan AM, Guest JE (1995) An historic subplinian/phreatomagmatic eruption: the 1630 AD eruption of Furnas volcano, São Miguel, Azores. *J Volcanol Geotherm Res* 69:117–135

- Cole PD, Guest JE, Queiroz G, Wallenstein N, Pacheco JM, Gaspar JL, Ferreira T, Duncan AM (1999) Styles of volcanism and volcanic hazards on Furnas volcano, São Miguel, Azores. *J Volcanol Geotherm Res* 92:39–53
- Cooper GF, Wilson CJ, Millet M, Baker JA, Smith EG (2012) Systematic tapping of independent magma chambers during the 1 Ma Kidnappers super eruption. *Earth Planet Sci Lett* 313:23–33
- Di Matteo V, Carroll MR, Behrens H, Vetere F, Brooker RA (2004) Water solubility in trachytic melts. *Chem Geol* 213:187–196
- Duncan AM, Queiroz G, Guest JE, Cole PD, Wallenstein N, Pacheco JM (1999) The Povoação Ignimbrite, Furnas volcano, São Miguel, Azores. *J Volcanol Geotherm Res* 92:55–65
- Elliott T, Blichert-Toft J, Heumann A, Koetsier G, Forjaz V (2007) The origin of enriched mantle beneath São Miguel, Azores. *Geochim Cosmochim Acta* 71:219–240
- Freire Luis J, Miranda JM, Galdeano A, Patriat P, Rossignol JC, Mendes Victor LA (1994) The Azores triple junction evolution since 10 Ma from an aeromagnetic survey of the Mid-Atlantic Ridge. *Earth Planet Sci Lett* 125:439–459
- Gardner JE, Rutherford M, Carey S, Sigurdsson H (1995) Experimental constraints on pre-eruptive water contents and changing magma storage prior to explosive eruptions of Mount St Helens volcano. *Bull Volc* 57:1–17
- Genske FS, Beier C, Haase KM, Turner SP, Krumm S, Brandl PA (2013) Oxygen isotopes in the Azores islands: crustal assimilation recorded in olivine. *Geology* 41:491–494
- Genske FS, Turner SP, Beier C, Chu MF, Tonarini S, Pearson NJ, Haase KM (2014) Lithium and boron isotope systematics in lavas from the Azores islands reveal crustal assimilation. *Chem Geol* 373:27–36
- Gente P, Dyment J, Maia M, Goslin J (2003) Interaction between the Mid-Atlantic Ridge and the Azores hot spot during the last 85 Myr: emplacement and rifting of the hot spot derived plateaus. *Geochem Geophys Geosyst* 4:8514. doi:10.1029/2003GC000527
- Giordano D, Mangiacapra A, Potuzak M, Russell JK, Romano C, Dingwell DB, Di Muro A (2006) An expanded non-Arrhenian model for silicate melt viscosity: a treatment for metaluminous, peraluminous and peralkaline liquids. *Chem Geol* 229:42–56
- Giordano D, Russell JK, Dingwell DB (2008) Viscosity of magmatic liquids: a model. *Earth Planet Sci Lett* 271:123–134
- Guest JE, Gaspar JL, Cole PD, Queiroz G, Duncan AM, Wallenstein N, Ferreira T, Pacheco JM (1999) Volcanic geology of Furnas volcano, São Miguel, Azores. *J Volcanol Geotherm Res* 92:1–29
- Guest JE, Pacheco JM, Cole PD, Duncan AM, Wallenstein N, Queiroz G, Gaspar JL, Ferreira T (2015) The volcanic history of Furnas Volcano, São Miguel, Azores. In: Gaspar JL, Guest JE, Duncan AM, Bariga FJAS, Chester DK (eds) 2015, Volcanic geology of São Miguel island (Azores archipelago). *Geol Soc Lond Mem* 44:125–134
- Harris C (1983) The petrology of lavas and associated plutonic inclusions of Ascension Island. *J Petrol* 24:424–470
- Hildreth W (1981) Gradients in silicic magma chambers: implications for lithospheric magmatism. *J Geophys Res* 86:10153–10192
- Hildreth W (2004) Volcanological perspectives on Long Valley, Mammoth Mountain, and Mono Craters: several contiguous but discrete systems. *J Volcanol Geotherm Res* 136:169–198
- Hildreth W, Wilson CJN (2007) Compositional zoning of the Bishop Tuff. *J Petrol* 48:951–999
- Huppert HE, Sparks RSJ (1984) Double diffusive convection due to crystallisation in magmas. *Annu Rev Earth Planet Sci* 12:11–37
- Jeffery AJ (2016) Petrogenesis and contrasting eruption styles of peralkaline silicic magmas from Terceira and São Miguel, Azores. Unpublished PhD thesis, Keele University, UK
