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Extensive CO₂ degassing in the upper mantle beneath oceanic basaltic volcanoes: First insights from Piton de la Fournaise volcano (La Réunion Island)

G. Boudoire^{a,b,*}, A.L. Rizzo^{c,d}, A. Di Muro^{b,e}, F. Grassa^c, M. Liuzzo^c

^a *Laboratoire Géosciences Réunion, Université de La Réunion, Institut de Physique du Globe de Paris (IPGP), Sorbonne Paris-Cité, UMR 7154 CNRS, F-97715 Saint-Denis, France*

^b *Observatoire Volcanologique du Piton de la Fournaise (OVPF), Institut de Physique du Globe de Paris (IPGP), Sorbonne Paris-Cité, UMR 7154 CNRS, Université Paris Diderot, Bourg Murat, France*

^c *Istituto Nazionale di Geofisica e Vulcanologia, Sezione di Palermo, Via Ugo La Malfa 153, 90146 Palermo, Italy*

^d *Dipartimento di Fisica e Scienze della Terra, Università degli Studi di Ferrara, Italy*

^e *Institut de Physique du Globe de Paris (IPGP), Sorbonne Paris-Cité, UMR 7154 CNRS, Université Paris Diderot, F-75005 Paris, France*

Abstract

In spite of its major role on the atmospheric volatile budget, climate, and tracking magmatic transfers, mantle (CO₂) degassing below volcanoes is still poorly understood. Most of the studies on this scientific topic lack constraint on the CO₂ concentration of primary melts, the depth at which it starts degassing, and the extent of this process in the mantle. In this study of Piton de la Fournaise (PdF) volcano, we couple geochemistry of low solubility gases (He, Ar, CO₂, δ¹³C) in fluid inclusions (FIs) and petro-chemistry of magmatic inclusions on a set of olivine and clinopyroxene crystals from basalts and ultramafic enclaves.

We constrain basaltic melt degassing at PdF over a large pressure range (from 4 GPa up to the surface). Based on CO₂-He-Ar systematics, we infer that extensive degassing occurs already in the upper mantle (4–1 GPa) and it is favored by multiple steps of magma ponding and differentiation up to the mantle-crust underplating depth (0.4 GPa). Thus, we calculate that basaltic melts injected at crustal depth (<0.4 GPa) have already exsolved ~94 ± 5 wt% of their primary CO₂ content in accordance with (1) the evolved and degassed signature of erupted lavas and (2) the weakness of inter-eruptive gas emissions in the active area bearing low-temperature vapor-dominated fumaroles. Our results at PdF strongly contrast with previous findings on other ocean island volcanoes having a higher magma production rate and faster magma ascent, like Kilauea (Hawaii), whose basalts experience only limited extent of differentiation and degassing. We propose that extensive degassing already in the upper mantle can be a common process for many volcanoes of the Earth and is tightly dependent on the dynamics of magma ascent and differentiation across multiple ponding zones.

Based on the modeling developed in this study, we propose a new estimation of the CO₂ content (up to 3.5 ± 1.4 wt%) in primary basaltic melts at PdF leading to a carbon content in the mantle source of 716 ± 525 ppm. This new estimation is considerably higher than the few previous calculations performed for Ocean Island Basalts (OIB) systems. Another implication of this work involves the possible bias between the δ¹³C measured in volcanic gas emissions (<-6‰) and that of primary vapour phase (-0.5 ± 0.5‰) constrained in this work. This bias would confirm the early step of extensive CO₂ degassing within the upper mantle and could represent an alternative for the hypotheses of carbon recycling or mantle heterogeneity in support of the low δ¹³C signature of some mantle reservoirs. This study bears significant implications on the global budget of volcanic volatile emissions, chiefly regarding the contribution of past and future emissions of volcanic CO₂ to climate dynamics, and on volcanic gas monitoring.

* Corresponding author at: Observatoire Volcanologique du Piton de la Fournaise (OVPF/IPGP), Impasse de l'Observatoire, Bourg Murat 97418, France.

E-mail address: guillaume.boudoire@gmail.com (G. Boudoire).

1. INTRODUCTION

Magmatic degassing is an exciting field of investigation to develop reliable models for forecasting volcanic activity (Tazieff and Tonani, 1963; Le Guern et al., 1982; Allard et al., 1991; Scarpa, 2001; Aiuppa and Federico, 2004; Liuzzo et al., 2013; Rizzo et al., 2006, 2009, 2015). Among major volatiles emitted from volcanoes, CO₂ has a low solubility in melts and is released at great depth strongly contributing to the deep carbon cycle (Marty and Jambon, 1987; Dasgupta and Hirschmann, 2010; Marty et al., 2013; Manning, 2014; Lee et al., 2016; Mason et al., 2017 and references therein). CO₂ is thus the major non-aqueous constituent of magmatic gases and the principal carrier for other trace volatiles like noble gases (Barry et al., 2014; Gilfillan et al., 2014). Consequently, it is extensively used to constrain deep degassing when coupled to noble gases (e.g., ⁴He/CO₂, CO₂/Ar, ⁴He/⁴⁰Ar*) due to the low but distinct solubility between these species (Jambon et al., 1986; Burnard, 2004; Burnard et al., 2003, 2014; Aubaud et al., 2004; Paonita and Martelli, 2007; Paonita et al., 2012). The isotope composition of carbon ($\delta^{13}\text{C}$) of CO₂ is also an important tracer of magmatic degassing, because it fractionates during this process moving towards lower values (Javoy et al., 1978; Matthey, 1991; Trull et al., 1993; Shaw et al., 2004).

Even if the volcanic contribution to the total CO₂ budget in the Earth atmosphere is minor with respect to anthropogenic emissions, it strongly influences the climate dynamics with disastrous consequences for the ecosystems (Le Guern et al., 1975; Allard et al., 1991; Oppenheimer et al., 2011; Burton et al., 2013; Carn et al., 2017). A critical issue to constrain deep CO₂ degassing below volcanoes and evaluate the primary magmatic volatile budget is that most of the studies: (1) focus on late-stage products (as residual glasses or free gases), which are more prone to be affected by secondary effects (Marty et al., 1992; Caracausi et al., 2003; Capasso et al., 2005; Aubaud et al., 2004, 2006); (2) deal with Mid Ocean Ridge Basalts (MORB) systems and only a few data concern Ocean Island Basalts (OIB) systems (see Barry et al. (2014) for a complete review); (3) lack of constraints on pressures of exsolution, most of barometric data being estimated from experimental and modeling studies on noble gases (Caracausi et al., 2003; Sarda and Guillot, 2005; Iacono-Marziano et al., 2010; Paonita et al., 2012).

In this study, we focus on a set of crystals (olivine, clinopyroxene) from various products (pyroclasts and ultramafic enclaves) and eruptive sites (central and peripheral areas) of the Piton de la Fournaise shield volcano (La Réunion Island), which is considered as a reference example of basaltic activity in an OIB geodynamic

setting (Courtilot et al., 2003). The eruptive products were selected with the aim at tracking a broad range of pressure, from the upper mantle to the crustal volcano plumbing system, and thus to constrain the extent of magma degassing and evolution during its ascent. We adopt an approach that couples gas geochemistry (He, Ar, CO₂, $\delta^{13}\text{C}$) of fluids inclusions and petro-chemistry of pyroclasts and enclaves (crystal composition, fluid inclusion composition and barometry) in order to (1) constrain the volatile content of primary basaltic melts and (2) estimate the extent of magmatic evolution and degassing below the volcano as a function of depth. The PdF volcano was targeted due to (1) its intense eruptive activity (one eruption each nine months since 1985; Roult et al., 2012) contrasting with limited gas emissions during inter-eruptive periods (see Di Muro et al. (2016) for a review), (2) the homogeneity of the mantle source limiting the initial variability in composition (noble gases, CO₂ and $\delta^{13}\text{C}$, Graham et al., 1990; Füre et al., 2011; Schiano et al., 2012; Vlastelic and Pietruzka, 2016), and (3) similarities between its plumbing system and those inferred for many other ocean basaltic volcanoes (Michon et al., 2015). These features make this volcano an interesting natural laboratory to study deep magma transfer and degassing from mantle to crustal level (Liuzzo et al., 2015; Di Muro et al., 2016; Boudoire et al., 2017a,b).

2. GEOLOGICAL SETTING

Located in the Indian Ocean, La Reunion Island (latitude: -21° ; longitude: 56°) is the youngest volcanic island of the chain attributed to the activity of hot spot, which produced the Deccan flood basalt province in India 65 Myrs ago (Mahoney et al., 2002). The volcanic edifice rises 7 km-high on the ocean floor, which consists of a Upper Cretaceous-to Paleocene oceanic crust. The Moho is located at 10.2–13 km below sea level (b.s.l.) according to seismic investigations (Gallart et al., 1999; Fontaine et al., 2015). A <3 km-thick layer of dense material interpreted as the underplating of ultramafic rocks between the crust and the mantle lithosphere was inferred by geophysical studies (Charvis et al., 1999; Fontaine et al., 2015). The island is composed by two volcanic edifices (Piton des Neiges: 3070 m a.s.l and PdF: 2632 m a.s.l.) that emitted geochemically similar magmas of the transitional series (Upton and Wadsworth, 1966).

The currently active volcano is the PdF, whose activity began ca. 530 kyrs ago on the southwest flank of the Piton des Neiges volcano (Gillot and Nativel, 1989). Basalts are the only products of PdF activity and they are inferred to originate from partial melting of the upper mantle (4 GPa; Fretzdorff and Haase, 2002). The mantle source

was described as homogeneous with constant noble gases and Sr-Nd-Hf-Os isotope signature (Graham et al., 1990; Staudacher et al., 1990; Albarède et al., 1997; Hanyu et al., 2001; Schiano et al., 2012; Peters et al., 2016). Only minor isotope variations are documented in the eruptive products and have been attributed to either (1) a distinct component intrinsic to the plume itself or to the incorporation of the ambient Indian asthenosphere into the upwelling plume (Pietruszka et al., 2009; Di Muro et al., 2014; Vlastelic and Pietruszka, 2016), and/or (2) small degrees of late-stage crustal contamination (Luais, 2004; Pietruszka et al., 2009). All basalts of PdF belong to the transitional series (Upton & Wadsworth, 1966; Albarède et al., 1997; Boivin and Bachèlery, 2009), with the exception of old evolved alkaline basalts bearing abundant plagioclase (Valer et al., 2017a).

Since 1640 (i.e., the beginning of permanent colonization of the island), most of the PdF eruptions (>95%) occurs inside the Enclos Fouqué caldera (Fig. 1; Villeneuve and Bachèlery, 2006; Michon et al., 2015), formed between ~5.5 and ~3.0 kyrs BP during the Bellecombe explosive events (Ort et al., 2016). There, seismic hypocenters are mostly located at crustal depth (i.e. from 0.5 km a.s.l. to 11 km b.s.l.) consistent with the presence of several shallow crustal reservoirs (Nercessian et al., 1996; Di Muro et al., 2014; Coppola et al., 2017). The Enclos Fouqué cliff marks a geomorphological limit between the central activity inside the caldera and the peripheral activity outside it (Fig. 1). The peripheral activity (<5% of historical eruptions) is mainly concentrated within three converging rift zones:

the SE rift zone (SERZ), the NE rift zone (NERZ) and the NW rift zone (NWRZ). SERZ and NERZ are almost aseismic and are mostly related to lateral magma transfers from the shallow part of the central plumbing system (Michon et al., 2015). Conversely the NWRZ, which connects the old PdN volcanic edifice with the western flank of PdF volcano is characterized by deep seismicity located within the lithospheric mantle (>11 km b.s.l.). Petrological studies highlight that melts erupted within the NWRZ are more primitive and richer in gas than evolved and degassed melts usually erupted in the central area (Bureau et al., 1998; Di Muro et al., 2014, 2016; Coppola et al., 2017). At least two main zones of magma ponding were inferred within the upper mantle below the PdF volcano and they have been linked to eruptions occurring within the NWRZ. The deepest one (~30–35 km-depth), holds rare petrological evidences, and has been inferred by seismic studies (Gallart et al., 1999; Massin, 2009; Fontaine et al., 2015) and modeling of stress fields related to the edifice loading (Gerbault et al., 2017). The second one corresponds to the geophysically defined underplating layer (~15 km b.s.l.) and is in accordance with most of the barometric data obtained on NWRZ eruptive products (Bureau et al., 1998; Boudoire, 2017). Based on the spatio-temporal location of recent seismic hypocenters, some authors have proposed that central crustal reservoirs are fed by lateral transfers of melts at mantle level below the NWRZ, starting from a 30 km-deep offset deep plumbing system located between PdF and Piton des Neiges volcanoes (Michon et al., 2015; Boudoire et al., 2017a, 2017b). Interestingly, this zone located between both volcanic edifices is also set by the highest soil CO₂ fluxes, which contrasts with the weakness of inter-eruptive gas emissions in the central area (Liuzzo et al., 2015; Di Muro et al., 2016).

3. SAMPLES AND METHODS

In this study, we target fast quenched eruptive products and ultramafic enclaves from both the central and the peripheral (NWRZ) area of PdF (Fig. 1). Eruptive products from the peripheral area are olivine phenocrysts from Piton de Caille (PDC; <5 kyrs) and Trous Blancs (TB047; <10 kyrs) tephtras. These products were targeted due to (1) the negligible amount of recycled xenocrysts and (2) their geochemical signature (major and traces elements, Sr-Nd isotopes) spanning the geochemical variability of recent PdF magma (Valer, 2016). Similarly, in order to limit the effect of crystal recycling, we selected among the products from the central area a set of olivine microphenocrysts from the 5th April 2007 and of phenocrysts from the 26th October 2015 eruptions. These products are representative of the largest recent eruptions of PdF and were emitted during the most intense phases of activity (Di Muro et al., 2014; Coppola et al., 2017).