- Jeffery AJ, Gertisser R, Jackson RA, O'Driscoll B, Kronz A (2016) On the compositional variability of dalyite, $K_2ZrSi_6O_{15}$: a new occurrence from Terceira, Azores. *Mineral Mag.* doi:10.1180/minmag.2016.080.018
- Jones G, Chester DK, Shoostariam F (1999) Statistical analysis of the frequency of eruptions at Furnas Volcano, São Miguel, Azores. *J Volcanol Geotherm Res* 92:31–38
- Kaula WM (1970) Earth's gravity field: relation to global tectonics. *Science* 169:982–985
- Krause DC, Watkins ND (1970) North Atlantic crustal genesis in the vicinity of the Azores. *Geophys J R Astro Soc* 19:261–283
- Larrea P, Galé C, Ubide T, Widom E, Lago M, França Z (2014) Magmatic evolution of Graciosa (Azores, Portugal). *J Petrol* 55:2125–2154
- Le Bas MJ, Le Maitre RW, Streckeisen A, Zanettin B (1986) A chemical classification of volcanic rocks based on the Total Alkali-Silica diagram. *J Petrol* 27:745–750
- Leake BE, Woolley AR, Arps CES, Birch WD, Gilbert MC, Grice JD, Hawthorne FC, Kato A, Kisch HJ, Krivovichev VG, Linthout K, Laird J, Mandarino JA, Maresch WV, Nickel EH, Rock NMS, Schumacher JC, Smith DC, Stephenson NCN, Ungaretti L, Whittaker EJW, Youzhi G (1997) Nomenclature of amphiboles: report of the subcommittee on amphiboles of the international mineralogical association, commission on new minerals and mineral names. *Can Mineral* 35:219–246
- Lepage LD (2003) ILMAT: an excel worksheet for ilmenite-magnetite geothermometry and geobarometry. *Comput Geosci* 29:673–678
- Lindsley DH, Spencer KJ (1982) Fe-Ti oxide geothermometry: reducing analyses of coexisting Ti-magnetite (Mt) and ilmenite (Ilm). *EOS Trans Am Geophys Union* 63:471
- Luis JF, Miranda JM (2008) Reevaluation of magnetic chrons in the North Atlantic between 35°N and 47°N: implications for the formation of the Azores Triple Junction and associated plateau. *J Geophys Res* 113:B10105
- Macdonald R (2012) Evolution of peralkaline silicic complexes: lessons from the extrusive rocks. *Lithos* 152:11–22
- Macdonald R, Sumita M, Schmincke HU, Bagiński B, White JC, Ilnicki SS (2015) Peralkaline felsic magmatism at the Nemrut volcano, Turkey: impact of volcanism on the evolution of Lake Van (Anatolia) IV. *Contrib Mineral Petrol* 169:34
- Machado F (1972) Acid volcanoes of San Miguel, Azores. *Bull Volc* 36:319–327
- Mahood GA (1984) Pyroclastic rocks and calderas associated with strongly peralkaline magmatism. *J Geophys Res* 89:8540–8552
- Markl G, Marks MAW, Frost BR (2010) On the controls of oxygen fugacity in the generation and crystallisation of peralkaline melts. *J Petrol* 51:1831–1847
- Marks MAW, Schilling J, Coulson IM, Wenzel T, Markl G (2008) The alkaline-peralkaline Tamazeght complex, High Atlas Mountains, Morocco: mineral chemistry and petrological constraints for derivation from a compositionally heterogeneous mantle source. *J Petrol* 49:1097–1131
- Marks MAW, Hettmann K, Schilling J, Frost BR, Markl G (2011) The mineralogical diversity of alkaline igneous rocks: critical factors for the transition from miaskitic to agpaitic phase assemblages. *J Petrol* 52:439–455
- Mollo S, Masotta M, Forni F, Bachmann O, De Astis G, Moore G, Scarlato P (2015) A K-feldspar-liquid hygrometer specific to alkaline differentiated magmas. *Chem Geol* 392:1–8
- Montesinos FG, Camacho AG, Vieira R (1999) Analysis of gravimetric anomalies in Furnas caldera (São Miguel, Azores). *J Volcanol Geotherm Res* 92:67–81
- Moore RB (1990) Volcanic geology and eruption frequency, São Miguel, Azores. *Bull Volc* 52:602–614
- Moore RB (1991) Geology of three Late Quaternary stratovolcanoes on São Miguel, Azores. *U.S. Geol Surv Bull* 1900:1–46
- Moore RB, Rubin M (1991) Radiocarbon dates for lava flows and pyroclastic deposits on São Miguel, Azores. *Radiocarbon* 33:151–164