Investigated olivine and clinopyroxenes crystals in enclaves from the NWRZ peripheral area are dunites (DUNCHI-1/2/3), wehrlites (WERCHI-1/2) and harzburgite (ILR84-4) from Piton Chisny (~381 yrs BP) and Piton du Pas des Sables cone (>5 kyrs). Enclaves from the central area are dunites from the April 2007 eruption, with

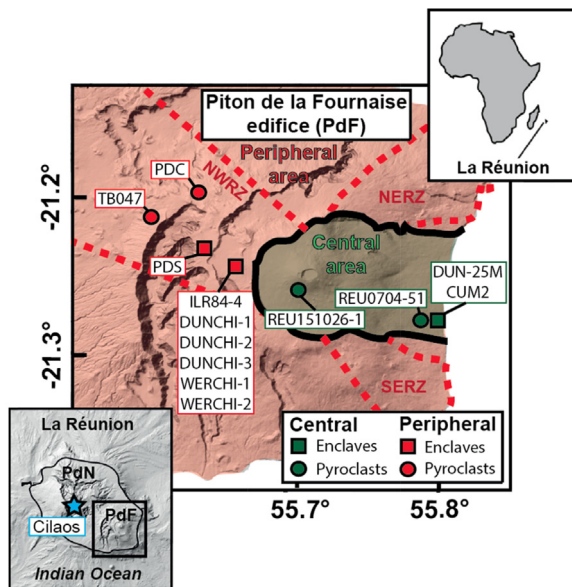


Fig. 1. Location of sampled sites (PdF for Piton de la Fournaise and PdN for Piton des Neiges) and of Cilaos free gases. TB047: Trous Blancs lapilli. PDC: Piton de Caille lapilli. PDS: dunite from Piton du Pas des Sables. ILR84-4: harzburgite from Piton Chisny. DUNCHI-1/2/3: dunites from Piton Chisny. WERCHI-1/2/3: wehrlites from Piton Chisny. CUM2: subaerial dunite sample from the April 2007 eruption. DUN-25M: submarine dunite sample (25 m bsl) from the April 2007 eruption. REU0704-51: 5th April 2007 lapilli. REU151026-1: 26th October 2015 scoria.

one subaerial (CUM2) and one submarine sample (DUN-25M; 25 m bsl).

The complete description of the analytical procedures used in this study (instrumentations, methodology, calculations, uncertainty and accuracy) is reported as Supplemental Information (Appendix A). The procedure of sample preparation and analysis for noble gases, TGC and carbon isotopes is also described in [Martelli et al. \(2014\)](#), [Rizzo et al. \(2015\)](#), [Gennaro et al. \(2017\)](#), and [Robidoux et al. \(2017\)](#). Below, we summarize the different methods used in this study to perform both the petrological and geochemical analyses.

Olivine and clinopyroxene crystals were handpicked from 13 samples previously fragmented and sieved in the size fractions of 0.5 and 1 mm.

Fluid inclusions (FIs) were analyzed for their composition with a Horiba T64000 Raman spectrometer manufactured by Jobin-Yvon with a 1800 grooves/mm gratings at IPGP (Paris, France). An argon ion laser at 488 nm wavelength was used at room temperature. Detailed descriptions of the composition of each fluid inclusion and of the procedure used for barometric calculations are reported in Supplemental Information (Appendices A and B).

Electron microprobe analyses on minerals chemistry were performed by using a CAMECA SX5 at Camparis (Paris, France). Further details are reported in Supplemental Information (Appendix A).

In total, 29 measurements have been carried out for noble gases and Total Gas Content (TGC), the latter being almost only CO₂ (“noble gases protocol”; more details are given in the text). These analyses were performed at the noble gas isotope laboratory of INGV, Sezione di Palermo (Italy). Helium (³He and ⁴He) isotopes were measured separately by two different split-flight-tube mass spectrometers (Helix SFT-Thermo). Argon isotopes (³⁶Ar, ³⁸Ar, and ⁴⁰Ar) were analyzed by a multicollector mass spectrometer (GVI Argus). The moles of TGC of the fluids trapped in the crystals were quantified by measuring the total pressure

of gas (by an IONIVAC Transmitters ITR 90) released during crushing in a known volume of the system.

A total of 30 measurements of the abundance and isotope composition of CO₂ were performed (“^δ¹³C protocol”). The moles of CO₂ released from crushing and purified from other components with cryogenic technique were quantified by measuring the pressure of gas within a section of a glass line with a known volume (~16 cm³) using a 626B Baratron[®] Absolute Capacitance Manometer made by MKS. The determination of ¹³C/¹²C (CO₂) isotope ratio (expressed as ^δ¹³C vs V-PDB) has been carried out by Thermo (Finnigan) Delta Plus XP CF-IRMS connected to a Trace GC gas chromatograph and a Thermo (Finnigan) GC/C III interface installed at the INGV-Palermo Stable Isotope Laboratory.

4. RESULTS

4.1. Petrology

4.1.1. Mineral chemistry in transitional lavas and enclaves

In samples from the central area, the composition of olivines (core and rims) from both pyroclasts and enclaves is consistent with the most magnesian mode (Fo₈₄) found in recent PdF central products ([Fig. 2](#); [Boivin and Bachelery, 2009](#); [Di Muro et al., 2016](#)). We find that olivines in peripheral pyroclasts are more magnesian (Fo_{85.5–86}) than central ones, consistently with previous findings ([Bureau et al., 1998](#)). The composition of olivines in peripheral enclaves spans a range from Fo₈₄ (DUNCHI-2) to Fo_{86.5} (DUNCHI-1) consistent with previous results on similar samples ([Brugier, 2016](#)).

Calcium contents in olivines cover a large range, from low-Ca olivines (CaO < 0.1 wt%) to high-Ca olivines (>0.4 wt%). Highest calcium contents are found in olivines of central pyroclasts and enclaves (0.38–0.41 wt%) and represent the highest Ca contents identified in PdF olivines

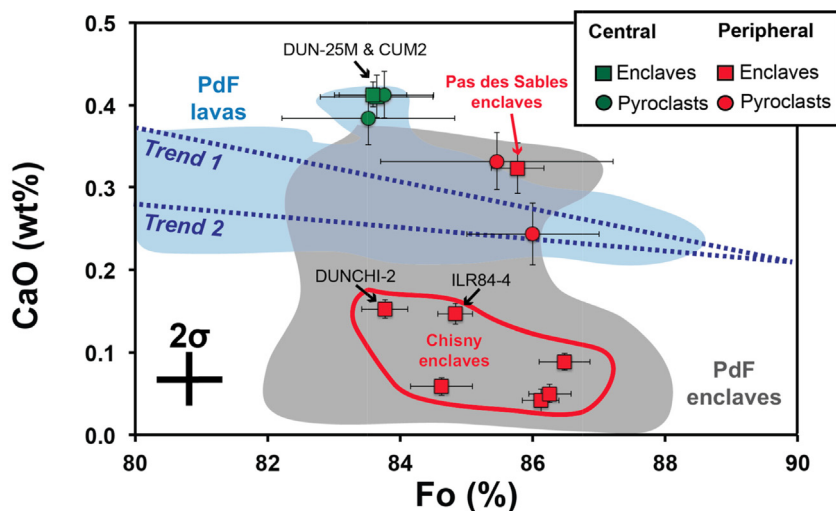


Fig. 2. Forsterite (mol%) vs. CaO content (wt%) in olivines hosting the fluid inclusions described in this study. Filled blue and grey fields represent published compositions of olivines from PdF lavas and enclaves, respectively ([Di Muro et al., 2015, 2016](#); [Brugier, 2016](#)). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

(Fig. 2). Olivines from peripheral pyroclasts and from the Piton des Sables dunite (PDS) have a lower CaO content with respect to central ones and fall close to the two trends (Trend 1 - high Ca and Trend 2 - low Ca on Fig. 2) that were previously recognized in PdF lavas (Di Muro et al., 2014). On the contrary, the main particularity of our dataset is that all olivines from Chisny enclaves present a strong depletion in calcium (<0.15 wt%) irrespective to their forsterite contents (from $F_{0.84.5}$ to $F_{0.86.5}$). Their composition falls in the lowest range of previously reported analyses (Fig. 2; Brugier, 2016).

4.1.2. Fluid inclusions in olivine crystals

Optically-resolvable FIs in olivine crystals studied here cover a large size range, from a few micrometers or less up to 100 μm in DUN-25M (Fig. 3). In olivines from both central and peripheral pyroclasts, their diameter rarely exceeds 1–2 μm (Supplemental Information: Fig. C1 in Appendix C). Conversely, a larger average size is observed in olivines from enclaves, mainly varying between 1 and 2 μm and 20 μm (up to 100 μm in DUN-25M). Following the nomenclature of Roedder (1984), we identify primary FIs (Fig. 3a) in olivines from peripheral and central pyroclasts (Table 1; REU151026-1a; TB047) and more rarely in olivines from enclaves (DUN-25M). Most of the FIs in olivine crystals from both central and peripheral enclaves are secondary FIs (multiple small FIs distributed along healed fractures that crosscut the entire crystal; Fig. 3b) and are sometimes associated with primary FIs (Table 1; PDS; WERCHI-1). Interestingly, in crystals hosting both types of FIs (e.g. PDS; WERCHI-1), neither compositional nor CO_2 densities differences were observed among small (<5 μm) FIs, independently of their primary or secondary origin.

No empty FIs were detected during Raman analysis. Only FIs smaller than 5 μm having a well-rounded shape without visible daughter minerals have been analysed following the recommendations of Frezzotti and Touret (2014). Large FIs (>5 μm) have often an irregular shape with optical evidences of daughter minerals which suggest the occurrence of post-trapping modification processes (Wanamaker and Evans, 1989; Frezzotti and Touret, 2014).

For each analysed FI, the low frequency region (<1500 cm^{-1}) of Raman spectra (200–4000 cm^{-1} ; Supplemental Information: Appendix B) shows the occurrence of bands related to (1) the main vibrational frequencies of the host crystal (i.e. olivine here; green arrows in Fig. 3) and, (2) the Fermi dyad resonance lines of gaseous CO_2 used for barometric purposes (1285 and 1388 cm^{-1}), together with the two shoulder peaks called “hot bands”.

In spite of the absence of visible daughter minerals, Raman spectra reveal the presence of microscopic solid phases in many FIs (1000–1500 cm^{-1} ; Supplemental Information: Appendix B):

- (1) Narrow bands (1083–1090 cm^{-1}), which can be attributed to Mg-Ca carbonate peaks and possibly to magnesite (MgCO_3 ; 1093 cm^{-1} ; Fig. 3b). These FIs mostly occur in crystals from Piton Chisny enclaves and TB047 peripheral pyroclasts. In other samples, we frequently observe weak narrow bands

at 1042–1044 cm^{-1} and 1057 cm^{-1} that are tentatively attributed to other C-bearing phases as nahcolite (NaHCO_3) and/or variably hydrous carbonates (Ruff spectral database),

- (2) Narrow bands in the range 1013–1038 cm^{-1} are attributed to sulfates. They were detected in FIs hosted in crystals of CUM2, PDS (Fig. 3a) and peripheral clinopyroxene-bearing enclaves (WERCHI-2). A narrow peak at 1150 cm^{-1} attributed to gaseous SO_2 was only observed in FIs of DUN-25M (Fig. 3c).

In the high frequency region (>1500 cm^{-1}), we identified three groups of spectra with either (1) a flat high frequency region (Fig. 3a; PDS); (2) a broad asymmetric band at 3430–3463 cm^{-1} and a weaker band at 3189–3192 cm^{-1} (Fig. 3b; DUNCHI-1 and essentially in Piton Chisny enclaves) that can be attributed to an aqueous phase ($\text{H}_2\text{O} \pm \text{CO}_2 \pm \text{salts}$) and; (3) a flat high frequency region with several weak narrow peaks (3159, 3222 and 3466 cm^{-1} for the most frequent) possibly related to an hydrous crystalline phase. Peaks related to interference of epoxy were clearly identified in a few samples (DUN-25M, CUM2, DUNCHI-1; Fig. 3c). Even if no narrow peak linked to water vapor (3640 cm^{-1}) was found in any sample, we cannot exclude that low amount of water vapor might occur in the fluid inclusions.

4.2. Geochemistry of fluid inclusions

4.2.1. He-Ar- CO_2 systematics

Helium, argon, carbon dioxide abundances and Total Gas Content (TGC) obtained by crystal crushing are given in Table 2. Good reproducibility is observed between replicates. Here we describe the general behavior of gas abundance between central and peripheral products and between pyroclasts and enclaves.

Helium abundance, here designated as ^4He , varies from $3.2 \times 10^{-13} \text{ mol g}^{-1}$ (ILR84-4) to $2.9 \times 10^{-11} \text{ mol g}^{-1}$ (REU151026-1) (Fig. 4). Pyroclasts and enclaves (CUM2) erupted in the central area show the lowest abundances (< $9 \times 10^{-13} \text{ mol g}^{-1}$). Pyroclasts emitted in the peripheral area on the volcano flanks are He-richer (up to $5 \times 10^{-12} \text{ mol g}^{-1}$) and the highest contents are measured in enclaves from the peripheral area (> $4 \times 10^{-12} \text{ mol g}^{-1}$).

The $^{40}\text{Ar}/^{36}\text{Ar}$ varies from 322 to 5226, which for most of the samples is sensibly higher than the atmospheric signature ($^{40}\text{Ar}/^{36}\text{Ar} = 295.5$). However, a correction from atmospheric contamination is necessary in order to obtain the so-called magmatic-derived $^{40}\text{Ar}^*$. This has been calculated, accordingly with the following formula that assumes that all the measured ^{36}Ar is of atmospheric origin:

$$^{40}\text{Ar}^* = ^{40}\text{Ar}_m - \left(\left(\frac{^{40}\text{Ar}}{^{36}\text{Ar}} \right)_{\text{air}} \times ^{36}\text{Ar}_m \right)$$

where $^{40}\text{Ar}^*$ represents the corrected ^{40}Ar , and the “m” subscript indicates “measured”. The recalculated $^{40}\text{Ar}^*$ ranges from $9 \times 10^{-14} \text{ mol g}^{-1}$ (REU151026-1) to $2 \times 10^{-11} \text{ mol g}^{-1}$ (ILR84-4; harzburgite). Like the ^4He abundance, $^{40}\text{Ar}^*$ contents generally increase from pyroclasts and enclaves

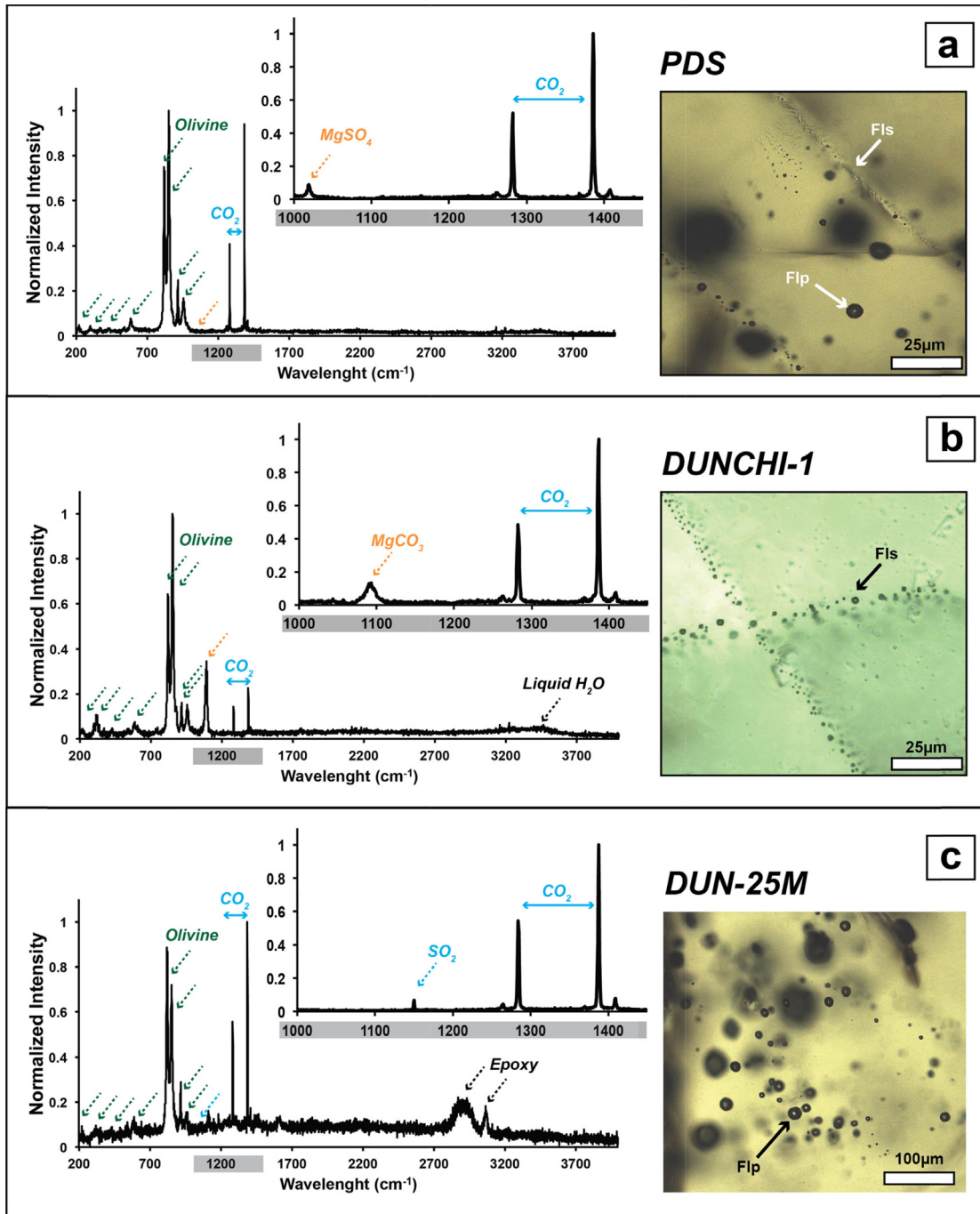


Fig. 3. Microphotographs and normalized Raman spectra (200 – 4000 cm^{-1}) of fluid inclusions. (a) Primary and secondary fluid inclusions in olivines of PDS. (b) Trails of small secondary fluid inclusions in olivines from DUNCHI-1. (c) Large primary fluid inclusions in olivines of DUN-25M. FIp: primary fluid inclusion. FIs: secondary fluid inclusion. Green arrows: Raman bands of the host crystal (olivine). Orange arrows: Raman bands of secondary mineral phases within fluid inclusions (mainly carbonates and sulfates). Light blue arrows: Raman bands of gaseous species. Black arrows: Raman bands of liquid H_2O and epoxy in the high frequency region ($>1500 \text{ cm}^{-1}$). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

of the central area ($<7.6 \times 10^{-12} \text{ mol g}^{-1}$) to pyroclasts emitted on the volcano flanks ($\sim 2 \times 10^{-12} \text{ mol g}^{-1}$) (Fig. 4a). Highest $^{40}\text{Ar}^*$ abundances are found in peripheral enclaves ($>4 \times 10^{-12} \text{ mol g}^{-1}$). The $^4\text{He}/^{40}\text{Ar}^*$ varies in the range between 0.7 and 8.5, which is slightly wider than that

of the typical production ratio of the mantle ($^4\text{He}/^{40}\text{Ar}^* = 1\text{--}5$; e.g., Ozima and Podosek, 1983; Marty, 2012).

CO_2 concentration (from TGC) ranges from $9 \times 10^{-10} \text{ mol g}^{-1}$ (CUM2) to $8 \times 10^{-7} \text{ mol g}^{-1}$ (ILR84-4) (Fig. 4b). The lowest contents are found in pyroclasts and rare

Table 1
Petrography, composition and Raman Fermi dyad shift of fluid inclusions with calculated pressures from the CO₂-density equations of Lamadrid et al. (2016). Composition of host olivines is reported (Fo: forsterite content).

Sample	Location	Type	Name	Petrography		Raman analysis		ΔFermiCO ₂ (cm ⁻¹)				Pressure estimations (MPa)				Olivine composition			
				Primary FIs	Secondary FIs	Number	Number	Min	Max	Avg	Min	Max	Avg	σ	Number	Fo (%)	σ	CaO (wt%)	σ
Central	Enclaves	DUN-25M	CUM2	X	X	10	10	103.21	103.32	103.27	89	108	100	6	9	83.6	0.5	0.41	0.01
								103.07	103.14	103.11	67	76	72	3	10	83.6	0.8	0.41	0.03
								102.81	102.91	102.86	26	42	34	5	9	83.5	1.3	0.38	0.03
Peripheral	Enclaves	ILR84-4	PDS	X	X	9	9	103.39	103.39	103.39	167	167	167	0	9	83.8	0.7	0.41	0.03
								104.78	104.32	104.15	183	345	290	41	10	84.8	0.3	0.15	0.01
								104.16	104.41	104.25	294	391	323	11	10	85.8	0.4	0.32	0.03
Peripheral	Enclaves	DUNCHI-1	DUNCHI-1	X	X	10	10	103.09	104.08	103.72	71	267	183	50	10	86.5	0.4	0.09	0.01
								103.23	104.14	103.96	94	285	237	50	6	83.8	0.3	0.15	0.01
								102.79	103.22	103.03	22	89	60	17	8	84.6	0.5	0.06	0.01
								102.81	103.52	103.28	26	151	105	42	6	86.1	0.3	0.04	0.01
								102.91	103.38	103.20	42	118	89	24	8	86.3	0.3	0.05	0.01
								104.15	104.39	104.27	298	397	349	35	48	85.5	1.8	0.33	0.03
Peripheral	Pyroclasts	WERCHI-2	WERCHI-2	X	X	5	5	104.15	104.39	104.27	320	570	450	50	86.0	1.0	0.24	0.04	

Microthermometry analysis

enclaves from the central area ($<1 \times 10^{-9}$ mol g⁻¹). Intermediate abundances are found in peripheral pyroclasts ($2\text{--}3 \times 10^{-8}$ mol g⁻¹) whereas the highest values are set in peripheral enclaves ($>3 \times 10^{-8}$ mol g⁻¹). As reflected by ⁴He, ⁴⁰Ar* and CO₂ abundances, the TGC released by crystal crushing during noble gases analysis increases from central to peripheral products (Table 2). The only exception is given by DUN-25M central enclave, a fast quenched sample recovered from a depth of 25 m below sea level that has a gas abundance (4×10^{-7} mol g⁻¹) in the range of peripheral products, close to the maximum value measured in our dataset (8×10^{-7} mol g⁻¹; ILR84-4) (Table 2; Fig. 6c).

In our samples, ⁴He vs. ⁴⁰Ar* and ⁴He vs. CO₂ are well fitted by power functions ($r^2 = 0.95$ and 0.96 , respectively; Fig. 4). With the exception of DUN-25M, it confirms the gas depletion in central products with respect to peripheral ones. It is worth noting that DUNCHI-1, DUNCHI-2 and DUN-25M are the only samples falling out of the 95%-confidence interval defined by the power functions. We note that previous analyses performed on Piton Chisny peripheral enclaves (Trull et al., 1993) present a maximum ⁴He content (3.0×10^{-11} mol g⁻¹; black squares on Fig. 4b) that is similar to the one we document in our study (2.9×10^{-11} mol g⁻¹). Meanwhile, CO₂ content strongly differs between both studies. For instance, for ILR84-4, we measure a content of 7.8×10^{-7} mol g⁻¹ that is lower than the value of 3.9×10^{-6} mol g⁻¹ reported by Trull et al. (1993). We argue that the observed discrepancy in the CO₂ estimations could be due to the distinct analytical protocols (Kendrick and Phillips, 2009). Actually, ⁴He contents reported by Trull et al. (1993) refer to crystal crushing measurements performed by Staudacher et al. (1990) whereas CO₂ contents were determined by step-heating procedure.

4.2.2. Carbon isotope composition of CO₂

Single (or individual) measurement of CO₂ concentrations and δ¹³C values obtained by the replicates for each sample are reported in Table 3. Average value and standard deviation data are also reported in Table 2. A good reproducibility is observed between CO₂ estimations from the “δ¹³C protocol” and the “noble gases protocol” (Supplemental Information: Fig. C2 in Appendix C). Importantly, a good reproducibility is also observed between replicates regarding both CO₂ concentrations and δ¹³C values.

The δ¹³C values span a broad range from -11.3‰ in pyroxene from WERCHI-1 to -1.8‰ in olivine from DUN-25M (Fig. 7). This range of values is comparable to that reported by Trull et al. (1993) ($-10.2\text{‰} < \delta^{13}\text{C} < -3.7\text{‰}$). These authors found that the lowest isotopic values are only observed in clinopyroxene from Chisny wehrlitic enclaves (-10.2‰), while olivines have less negative δ¹³C in agreement with our measurements. We note that the measurements we made in clinopyroxene from WERCHI-2 (-3.7‰) does not show a ¹³C depletion like in WERCHI-1. These values while fall within the range of δ¹³C values found in olivines (-2.6‰ to -4.6‰) reported for other enclaves in this study and by Trull et al. (1993). Lower values are obtained for peripheral pyroclasts (-5‰ in TB047 and -6.4‰ in PDC).