- Morimoto N, Fabries J, Ferguson AK, Ginzburg IV, Ross M, Seifert FA, Zussman J, Aoki K, Gottardi G (1988) Nomenclature of pyroxenes. *Mineral Mag* 52:535–550

- Mungall JE, Martin RF (1995) Petrogenesis of basalt-comendite and basalt pantellerite suites, Terceira, Azores, and some implications for the origin of oceanic-island rhyolites. *Contrib Mineral Petrol* 119:43–55
- Neave DA, Fabbro G, Herd RA, Petrone CM, Edmonds M (2012) Melting, differentiation and degassing at the Pantelleria volcano, Italy. *J Petrol* 53:637–663
- Nielsen R (2006) Geochemical earth reference model (GERM) partition coefficient (Kd) database. www.earthref.org/KDD/
- Papale P, Moretti R, Barbato D (2006) The compositional dependence of the saturation surface of H₂O + CO₂ fluids in silicate melts. *Chem Geol* 229:78–95
- Peccerillo A, Donati C, Santo AP, Orlando A, Yirgu G, Ayalew D (2007) Petrogenesis of silicic peralkaline rocks in the Ethiopian rift: geochemical evidence and volcanological implications. *J Afr Earth Sci* 48:161–173
- Pensa A, Giordano G, Cas RAF, Porreca M (2015) Thermal state and implications for eruptive styles of the intra-Plinian and climactic ignimbrites of the 4.6 ka Fogo A eruption sequence, São Miguel, Azores. *Bull Volcanol* 77:99
- Pimentel A, Pacheco J, Self S (2015) The ~1000-years BP explosive eruption of Caldeira Volcano (Faial, Azores): the first stage of incremental caldera formation. *Bull Volc* 77:42
- Pistone M, Arzilli F, Dobson KJ, Cordonnier B, Reusser E, Ulmer P, Marone F, Whittington AG, Mancini L, Fife JL, Blundy JD (2015) Gas-driven filter pressing in magmas: insights into in situ melt segregation from crystal mushes. *Geology* 43:699–702
- Putirka KD (2008) Thermometers and barometers for volcanic systems. *Rev Mineral Geochem* 69:61–120
- Renzulli A, Santi P (2000) Two-stage fractionation history of the alkali basalt-trachyte series of Sete Cidades volcano (São Miguel island, Azores). *Eur J Mineral* 12:469–494
- Ridley WI, Watkins ND, MacFarlane DJ (1974) Chapter 12: The Oceanic Islands: Azores. In: Nairn AEM, Stehli FG (eds) *The ocean basins and margins, vol 2: The North Atlantic*. Plenum Press, London, pp 445–484
- Ridolfi F, Renzulli A, Santi P, Upton BGJ (2003) Evolutionary stages of crystallization of weakly peralkaline syenites: evidence from ejecta in the plinian deposits of Agua de Pau volcano (São Miguel, Azores Islands). *Mineral Mag* 67:749–767
- Rooney TO, Hart WK, Hall CM, Ayalew D, Ghiorso MS, Hidalgo P, Yirgu G (2012) Peralkaline magma evolution and the tephra record in the Ethiopian Rift. *Contrib Mineral Petrol* 164:407–426
- Scaillet B, Macdonald R (2001) Phase relations of peralkaline silicic magmas and petrogenetic implications. *J Petrol* 42:825–845
- Scaillet B, Macdonald R (2003) Experimental constraints on the relationships between peralkaline rhyolites of the Kenya Rift Valley. *J Petrol* 44:1867–1894
- Scaillet B, Holtz F, Pichavant M (1998) Phase equilibrium constraints on the viscosity of silicic magmas 1. Volcanic-plutonic comparison. *J Geophys Res* 103:27257–27266
- Schmincke HU, Weibel M (1972) Chemical study of rocks from Madeira, Porto Santo and São Miguel, Terceira (Azores). *Neues Jahrb Mineral Abh* 117:253–281
- Searle R (1980) Tectonic pattern of the Azores spreading centre and triple junction. *Earth Planet Sci Lett* 51:415–434
- Self S, Gunn BM (1976) Petrology, volume and age relations of alkaline and saturated peralkaline volcanics from Terceira, Azores. *Contrib Mineral Petrol* 54:293–313
- Sibrant ALR, Hildenbrand A, Margues FO, Wiess B, Boulesteix T, Hübscher C, Lüdmann T, Costa ACG, Catalão JC (2015) Morpho-structural evolution of a volcanic island developed inside an active oceanic rift: S. Miguel Island (Terceira Rift, Azores). *J Volcanol Geotherm Res* 301:90–106