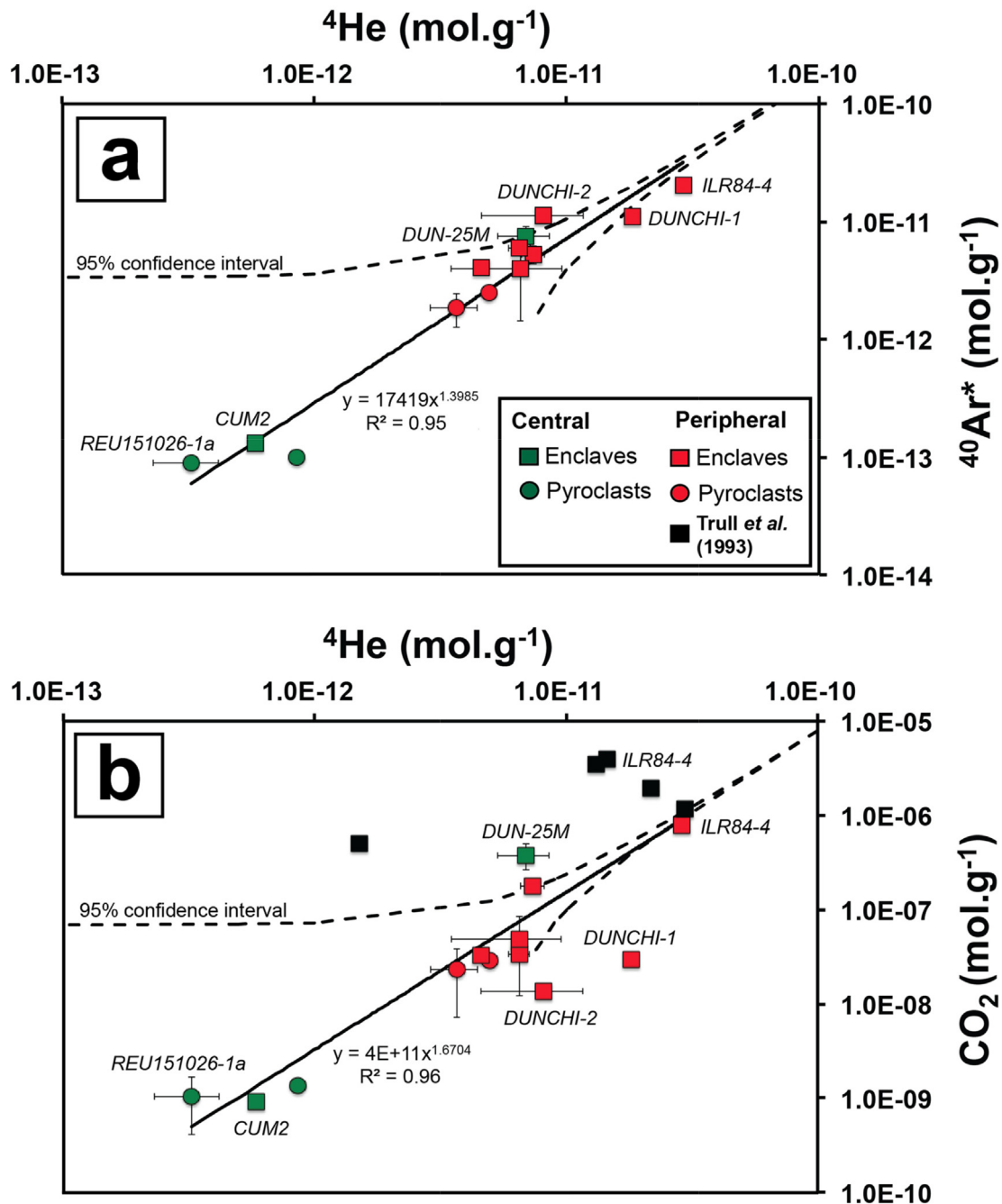


Fig. 4. Gas contents released by crystal crushing and best fitting functions (the 95% confidence intervals are reported by dashed lines and delimit the field where 95% of the measured values are expected to fall with respect to the best fitting functions). (a) ^4He vs. $^{40}\text{Ar}^*$. (b) ^4He vs. CO_2 . CO_2 contents from Trull *et al.* (1993) (black squares) measurements were obtained by a step-heating procedure whereas He contents refer to crushing measurements on the same samples by Staudacher *et al.* (1990).

Unfortunately, no measurements were possible in central pyroclasts due to the very low amount of CO_2 in these products.

ILR84-4 harzburgite ($\delta^{13}\text{C}$: -3.2‰) was also analysed by Trull *et al.* (1993) that found a similar ratio ($\delta^{13}\text{C}$: -3.7‰), the difference falling in the range including both the uncertainty of the analytical procedure and the natural

variability of the samples. If CO_2 contents measured by step-heating (Trull *et al.*, 1993) are higher than those measured in this study by crystal crushing, $\delta^{13}\text{C}$ values are independent on the used protocol. Consequently, in order to avoid any bias in the following parts, we will take into consideration only $\delta^{13}\text{C}$ values (and not the CO_2 contents) documented by these authors.

Table 3

CO₂ contents and $\delta^{13}\text{C}$ from mineral crushing. Average and standard deviation from replicates are reported. Crushed mineral (ol: olivine; px: clinopyroxene) between parenthesis).

Sample			Replicates			Average			
Location	Type	Name	Label	CO ₂ (mol g ⁻¹)	$\delta^{13}\text{C}$ (‰)	CO ₂ (mol g ⁻¹)	σ	$\delta^{13}\text{C}$ (‰)	σ
Central	Enclaves	DUN-25M (ol)	1	4.52E-07	-3.3	4.04E-07	1.03E-07	-2.6	0.5
			2	4.21E-07	-2.9				
			3	5.78E-07	-2.7				
			4	3.09E-07	-1.8				
			5	4.06E-07	-2.7				
			6	2.55E-07	-2.0				
Peripheral	Enclaves	ILR84-4 (ol)	1	4.67E-07	-3.1	5.96E-07	1.24E-07	-3.2	0.1
			2	5.74E-07	-3.2				
			3	5.14E-07	-3.3				
			4	5.95E-07	-3.3				
			5	8.27E-07	-3.3				
		PDS (ol)	1	2.14E-07	-3.4	1.47E-07	4.99E-08	-3.1	0.4
			2	9.35E-08	-2.5				
			3	1.35E-07	-3.5				
		DUNCHI-1 (ol)	1	2.95E-08	-3.6	1.29E-08	3.37E-09	-4.6	0.1
			1	1.62E-08	-4.5				
		DUNCHI-2 (ol)	1	1.62E-08	-4.5	1.29E-08	3.37E-09	-4.6	0.1
			2	9.49E-09	-4.7				
		DUNCHI-3 (ol)	1	3.34E-08	-3.2	3.19E-08	3.26E-09	-3.8	0.4
			2	2.74E-08	-4.1				
			3	3.50E-08	-4.0				
	WERCHI-1 (px)	1	5.44E-08	-11.3	6.41E-08		-3.7	0.6	
		1	6.41E-08	-4.3					
	WERCHI-2 (px)	1	6.41E-08	-4.3	6.41E-08		-3.7	0.6	
		2	6.41E-08	-3.2					
	Pyroclasts	TB047 (ol)	1	2.57E-08	-5.0	9.64E-08	3.84E-08	-6.4	0.5
			1	1.31E-07	-5.5				
PDC (ol)		2	6.10E-08	-6.0					
		3	1.62E-07	-6.8					
		4	6.20E-08	-6.4					
		5	9.58E-08	-6.6					
		6	6.66E-08	-7.2					

5. DISCUSSION

5.1. Inferences on magma ponding zones

5.1.1. Crystallization depth

Entrapment pressures of FIs were estimated from their CO₂ density (Table 1; see Methods in Supplemental Information). Since numerous processes can affect the initial CO₂ density of FIs and thus modify their pristine pressure, it is generally assumed that the estimated pressure corresponds to the minimum entrapment pressure and depth (Viti and Frezzotti, 2000). In volcanic contexts, the main processes of post-trapping modification and leading to major underestimation of the original density in FIs are: (1) the partial or total fluid leakage, (2) the formation of volatile bearing daughter minerals like carbonates and, (3) volumetric resetting by crystal brittle (decrepitation) or plastic failure (stretching) (Wanamaker and Evans, 1989; Frezzotti and Touret, 2014). These phenomena are mainly dependent on the strength of the mineral host (Wanamaker and Evans, 1989) and on the ascent history (see Hansteen and Klügel (2008) for a complete review). In Supplemental Information (Appendix D), we evaluate

their influence on our pressure estimations. More specifically, we point out that (1) fluid leakage is quite limited in our samples set and (2) late-stage carbonates formation only plays a minor role on the estimated densities, close to the uncertainty related to the analytical procedure (Fig. 5). We infer that (3) volumetric resetting processes could be the main factor leading to an underestimation of the initial trapping pressure, and have mainly operated in crystals from enclaves (see below).

Meanwhile, in our study (Table 1), we show that central products (pyroclasts: REU0704-51, REU151026-1; dunite enclaves: CUM2, DUN-25M) record low crustal pressures (<167 MPa and centered at 80–120 MPa; Fig. 5c; Table 1) broadly consistent with previous data (Famin et al., 2009; Di Muro et al., 2016). FIs from central samples sometimes contain S-bearing phases as sulfates (CUM2), consistent with the exsolution of sulfur species at shallow pressure (<200 MPa) due to the high solubility of sulfur in PdF basaltic melts (Di Muro et al., 2016). Highest (mantle) pressures are recorded in peripheral pyroclasts (570 MPa and always >310 MPa; Fig. 5c). This is in accordance with the high pressures previously documented for peripheral products emitted within the NWRZ (Bureau et al., 1998).

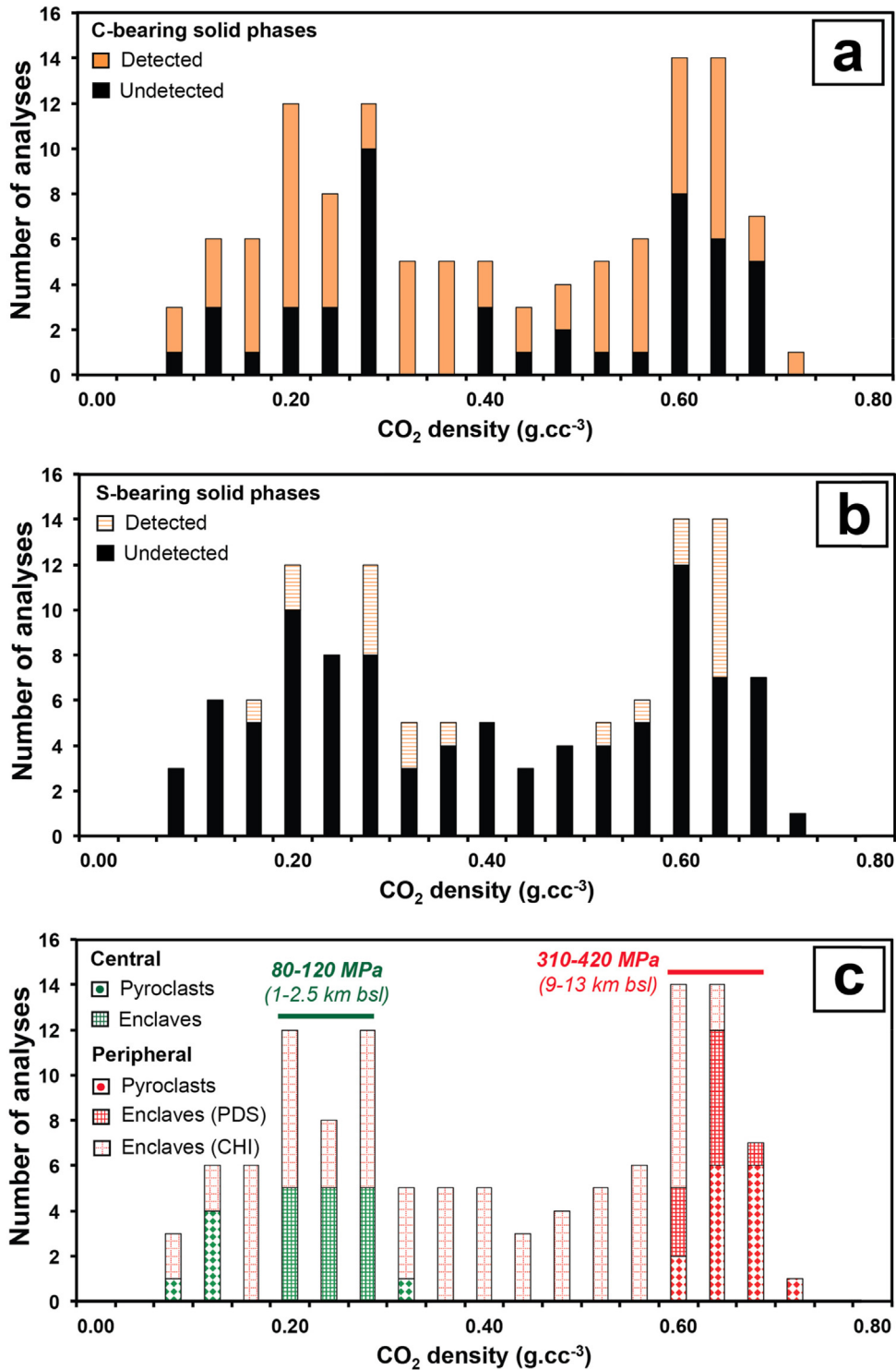


Fig. 5. Frequency distribution of CO₂ densities (see text for explanations) in fluid inclusions according to (a) the presence of carbon-bearing solid phases (mainly Mg-carbonates), (b) the presence of S-bearing solid phases (mainly Mg-sulphates) and, (c) the type of sample (pyroclasts or enclaves) and its location on the volcano (central or peripheral area).

Peripheral enclaves (dunites: DUNCHI-1/2/3, PDS; wehrlites: WERCHI-1/2; harzburgite: ILR84-4) span a large range of pressures. On one hand, PDS FIs record pressures consistent with those of peripheral pyroclasts

(310–420 MPa; Fig. 5c). On the other hand, FIs from Piton Chisny enclaves span a large domain of pressure, ranging from mantle pressures (>300 MPa; ILR84-4) to crustal pressures (<300 MPa and mostly <151 MPa), and thus

overlapping the density values recorded in central products. Our large dataset allows us to define two distinct barometric modes: (1) at 80–120 MPa that is within the volcano edifice and close to the sea level. This mode is best recorded by central products and some FIs in Piton Chisny enclaves; (2) at 310–420 MPa, that is the depth of the mantle-crust underplating layer, essentially recorded by peripheral pyroclasts, PDS enclave and part of the Piton Chisny enclaves.

Interestingly, in olivines from pyroclasts, as pressure of hosted FIs decreases, the forsterite content decreases (Fig. 6a), while calcium content increases (Fig. 6b). This positive correlation between olivine Mg content and pressure as well the calcium enrichment mirrors the PdF melt differentiation (dashed line in Fig. 6a and b; Di Muro et al., 2015, 2016; Boudoire, 2017). Such a compositional evolution is mainly controlled by early olivine fractionation followed by clinopyroxene (Famin et al., 2009). Central (CUM2, DUN-25M) and part of peripheral enclaves (PDS from Pas des Sables; ILR84-4 harzburgite and DUNCHI-2 from Chisny) fall on the expected trend of evolution of olivine composition with pressure. Olivines from the Chisny enclaves hosting FIs with a large range of densities have a very low Ca content (<0.15 wt% and mostly <0.1 wt%) which does not correlate with the olivine forsterite content. This feature was previously attributed to a diffusive re-equilibration of Ca either in hot melts or during long residence time in the PdF plumbing system. It may thus reflect the long-lasting and potentially complex thermal history of Chisny enclaves (Brugier, 2016).

5.1.2. Gas contents released by crystal crushing

Interestingly, the TGC from crushed crystals from pyroclasts decreases together with pressure (dashed line in Fig. 6c). This means that for a given mineral phase, gas-rich samples are preferentially trapped at greater pressures. Conversely, most of the enclaves differ from this trend, being gas-rich in spite of low recorded densities. We link this behavior to a preferential volumetric resetting of FIs in olivines from enclaves during multistage magma ascent and ponding, registering the pressure range crossed by the volcanic conduit (Supplemental Information: Appendix D). As a matter of fact, TGC in these enclaves (<300 MPa and mostly <151 MPa) are similar to those in peripheral pyroclasts (>300 MPa), suggesting a deeper storage conditions than that recorded by FI density. Partial resetting of FI densities in Chisny enclaves is also in agreement with the inferred occurrence of a dunitic-wehrlitic cumulative zone in the depth range of the underplating layer (>300 MPa; Charvis et al., 1999; Fontaine et al., 2015; Brugier, 2016; Boudoire, 2017). The highest TGC measured in the dunitic enclave from Piton Pas des Sables (PDS) and from the harzburgitic enclave from Chisny (ILR84-4) could record an even deeper origin within the upper lithospheric mantle (Fig. 6c). We point out here that crystals from enclaves are more likely to undergo diffusive re-equilibration processes (e.g., low Ca content, volumetric resetting of fluid inclusions) and favor the use of pressure estimations from fluid inclusions in crystals from lavas and pyroclasts in the following discussion.

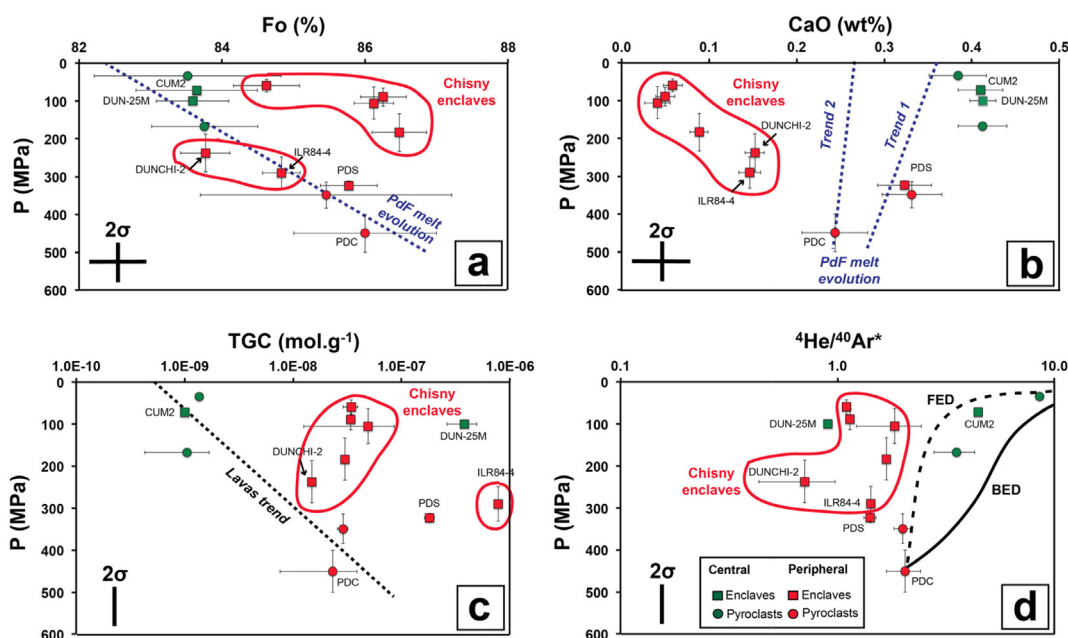


Fig. 6. Average host olivine composition (a, b) and total gas contents (TGC) or elements ratios (c, d) vs. average pressure estimations from fluid inclusion barometry. (a) Predicted olivine composition in equilibrium with PdF melt from Boudoire (2017) (b) Calcium content (wt%) of olivines and trends from Fig. 3. (d) Fractional Equilibrium Degassing (FED) and Batch Equilibrium Degassing (BED) paths were modeled using gases solubility from Piton de Caille (PDC) melt inclusions (see text for explanation and Table 4; Iacono-Marziano et al., 2010; Duan, 2014). For each sample, the error bars represent the variability between replicates. Analytical and methodological uncertainty are also represented as 2σ for pressure.

If absolute gas contents may be influenced by the frequency of FIs trapped within each crystal, we expect that the ratios between noble gases are not affected. $^4\text{He}/^{40}\text{Ar}^*$ is a tracer of magmatic degassing processes due to low solubility of Ar in silicate melts with respect to He (Jambon et al., 1986; Burnard, 2004; Burnard et al., 2003, 2014; Aubaud et al., 2004; Yamamoto and Burnard, 2005; Paonita and Martelli, 2007; Iacono-Marziano et al., 2010). $^4\text{He}/^{40}\text{Ar}^*$ higher than 3.5 have been found in central pyroclasts and enclaves (CUM2), consistent with the shallow origin of these products (Fig. 6d). Conversely lower ratios (≈ 2) are found in peripheral (i.e., deep) pyroclasts. In order to test this pressure-dependence of the $^4\text{He}/^{40}\text{Ar}^*$ in our products, we model batch and fractional equilibrium degassing trends (BED and FED in Fig. 6d), corresponding to closed- and open-system conditions, respectively. We start from PDC melt composition (Table 4; see explanation in the following parts), focusing on the compositional evolution of the vapor phase between 450 MPa and 5 MPa (Duan, 2014). This approach suggests that $^4\text{He}/^{40}\text{Ar}^*$ variability between deep peripheral and shallow central products is mainly modulated by FED. Both melts degassing and differentiation are thus strongly dependent on pressure at PdF. Low $^4\text{He}/^{40}\text{Ar}^*$ recorded by peripheral enclaves are also consistent with our previous considerations suggesting a deeper origin for these enclaves. However, these ratios are systematically lower than the rest of the dataset (< 1.9). Potential effects accounting for this difference will be considered and discussed in the following sections.

Nevertheless, DUN-25M is the only sample in our dataset that is not consistent with these results. In this sample, olivine composition, density of FIs, and the presence of solid or solute phases in FIs fully overlap the ranges observed in other central products (Fig. 6a and b). The TGC and $^4\text{He}/^{40}\text{Ar}^*$ are instead more in accordance with a deeper origin of this fast quenched enclave with respect to typical central products (Fig. 6c and d). We also point out that this sample has the largest size range of FIs described in this study (up to 100 μm) with an uncommon proportion of very large FIs (Fig. 2a) and it is the only sample collected below sea level. Our data suggests a probable volumetric resetting of FIs, leading to a strong underestimation of the entrapment pressure. However, further investigations are required to explain the processes accounting for the peculiar signature of this sample.

5.2. Modeling of magma degassing

5.2.1. $\text{CO}_2 - \delta^{13}\text{C}$ systematics

Even if CO_2 densities in FIs can be affected by volumetric resetting, which leads to pressure underestimation, this process does not affect the absolute CO_2 contents trapped in FIs. Only CO_2 leakage or sequestration in solid and liquid phases can affect the gas content released by crushing. However, we did not identify empty FIs in this study, thus limiting the effect of leakage, and we observed that liquid phases were absent or have small volume with a limited role on the CO_2 sequestration. We have previously highlighted

Table 4

Starting conditions for degassing modeling. Model A: most primitive primary melt inclusion from Piton de Caille peripheral eruption (PDC; Bureau et al., 1998) and initial pressure of 570 MPa (maximum pressure from fluid inclusions barometry). Model B: most primitive melt inclusions from Trouis Blancs (TB047) peripheral eruption at 397 MPa (maximum pressure from fluid inclusions barometry). Model C: primary melt inclusion from August 2015 central eruption at 167 MPa. $^*\text{CO}_2$ estimation based on the pressure-composition-water dependent model of Duan (2014). Temperature estimated from the hydrous-composition dependent equation of Putirka (2008). Noble gases solubility calculated from Iacono-Marziano et al. (2010). Densities estimated from Lange and Carmichael (1990). The pressure and compositional range used for the modeling fully overlap the variability described by PdF melt.

		Model A IR98 5	Model B TB047-9-MI2	Model C A2015
Primary melt inclusions composition (wt%)	SiO_2	49.67	47.71	49.53
	TiO_2	2.69	2.61	2.71
	Al_2O_3	13.33	13.29	14.50
	FeO_T	10.03	10.62	10.03
	Fe_2O_3	1.19	1.27	1.32
	FeO	8.96	9.48	8.85
	MnO	0.16		0.14
	MgO	9.97	10.90	8.12
	CaO	10.93	10.84	11.65
	Na_2O	2.43	2.77	2.66
	P_2O_5	0.25	0.34	0.00
	K_2O	0.55	0.92	0.65
	H_2O	1.12	1.42	0.69
CO_2^*	0.35	0.24	0.08	
Pressure (MPa) & Temperature ($^\circ\text{C}$)	P	570	397	167
	T	1257	1281	1218
Solubility (ccSTP g^{-1} bar $^{-1}$)	^4He	5.23E-04	4.58E-04	4.81E-04
	CO_2	3.08E-04	3.08E-04	2.46E-04
	$^{40}\text{Ar}^*$	5.16E-05	4.46E-05	5.16E-05
Density (g cc^{-1})	d	2.72	2.69	2.68

that the TGC is more prone to reflect the original pressure of crystallization. Similar conclusion can be attained for the CO₂ content released by crushing, considering that CO₂ contributes to more than 95% of the TGC (Table 2).

To constrain degassing processes and extents, it is a common approach to evaluate how CO₂ concentration dissolved in glasses or trapped in vesicles varies respect to the δ¹³C signature (Macpherson and Matthey, 1994; Shaw et al., 2004; Aubaud et al., 2005, 2006; Barry et al., 2014). Here we investigate the evolution of CO₂ content and related carbon isotopic signature by assuming that the evolution of the CO₂ content trapped by fluid inclusions in crystals is positively correlated with the CO₂ content dissolved in the melt. This assumption is supported by the pressure-TGC and pressure-Fo correlations described by the crystals we analysed in pyroclasts (Fig. 6): the deeper (higher pressure) and more magnesian (high Fo) is the basaltic melt, the larger is the amount of CO₂ (e.g., TGC) released by crystal crushing (Fig. 6). Similarly, dissolved CO₂ content increases with increasing pressure and magnesium content in basaltic melts (Aubaud et al., 2004; Papale et al., 2006).

To model magma degassing and the isotopic composition of the vapour phase, we have taken into account the carbon isotope fractionation factor between vapour and melt ($\Delta = \delta^{13}C_{melt} - \delta^{13}C_{vapor}$) in order to modify the initial BED and FED degassing equations of Macpherson and Matthey (1994) available for the melt phase:

$$\delta^v = \delta^p - \Delta \times \left(\frac{C_{BED}^r}{C^p} \right) \text{ for batch equilibrium degassing model} \quad (1)$$

and

$$\delta^v = \delta^p + \Delta \times \left(\ln \left(\frac{C_{FED}^r}{C^p} \right) - 1 \right) \text{ for fractional equilibrium degassing model} \quad (2)$$

where BED and FED subscript refers to batch equilibrium degassing and fractional equilibrium degassing model, respectively, δ^v represent the isotope composition (δ¹³C) of the CO₂ in the vapor phase (i.e., trapped by FIs), C^r is the CO₂ contents from crushed crystals (here considered as a proxy of the CO₂ contents dissolved in melt), C^p and δ^p are the initial concentrations and isotope compositions of the exsolved phase of CO₂ that should be released by fluid inclusions during crystal crushing, and $\Delta = \delta^{13}C_{melt} - \delta^{13}C_{vapor}$.

A melt-vapor fractionation factor (Δ) of $+2.2 \pm 0.2$ and the uncertainty related to the analytical procedure were determined from Eq. (3) (see below) by calculating the slope of the best fit regression line through the analysis of the sample with the highest gas content (ILR84-4) (a) and PDC (b), which are best fitted by a FED equation (Barry et al., 2014):

$$\Delta = \frac{(\delta_b^v - \delta_a^v)}{\ln \left(\frac{C_b^r}{C_a^r} \right)} \quad (3)$$

The factor of melt-vapor fractionation we estimate falls in the range of experimental data and values obtained for other volcanoes (+1.8 to +4.5; Javoy et al., 1978; Matthey,

1991; Aubaud et al., 2005); moreover, our Δ is close to the value assumed at Mount Etna and Stromboli for similar modeling ($\Delta = +3$; Paonita et al., 2012; Gennaro et al., 2017).