- Sigmundsson F, Tryggvason E, Alves MM, Alves JL, Pálsson K, Ólafsson H (1995) Slow inflation of the Furnas volcano, São Miguel, Azores, suggested from initial levelling and global positioning system measurements. *Geophys Res Lett* 22:1681–1684
- Sisson TW, Bacon CR (1999) Gas-driven filter pressing in magmas. *Geology* 27:613–616
- Snyder DC, Widom E, Pietruszka AJ, Carlson RW (2004) The role of open-system processes in the development of silicic magma chambers: a chemical and isotopic investigation of the Fogo A trachyte deposit, São Miguel, Azores. *J Petrol* 45:723–738
- Storey M (1981) Trachytic pyroclasts from Agua de Pau volcano, São Miguel, Azores: evolution of a magma body over 4000 years. *Contrib Mineral Petrol* 78:423–432
- Storey M, Wolff JA, Norry MJ, Marriner GF (1989) Origin of hybrid lavas from Agua de Pau volcano, São Miguel, Azores. In: Saunders AD, Norry MJ (eds) *Migmatism in the ocean basins*. Blackwell, London, pp 161–180
- Stormer JC Jr (1983) The effects of recalculation on estimates of temperature and oxygen fugacity from analyses of multi-component iron-titanium oxides. *Am Mineral* 68:586–594
- Sun S, McDonough WF (1989) Chemical and isotopic systematics of ocean basins: implications for mantle composition and processes. In: Saunders AD, Norry MJ (eds) *Migmatism in the ocean basins*. *Geol Soc Lond Spec Publ* 42:313–346
- Trua T, Deniel C, Mazzuoli R (1999) Crustal control in the genesis of Plio-Quaternary bimodal magmatism of the Main Ethiopian Rift (MER): geochemical and isotopic (Sr, Nd, Pb) evidence. *Chem Geol* 155:201–231
- Venezky DY, Rutherford MJ (1999) Petrology and Fe–Ti oxide reequilibration of the 1991 Mount Unzen mixed magma. *J Volcanol Geotherm Res* 89:213–230
- Vogt PR, Jung WY (2004) The Terceira Rift as hyper-slow, hotspot-dominated oblique spreading axis: a comparison with other slow-spreading plate boundaries. *Earth Planet Sci Lett* 218:77–90
- Walker GPL, Croasdale R (1971) Two plinian-type eruptions in the Azores. *J Geol Soc London* 127:17–55
- Wallenstein N (1999) Estudo da história recente e do comportamento eruptivo do vulcão do Fogo (S. Miguel, Açores). Avaliação preliminar do hazard. Unpublished PhD thesis, Departamento de Geociências Universidades dos Açores, São Miguel Island (Portugal)
- White JC, Parker DF, Ren M (2009) The origin of trachyte and pantellerite from Pantelleria, Italy: insights from major element, trace element, and thermodynamic modelling. *J Volcanol Geotherm Res* 179:33–55
- Widom E, Farquhar J (2003) Oxygen isotope signatures in olivines from São Miguel (Azores) basalts: implications for crustal and mantle processes. *Chem Geol* 193:237–255
- Widom E, Schmincke HU, Gill JB (1992) Processes and timescales in the evolution of a chemically zoned trachyte: Fogo A, São Miguel, Azores. *Contrib Mineral Petrol* 111:311–328
- Widom E, Gill JB, Schmincke HU (1993) Syenite nodules as a long-term record of magmatic activity in Agua de Pau Volcano, São Miguel, Azores. *J Petrol* 34:929–953
- Wolff JA (1987) Crystallisation of nepheline syenite in a subvolcanic magma system: Tenerife, Canary Islands. *Lithos* 20:207–223
- Zanon V (2015) Conditions for mafic magma storage beneath fissure zones at oceanic islands. The case of São Miguel island (Azores archipelago). In: Caricchi L, Blundy JD (eds) *Chemical, physical and temporal evolution of magmatic systems*. *Geol Soc Lond Spec Publ* 422. doi:10.1144/SP422.4
- Zanon V, Pimentel A (2015) Spatio-temporal constraints on magma storage and ascent conditions in a transtensional tectonic setting: the case of the Terceira Island (Azores). *Am Mineral* 100:795–805
- Zanon V, Kueppers U, Pacheco JM, Cruz I (2013) Volcanism from fissure zones and the Caldeira central volcano of Faial Island, Azores archipelago: geochemical processes in multiple feeding systems. *Geol Mag* 150:536–555