Then, considering that one δ^v value may correspond to one C_{BED}^r and one C_{FED}^r , it is possible to combine the Eqs. (1) and (2) in order to obtain C^p :

$$\ln(C^p) - \frac{C_{BED}^r}{C^p} = \ln(C_{FED}^r) - 1 \quad (4)$$

Using two samples (ILR84-4 and PDS) following a BED trajectory (here based on PDS that is the only enclave sample that does not present re-equilibration effects, neither the presence of carbonates in fluid inclusions; Supplemental Information: Appendix B; Fig. 6), we finally obtain from (4):

$$C^p = \frac{(C_{BED}^{ILR84-4} - C_{BED}^{PDS})}{\ln \left(\frac{C_{FED}^{ILR84-4}}{C_{FED}^{PDS}} \right)} \quad (5)$$

where $C_{BED}^{ILR84-4, PDS}$ are the CO₂ contents measured by crystal crushing and $C_{FED}^{ILR84-4, PDS}$ the calculated values using the best fit regression line considered to represent the FED trajectory. Using the obtained values for C^p , Δ , $C_{BED}^{ILR84-4, PDS}$, $C_{FED}^{ILR84-4, PDS}$ and $\delta^{a,b}$, the Eqs. (1) and (2) lead to $\delta^p = -0.8 \pm 1.0\text{‰}$ considering the average and the uncertainty related to both the analytical procedure and the natural variability of the samples (see Barry et al. (2014)). Using the melt-vapor fractionation factor, it leads to an initial $\delta^{13}C_{melt} = -3.0 \pm 1.2\text{‰}$ at PdF. This estimation is fully consistent with values reported for other hotspots (-3.4 to -4‰ at Kilauea; Gerlach and Taylor, 1990; -3 to -4‰ for Society Islands; Aubaud et al., 2005; Deines, 2002) as well as for very little degassed popping rocks from mid-Atlantic ridge (-3.7‰ ; Javoy and Pineau, 1991; Michael and Graham, 2015).

Using these calculated parameters, our whole dataset is well-fitted by the range of BED and FED degassing paths modeled from the starting conditions defined above (Model 1; Fig. 7a). PDC products (peripheral pyroclasts) are closer to a FED path. Other data (mostly related to enclaves) lie between FED and BED trajectories. Commonly such a behavior is interpreted as related to different stages of degassing (Cartigny et al., 2001, Sarda and Moreira, 2002, Shaw et al., 2004). However, other processes may account for the δ¹³C variability as (1) source variability (Deines, 2002), (2) assimilation of a crustal component (Parks et al., 2013) and/or, (3) calcite precipitation (Ray et al., 2009; Dubacq et al., 2012; Paonita et al., 2012; Barry et al., 2014). At PdF, it has been inferred that source variability (1) and contamination (2) play only a minor role in the melt geochemistry and cannot account for the observed variability in bulk rock composition [see Vlastelic and Pietruzka (2016) for a review]. Nevertheless, we have also seen that carbonates are systematically present in our FIs (3). Carbon isotope fractionation due to calcite precipitation is highly temperature-dependent (Bottinga, 1969; Hoefs, 2015) inducing a ¹³C-enrichment in the residual vapour phase at temperature higher than 192 °C. Assuming that about 10% of the initial CO₂ could precipitate as carbonate (Bolder-Schrijver et al., 2000) we estimate

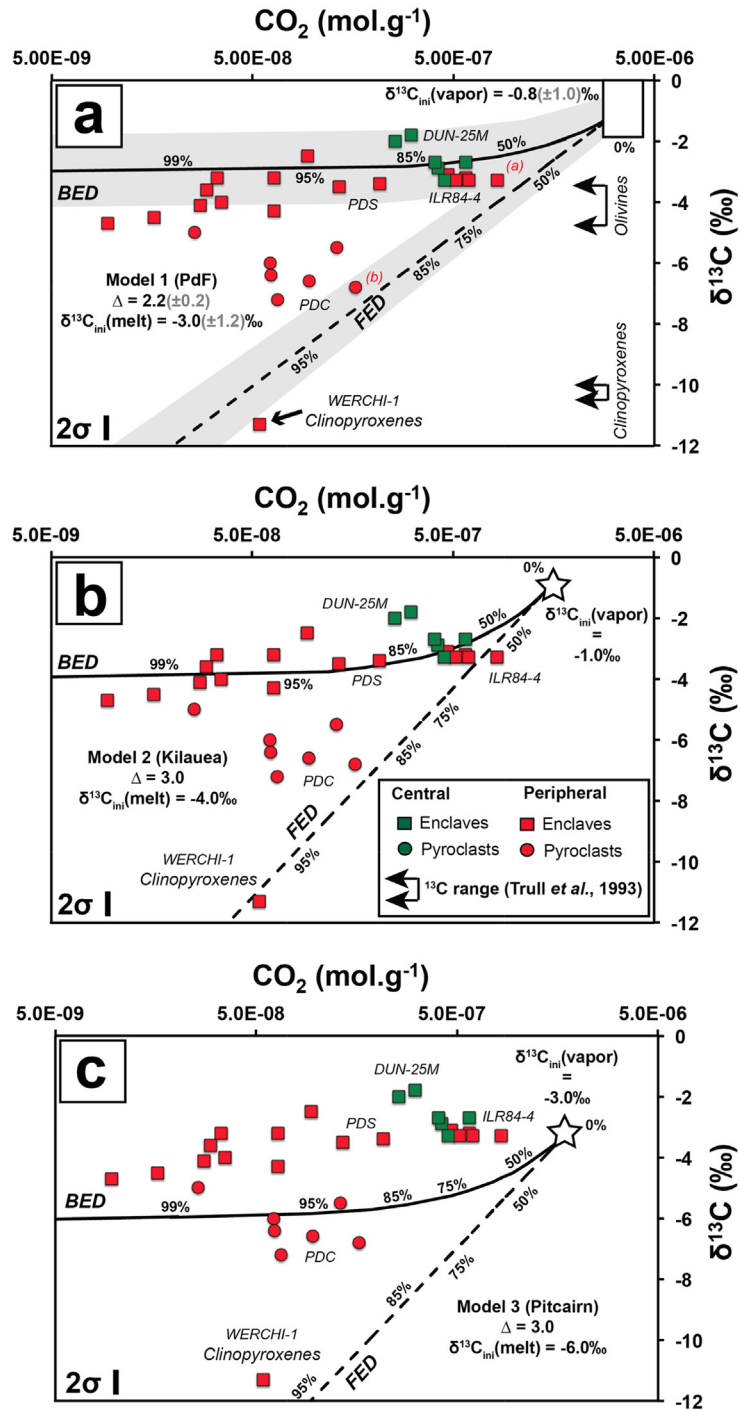


Fig. 7. CO_2 contents vs. $\delta^{13}\text{C}$ from crushed minerals. Batch Equilibrium Degassing (BED) and Fractional Equilibrium Degassing (FED) are modeled with three different starting conditions (a, b, c). The extent of degassing (0–50–75–85–95–99%) is reported along the curves. (a) Fractionation factor ($\Delta = 2.2 \pm 0.2$) and initial melt $\delta^{13}\text{C}$ ($= -3.0 \pm 1.2 \text{‰}$) computed from the best fit on PdF products and taking into account the uncertainties related to the analytical and natural variability (see text for explanation; Barry et al., 2014). Grey field highlights the expected range of BED and FED considering the uncertainty related to the starting conditions (Δ and $\delta^{13}\text{C}$). (b) Fractionation factor ($\Delta = 3$) corresponding to the average reported in the literature (see text for explanations) and to an initial melt $\delta^{13}\text{C}$ ($= -4 \text{‰}$) for Kilauea hotspot (Gerlach and Taylor, 1990). (c) Fractionation factor ($\Delta = 3$) corresponding to the average reported in the literature (see text for explanations) and to a primitive melt $\delta^{13}\text{C}$ ($= -6 \text{‰}$) for Pitcairn hotspot (Aubaud et al., 2006). $\delta^{13}\text{C}$ range from -4 to -6‰ is considered to overlap the mantle variability (Cartigny et al., 2001). Black arrows correspond to the $\delta^{13}\text{C}$ variability obtained by step-heating olivines and clinopyroxenes from Piton Chisny peripheral enclaves and determined by Trull et al. (1993).

a maximum variation from -1‰ to $+0.3\text{‰}$ (between 25 and 1000 °C; Hoefs, 2015) that is broadly overlapped by the uncertainty related to the method of FIs extraction and by the natural variability for a same sample (Table 3). Such process may be sufficient to explain the few high $\delta^{13}\text{C}$ values in DUN-25M (Fig. 7a), but cannot be considered as a major process influencing our dataset.

In order to compare our result with a more classical approach, we modeled BED and FED paths with parameters constrained for other well-known hotspots environments: (1) Model 2 (Fig. 7b) refers to Kilauea starting conditions with an initial $\delta^{13}\text{C}_{\text{melt}} = -4\text{‰}$ that is the lowest estimation made by Gerlach and Taylor (1990); (2) Model 3 (Fig. 7c) refers to Pitcairn starting conditions with an initial $\delta^{13}\text{C}_{\text{melt}} = -6\text{‰}$ (Aubaud et al., 2006). In both models we assumed a melt-vapor fractionation factor $A = +3$ in accordance with the average from literature and close to our estimation for Piton de la Fournaise (Javoy et al., 1978; Matthey, 1991; Aubaud et al., 2006). With these models, we fully overlap the variability of starting conditions for hotspots (Barry et al., 2014). In fact, there is a good consistency between our model for PdF (Model 1) and the Kilauea model (Model 2). Conversely, BED and FED trajectories predicted from the Pitcairn model (Model 3) does not fit our data, most of them lying above the BED trajectory, even above the starting conditions for some of them (Fig. 7c) that make this model unrealistic for PdF.

Therefore, in our case, we stress that (1) $\delta^{13}\text{C}$ variability observed in our dataset is consistent with different stages of degassing and (2) estimated starting conditions are fully in accordance with values reported at Hawaii in a similar geodynamic context. It is important to note that if the

modeling proposed in Model 1 (Fig. 7a) is linked to the amount of CO_2 released by crushing instead of the amount of CO_2 dissolved in the melt, it cannot in any way affect the interpretation of the $\delta^{13}\text{C}$ signature. For instance, the $\delta^{13}\text{C}$ signature of PDC ($-6.4 \pm 0.5\text{‰}$) can only be reached by a FED trajectory and highlights an important extent of degassing with respect to the source ($90 \pm 7 \text{ wt}\%$) whatever is the initial CO_2 content assumed for modeling FED and BED trajectories.

5.2.2. He-Ar- CO_2 systematics

We report that olivine crystal composition (Fo; Ca) and extracted gas content from PdF pyroclasts are consistently record melt differentiation and degassing at variable depths in the volcano plumbing system. Enclaves exhibit a more complex signature, which could result from a combination of both magma degassing and re-equilibration processes during ponding and long-lasting residence at depth. Actually, even if CO_2 content is not significantly affected by diffusion across the host crystals, diffusive fractionation could affect the variability of other gaseous species trapped in FIs like H_2O and noble gases (Danyushevsky et al., 2002; Burnard, 2004; Yamamoto et al., 2009). We will now investigate the processes affecting He, Ar and CO_2 contents released by crystal crushing.

We first consider our data in terms of classical modeling of magmatic degassing, controlling noble gas and CO_2 ratios. Actually, $^4\text{He}/\text{CO}_2$ and $^4\text{He}/^{40}\text{Ar}^*$ bring important information on magmatic degassing process (e.g., Burnard, 2004; Burnard et al., 2003, 2014; Paonita et al., 2012; Barry et al., 2014; Correale et al., 2014). The degassing trends potentially recorded in the vapour phase

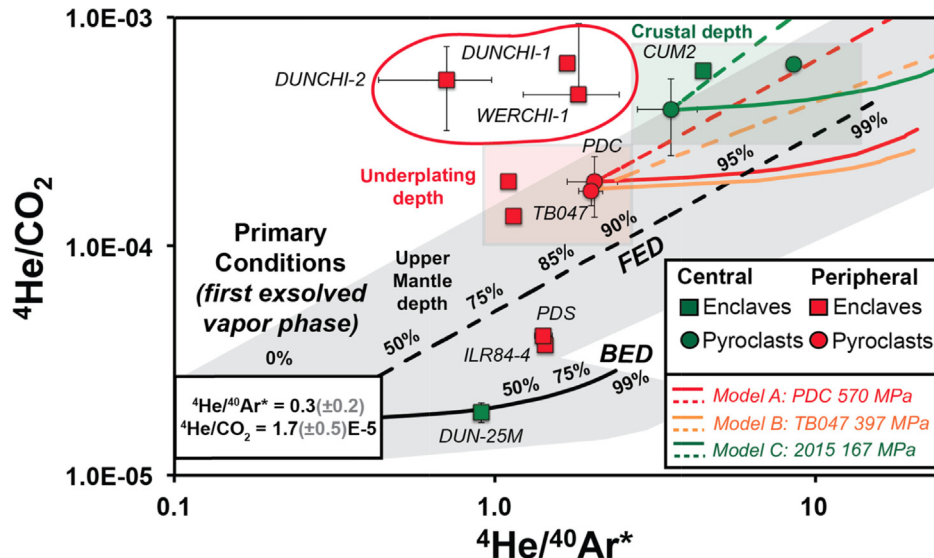


Fig. 8. Modeling of $^4\text{He}/^{40}\text{Ar}^*$ vs. $^4\text{He}/\text{CO}_2$ in the fluid phase following Fractional Equilibrium Degassing (FED) and Batch Equilibrium Degassing (BED) paths. The extents of degassing (0–50–75–85–95–99%) are reported along the curves. Primary conditions are calculated assuming that PDC has undergone 89% of open-system degassing with respect to primary melts as suggested by the $\delta^{13}\text{C}$ analysis in this study. Starting conditions used for Models A-B-C are defined in Table 4 (see text for explanations). Enclaves deviating from equilibrium degassing paths are identified (red contour line). Color filled areas correspond to specific trapping depth ranges and relative position in the magmatic plumbing system of PdF. Grey field highlights the expected range of BED and FED considering the uncertainty related to the starting conditions ($^4\text{He}/^{40}\text{Ar}^*$ and $^4\text{He}/\text{CO}_2$). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

extracted from the melt were modeled using the BED and FED equations (Fig. 8; Jambon et al., 1986; Shaw et al., 2004, Hoefs, 2015). We constrain starting conditions by taking into account (1) the ${}^4\text{He}/{}^{40}\text{Ar}^*$ measured in PDC products (2.05 ± 0.37), (2) that PDC products suffered 90 ± 7 wt% of FED as suggested by the $\delta^{13}\text{C}$ study, (3) Ar-He solubility ($S_{\text{He}}/S_{\text{Ar}} = 10$) from representative melt inclusion composition from PDC olivines (Bureau et al., 1998; Table 4). In the case of $S_{\text{He}}/S_{\text{Ar}}$, we assume the linearity of Ar-He solubility up to 5 GPa (Sarda and Guillot, 2005), i.e. compatible with source melting conditions for La Réunion hotspot (partial melt production at 4 GPa; Fretzdorff and Haase, 2002). Consequently, we have calculated that the first exsolved vapour phase has an initial ${}^4\text{He}/{}^{40}\text{Ar}^*$ of 0.3 ± 0.2 (Fig. 8). Having in mind that the typical production ratio in the mantle is 2–5 (Marty, 2012) and that $S_{\text{He}}/S_{\text{Ar}} = 10$ (Poreda and Farley, 1992; Matsuda and Marty, 1995; Iacono-Marziano et al., 2010; Tedesco et al., 2010; Marty, 2012), our estimated ratio is within the range expected for the first degassing vapour (${}^4\text{He}/{}^{40}\text{Ar} = 0.2\text{--}0.5$). Our carbon study has suggested an initial $\delta^{13}\text{C}$ of $-0.8 \pm 1.0\text{‰}$ in the first exsolved fluid phase and a potential maximum CO_2 content of $3.6 \pm 0.7 \times 10^{-6}$ mol g^{-1} from the crushing procedure (Fig. 7a). Using the fitting power function from Fig. 4b, we estimate a corresponding initial ${}^4\text{He}$ content of $6.2 \pm 0.7 \times 10^{-11}$ mol g^{-1} and thus an initial ${}^4\text{He}/\text{CO}_2$ of $1.7 \pm 0.5 \times 10^{-5}$.

${}^4\text{He}/{}^{40}\text{Ar}^*$ vs. ${}^4\text{He}/\text{CO}_2$ degassing trends suggest that FED is predominant in the evolution from the source to deep magmas feeding peripheral eruption to shallow magmas producing the central activity (Fig. 8). Conversely, the data of some enclaves (PDS, ILR84-4, DUN-25M) lie between FED and BED trajectories. These modeling paths are fully consistent with those predicted by the $\delta^{13}\text{C}$ (Fig. 7). Deviations from these degassing trends are mainly observed in Piton Chisny enclaves (DUNCHI-1, DUNCHI-2 and WERCHI-1), characterized by apparently higher ${}^4\text{He}/\text{CO}_2$ or lower ${}^4\text{He}/{}^{40}\text{Ar}^*$ (Fig. 8).

Noble gases and CO_2 contents in the vapour phase are dependent on their initial abundance, diffusivity and solubility in melts (Carroll and Stolper, 1991, 1993; Shibata et al., 1996, 1998; Chamorro-Perez et al., 1996, 1998; Paonita et al., 2000; Schmidt and Keppler, 2002; Miyazaki et al., 2004; Marrocchi and Toplis, 2005; Paonita, 2005 and references therein; Bouhifd and Jephcoat, 2006; Gonnermann and Mukhopadhyay, 2007; Iacono-Marziano et al., 2010; Shishkina et al., 2014). CO_2 and Ar diffusivities in basaltic melts are similar, while that of He is 100 times higher (Gonnermann and Mukhopadhyay, 2007). In case of disequilibrium degassing, we would expect a similar enrichment in Ar and CO_2 of the vapour phase but a more extensive enrichment in He, and that is not observed in our samples (Fig. 4). In our dataset, only DUNCHI-1 could be the result of disequilibrium degassing, however this process would imply a fast degassing dynamics. We argue that DUNCHI-1 is a cumulate that should have been formed from melts ponding during long periods, a process that should favor equilibrium degassing. Therefore, we exclude disequilibrium degassing or it has a restricted role in the interpretation of our dataset.

Gas solubility in silicate melt is affected by numerous factors, as temperature, pressure, oxygen fugacity and melt composition. At PdF, the (slight) compositional variability of both peripheral and central melts was mainly attributed to crystallization-assimilation processes and only subordinately to source heterogeneities (see Vlastelic and Pietruzka (2016) for a review). In order to cover the variability of PdF melts in terms of chemical composition and pressure-temperature domains, we have tested three different models based on different initial melt inclusion compositions (corrected from post-trapping effects; Boudoire, 2017). Melt inclusions were measured in olivines extracted from the same products analysed for noble gases in this study (Table 4; Bureau et al., 1998). In order to model magma degassing, we choose the maximum pressure recorded in the FIs of three distinct eruptive products as starting pressure for each model. Similarly, Viti and Frezzotti (2000) suggest that the maximum pressure estimated from FIs is more likely to provide a reliable estimation of the crystallization depth. Model A: based on melt inclusions in olivines from the peripheral PDC eruptive site, with a starting pressure of 570 MPa. Model B: based on melt inclusions in olivines from the peripheral Trouis Blancs (TB047) eruptive site, with an initial pressure of 397 MPa. Model C: based on melt inclusions from the central August 2015 (REU151026-1a) eruption and a starting pressure of 167 MPa. These three models cover a large range of pressure (167–570 MPa), temperature (1218–1281 °C; from the hydrous-composition dependent equation of Putirka (2008)), and melt compositions (MgO: 8.1–10.9 wt%; K_2O : 0.5–0.9 wt%; SiO_2 : 47.7–49.7 wt%; H_2O : 0.7–1.4 wt%). They broadly overlap the geochemical variability of PdF melts in both major, traces and isotope signature in the whole volcano plumbing system (Valer, 2016). Noble gases and CO_2 solubility were calculated using experimental results (e.g., Iacono-Marziano et al., 2010; Duan, 2014 and references therein), well adapted for this compositional and pressure-temperature range, and are thus representative of the range of solubility expected for PdF melts. These calculations show that solubility in all studied melts decreases from He to CO_2 to Ar (Table 4; e.g., Iacono-Marziano et al., 2010; Paonita et al., 2012 and references therein).

Interestingly, the degassing trends indicate no major difference between the three models (Fig. 8) suggesting that melt composition, temperature and pressure variability play thus a minor role in the evolution of He-Ar- CO_2 ratios by degassing at PdF. For instance, Model A-B-C present a very similar behavior in spite of (1) almost 400 MPa of pressure change and, (2) geochemical variability in melt composition spanning the full range reported for PdF (Table 4; Valer, 2016). This suggests that the deviations from the predicted degassing trends recorded by Piton Chisny enclaves (DUNCHI-1, DUNCHI-2 and WERCHI-1) reveal the occurrence of processes other than the geochemical variability of PdF magmas. Actually, we have documented that these products (1) fall out of the 95%-confidence interval of the gas fitting equations (Fig. 4), (2) generally record a shallower pressure mode than other peripheral products (Fig. 5c) and, (3) present a distinct petrological signature (Fig. 6b). This could

highlight an older/distinct magmatic history with respect to other samples analysed in this study (Matsuda and Marty, 1995; Burnard, 2004; Gautheron et al., 2005; Yamamoto et al., 2009).

5.3. The architecture of the magmatic plumbing system beneath Piton de la Fournaise

Deciphering deep degassing in volcanic systems is challenging and is generally based on experimental results and modeling. At PdF, we have based our study on a multiparametric approach coupling barometric data, olivine composition, melt and fluid inclusion chemistry, noble gases, CO₂, and carbon isotopes systematic. Consistently with previous studies, our result confirm that central (intracaldera) eruptions emit evolved and degassed melts stored at shallow crustal levels, whereas the products of peripheral eruptions are more primitive, gas-richer and record the signature of deeper levels, whose pressure fits with that of the mantle-crust underplating layer or even higher depth (see Di Muro et al., 2016 for a review). Peripheral enclaves (PDS and ILR84-4) testify of the deepest origin (upper lithospheric mantle) in our dataset.

Based on these results, we put new constrains on the plumbing system and magma degassing at PdF. PDC peripheral pyroclasts is the sample for which the largest dataset is available (Bureau et al., 1998; Welsch et al., 2013). By using melt inclusions composition (Table 4) with 1.1 wt% of dissolved H₂O (measured in the PDC melt inclusion and close to the primary melt estimation of Bureau et al. (1998) and Di Muro et al. (2014)), H₂O-CO₂ saturation pressure modeling (Duan, 2014) predicts that 0.35 wt% of CO₂ is dissolved in the melt at 570 MPa (maximum pressure from fluid inclusions barometry; Table 1). Our study suggests that this pressure is representative of the crystallization depth. The calculated amount of dissolved CO₂ is fully in accordance with results from other H₂O-CO₂ saturation models (Papale et al., 2006), and is very similar to the mean estimation of 0.30 wt% of CO₂ in MORB-like undegassed magmas and in the mantle (Gale et al., 2013; Michael and Graham, 2015). This value is higher than the maximum concentration of dissolved CO₂ measured in PDC melt inclusions by FTIR (i.e., 0.18 wt%; 280 MPa; Bureau et al.,

1998; Di Muro et al., 2016). However, many post-trapping process may affect the CO₂ estimation in melt inclusions (Wallace et al., 2015; Maclennan, 2017). In fact, Maclennan (2017) has recently shown that most of the estimated H₂O-CO₂ saturation pressures reflect extensive decrepitation of melt inclusions during melt ascent (when the differential pressure between the melt inclusion and the external environment is higher than 250 MPa). An amount of CO₂ dissolved in PDC melt of about 0.35 wt% is thus a reliable assumption. Based on these starting conditions and on the predominant role of fractional equilibrium degassing (FED) in our dataset, we estimate that (Table 5):

- (1) The primary melt formed at 4 GPa has up to 3.5 ± 1.4 wt% of dissolved CO₂ and the first exsolved vapor phase an initial $^4\text{He}/^{40}\text{Ar}^*$ of 0.3 ± 0.2 (considering the $90 \pm 7\%$ of FED degassing suggested for PDC on Figs. 7 and 8). The initial CO₂ content is much higher than previous estimation made by Bureau et al. (1998) at PdF (1.0 wt%) based on melt inclusions H₂O-CO₂ saturation pressure modeling (Table 5) predicts pressure higher than 3 GPa (limit of the model; Duan, 2014), in agreement with the depth of partial melt extraction from the mantle source beneath PdF (ca. 4 GPa; Fretzdorff and Haase, 2002),
- (2) The parental melt triggering the crystallization of our most primitive sample (ILR84-4 harzburgite) could have an amount of dissolved CO₂ of 1.1 ± 0.9 wt% (average between BED and FED trajectories for ILR84-4 on Figs. 7 and 8), fully consistent with previous estimations made by Bureau et al. (1998) at PdF (1.0 wt%). In the fluids related to this primitive magma, the $^4\text{He}/^{40}\text{Ar}^*$ should be about 0.8 ± 0.6 . Based on CO₂-H₂O saturation pressures modeling (Duan, 2014), we estimate that the parental melt that formed ILR84-4 ponded at pressures close to 1 GPa (Table 5). This pressure is consistent with the deep melt horizon identified beneath La Réunion Island by seismic studies (Gallart et al., 1999; Massin, 2009; Fontaine et al., 2015), stress-modeling (Gerbault et al., 2017) and petrological models (Albarède et al., 1997; Fretzdorff and Haase, 2002).

Table 5

Reconstruction of initial gas composition and conditions (in bold) for distinct PdF products representative of different levels of the plumbing system (taking into account the analytical uncertainty and the variability of the analysis). (1) Estimation of the degassing extent from the initial $\delta^{13}\text{C}$ calculated in this study (from -4.2‰ to -1.8‰ considering all uncertainties during the modeling). (2) Estimation of initial CO₂ content is obtained from the degassing extent for an initial 0.35 wt% CO₂ dissolved content in Piton de Caille melt inclusion at 5.7 kbar (maximum pressure) (3) Estimation of the saturation pressure of CO₂-H₂O (Duan, 2014) based on the inferred composition of the PdF primary melt (Fretzdorff and Haase, 2002). (4) Recalculation of the initial $^4\text{He}/^{40}\text{Ar}^*$. (5) Estimation of $\delta^{13}\text{C}$ values in central products.

	Crust-edifice Central Pyroclasts REU151026-1a		Underplating depth Peripheral Pyroclasts PDC		Upper mantle Peripheral Enclaves ILR84-4		Source Primary Melt Primary Melt	
	Average	σ	Average	σ	Average	σ	Average	σ
(1) Degassing extent	94	5	90	7	68	13	0	0
(2) Max dissolved CO ₂ (wt%)	0.08		0.35		1.1	0.9	3.5	1.4
(3) Max P (GPa)	0.17		0.57		1.3	0.7	>3	
(4) $^4\text{He}/^{40}\text{Ar}^*$	3.5	0.8	2.1	0.4	0.8	0.6	0.3	0.2
(5) $\delta^{13}\text{C}$ FED (‰)	-8.3	3.1	-6.4	0.5	-3.2	0.1	-0.8	1.0

(3) The low CO_2 content of olivine from central pyroclasts hinders the measurement of $\delta^{13}\text{C}$ by crystal crushing. The open-degassing paths suggested by noble gases ratios from FIs of PDC olivines and central products (Figs. 5 and 7) argue for at least $94 \pm 5\%$ of deep degassing for these products with respect to the primary melt. Starting from the $\delta^{13}\text{C}$ of the initial vapor phase calculated in this study ($-0.8 \pm 1.0\text{‰}$), we consequently estimate that the $\delta^{13}\text{C}$ of the central products (here based on the August 2015 eruption: REU151026-1a) could be as low as $-8.3 \pm 3.1\text{‰}$ just following an open-degassing path ($94 \pm 5\%$; FED). These estimations are more negative than $\delta^{13}\text{C}$ measurements performed by Marty et al. (1992) in summit fumaroles of PdF (-4.1‰) and in the thermal springs located on PdN volcano (-4.7‰ ; Cilaos in Fig. 1; 30 km-far from the PdF summit), during a quiescent period 25 yrs ago. However, magmatic dynamics could have changed between 1992 and 2015, especially after the 2007 caldera-collapse eruption. Actually, fumaroles sampled by Marty et al. (1992) are not accessible anymore, while current analysis (2016–2017) in gases from Cilaos thermal springs indicate lower $\delta^{13}\text{C}$ values (around -6‰ ; OVPF database) than fumaroles and thermal springs collected by Marty et al. (1992).

5.4. Implications on the extensive degassing in the upper mantle beneath Piton de la Fournaise

Regarding these new results, we now propose a model of magma degassing from mantle partial melting up to the

crustal plumbing system feeding the activity of PdF volcano (Fig. 9). Partial melt extracted from the mantle source can contain up to $3.5 \pm 1.4 \text{ wt}\%$ of dissolved CO_2 . Assuming $\approx 7.5 \pm 2.5\%$ of melting of the mantle plume source (Valer et al. (2017b) and references therein) and the incompatible nature of carbon during melting (see Rosenthal et al. (2015) for a review), we estimate a minimum of $716 \pm 525 \text{ ppm}$ of C in the mantle (i.e., considering C as fully incompatible element). This value is much larger than estimations made for MORB source mantle ($<140 \pm 30 \text{ ppm C}$; Cartigny et al., 2008; Hirschmann and Dasgupta, 2009; Marty, 2012; Rosenthal et al., 2015), but is consistent with (1) the estimation made for the bulk mantle ($765 \pm 300 \text{ ppm C}$; Marty, 2012) and (2) recent studies pointing out that carbon content in the mantle may be strongly underestimated (Anderson and Poland, 2017). According to our calculations, most of the CO_2 ($68 \pm 13\%$) is lost early, leading to partially degassed primary melts ($\text{CO}_2 = 1.1 \pm 0.9 \text{ wt}\%$) ponding at around 1 GPa in the upper mantle.

In the upper mantle (30–11 km bsl), melts undergo early differentiation and extensive degassing as suggested by the composition of related melt inclusions ($\text{MgO} < 10 \text{ wt}\%$ in PDC; Bureau et al., 1998), the low-magnesium olivines ($\text{Fo}_{<87}$; this study; Bureau et al., 1998) and the low $\delta^{13}\text{C}$ values measured in FI from olivines of peripheral pyroclasts ($< -5\text{‰}$). At the depth of the mantle-crust underplating layer, $90 \pm 7\%$ of the initial CO_2 is already lost (Figs. 8 and 9). Long lasting magma storage and complex thermal paths of magmatic cumulates emplaced at the depth of the underplating layer may favor (i) FI volumetric resetting and re-equilibration processes, and (ii) melt homogenization (Jennings et al., 2017).

Melts reaching the central magmatic reservoirs at crustal level are even more evolved ($\text{Fo}_{<85}$) and degassed (Figs. 6, 8

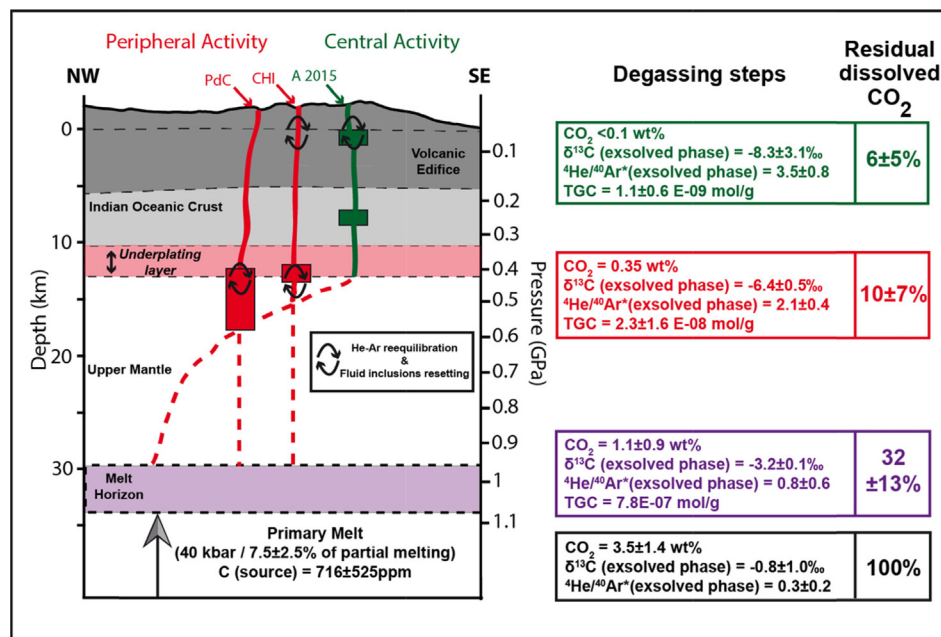


Fig. 9. Conceptual model of degassing in the magmatic plumbing system of PdF. Multistep magma ponding enhance extensive degassing, re-equilibration processes, melt differentiation and volumetric resetting of fluid inclusions. Thick coloured dashed lines highlight potential magmatic pathways (Michon et al., 2015; Boudoire, 2017).

and 9) than peripheral melts. This is consistent with the fact that the central and frequent activity of PdF volcano emits degassed magmas and is associated with very weak gas emissions (Liuzzo et al., 2015; Di Muro et al., 2016). Our findings contribute to explain the absence of permanent and sustained CO₂-rich fumarolic and soil emissions inside the PdF caldera as well as in most of the parts of La Réunion island. We have estimated that early and extensive degassing ($94 \pm 5\%$) can lower $\delta^{13}\text{C}$ values of magmatic fluids down to about -8% . Such negative values raise important issues regarding the inferences made about the contribution of exogenous processes in low carbon isotopic signatures of CO₂. In fact low carbon isotope values in volcanic gas emissions are often attributed to dissolution process or carbonate precipitation (Hoefs, 2015). Interestingly, the highest $\delta^{13}\text{C}$ values measured at PdF (in a tunnel) since the 2007 major eruption has similar values ($-8.2 \pm 0.3\%$; Liuzzo et al., 2015), and could thus correspond to a late stage of degassing in intracrustal conditions (Fig. 9).

5.5. Comparison with other ocean basaltic volcanoes and involvements regarding mantle CO₂

The extensive degassing we argue below PdF is unusual and appears in contrast with observations made on other ocean basaltic volcanoes like Kilauea. As a matter of fact, at Kilauea (1) high CO₂ fluxes are reported at the surface with a high carbon isotope signature, ranging from -3.4 to -4% (Gerlach and Taylor, 1990; Gerlach et al., 2002; Burton et al., 2013; Poland et al., 2014), and (2) erupted melts are primitive (MgO-rich melts and Fo-rich olivines; Helz et al., 2014) leading to the emission of tholeiitic basalts. Actually, the main similarity highlighted by this study between both volcanoes, is the presence of a deep magma ponding zone (ca. 30 km-depth) characterized by (1) a similar dissolved CO₂ content (1.1 ± 0.9 wt% at PdF; $0.7\text{--}1.2$ wt% at Kilauea; Gerlach et al., 2002; Wech and Thelen, 2015; Anderson and Poland, 2017) and (2) a similar carbon isotopic signature in the exsolved vapor phase ($-3.2 \pm 0.1\%$ at PdF; -3.4 down to -4% at Kilauea; Gerlach et al., 2002; Poland et al., 2014). These features suggest that the main differences regarding magma evolution of both volcanoes are related to magmatic processes occurring during magma ascent and ponding in the last 30 km-depth. Recent studies have shown that PdF melts undergo numerous stages of differentiation in the upper mantle and the crust due to the presence of multiple magma ponding zones (Bureau et al., 1998; Michon et al., 2015; this work). Conversely, at Kilauea, Rae et al. (2016) have shown that magma ascent could be extremely fast from 45 to 65 km-depth, as short as weeks to months that could be partially related to the more important magma supply at Kilauea (Poland et al., 2014). Jaupart (1998) has shown that slow magma ascents favor extensive degassing whereas fast magma ascents are more able to limit the degassing. Similarly, we argue that multiple magma ponding zones in the upper mantle, then in the crust, may account for an early extensive degassing and differentiation, characterizing PdF melts.

We infer that the mantle/crust underplating layer seems to play a critical role in the differentiation and degassing of PdF melts, and opens new insights on the following issues:

- (1) the high extent of CO₂ degassing during magma ponding at the depth of the mantle-crust underplating layer may represent a worldwide feature rather than a process specific to La Réunion Island. Actually, our model (Fig. 8) might be a classical model for numerous other ocean basaltic volcanoes, where barometric estimations evidence the presence of multiple deep magma ponding zones playing a major role in melt differentiation as e.g. Azores, Cape Verde, Canarias Islands (Zanon and Frezzotti, 2013; Klügel et al., 2015; Longpré et al., 2017),
- (2) the low $\delta^{13}\text{C}$ signature inferred for mantle reservoirs (-6% ; Loihi, Pitcairn; down to -8% in MORB mantle, e.g., Sano and Marty, 1995; Deines, 2002; Aubaud et al., 2004 and references therein) could already reflect an early stage of degassing in the upper mantle rather than a contribution of recycled carbon (Exley et al., 1986; Aubaud et al., 2006), or a mantle heterogeneity.

Consequently, the total budget of mantle CO₂ may be strongly underestimated when based on gas emission at the surface, on dissolved CO₂ from melt inclusions or calculated considering the low carbon isotopic signature of CO₂ in MORB glasses as starting conditions (Aubaud et al., 2006; Burton et al., 2013; Barry et al., 2014). For instance, previous estimations at PdF were based on a maximum dissolved CO₂ content of 1 wt% in melt (Bureau et al., 1998), i.e. 2–5 times less than our estimation (3.5 ± 1.4 wt%). Our study fully supports similar recent reevaluations of mantle CO₂ like (i) at Kilauea by coupling volcanic CO₂ emission rates with probabilistic magma supply rates (Anderson and Poland, 2017) and, (ii) at El Hierro based on the highest CO₂ contents ever measured in melt inclusions from ocean island volcano (Longpré et al., 2017). Note that at El Hierro, these authors have estimated a primitive dissolved CO₂ content in the range 2.5–5.0 wt% (by extrapolation to undegassed CO₂/Nb), thus overlapping our estimation for PdF melts (3.5 ± 1.4 wt%).

Our reevaluation of the mantle CO₂ beneath PdF opens new prospects regarding past climate dynamics and its links with the volcanic activity (Anderson and Poland, 2017; Gutjahr et al., 2017; Hasenclever et al., 2017; Kobashi et al., 2017; Tobin et al., 2017). More generally, this large amount of mantle CO₂ opens up the possibility that deep Earth volatile cycle related to OIB needs a complete reevaluation, as suggested by some high H/C OIB sources that are probably overestimated (Hirschmann & Dasgupta, 2009). Therefore, we strongly encourage further studies to reevaluate the global volcanic volatile budget induced by these new estimations of the initial mantle carbon and investigate past and potential future effects on the Earth atmosphere during major eruptions.

Finally, but not least important, this works suggest that monitoring of $\delta^{13}\text{C}$, $^4\text{He}/^{40}\text{Ar}^*$, $^4\text{He}/\text{CO}_2$ temporal variations in surface gas emissions of volcanoes characterized by very frequent eruptions may track deep magma arrival

and constrain the degassing pressure from various parts of a plumbing system. Actually, the highest $\delta^{13}\text{C}$ values recognized in gas emissions at La Réunion are around -6‰ with $^4\text{He}/^{40}\text{Ar}^* \sim 1.75$ (Cilaos free gases; OVPF database; Marty et al., 1992), probably tracking a contribution of late-degassed melts stored at the depth of the underplating layer as inferred by PDC melts.

6. CONCLUSIONS

Deep degassing below ocean basaltic volcanoes is still poorly constrained and only a few studies quantify the degassing extent as a function of depth. Here, we focused on one of the most active volcanoes on Earth, the Piton de la Fournaise volcano, whose gas emissions at the surface are very weak in spite of its frequent eruptions. This anomalous behavior opens an important issue whether gases released from magma may escape from lateral and submarine paths of ascent or whether the subaerial weak measured emissions reflect a very degassed system at depth. In order to address this question, we have studied a set of crystals from fast quenched pyroclasts and enclaves sampled both in the peripheral and in the currently active central area. Coupling barometric estimations from olivine FIs with mineral chemistry, carbon isotopes of CO_2 and noble gases in FIs we infer that peripheral products are generally more primitive, gas-rich, and have a deeper (mantle) origin whereas, the central (intracaldera) ones are fed by more evolved and degassed magmas stored at shallower crustal levels.

Based on He-Ar- CO_2 systematics, we quantify the extent of degassing from the upper mantle to the magmatic plumbing system beneath the volcano. At PdF, our data and modeling are in agreement with a predominant open system equilibrium degassing. Coupling barometric estimations and melt inclusions composition with carbon isotopes and noble gases systematics permit us to propose that:

- (1) primary basaltic melts of PdF have a dissolved CO_2 content around $3.5 \pm 1.4 \text{ wt}\%$; (this study), significantly higher than previous estimated ($1 \text{ wt}\%$, Bureau et al., 1998) and involving a carbon-rich mantle plume source ($716 \pm 525 \text{ ppm C}$),
- (2) $\delta^{13}\text{C}$ of primary vapor phase is estimated to be $-0.5 \pm 0.5\text{‰}$,
- (3) melt ponding at mantle lithospheric depth (30 km-depth) has already undergone almost $68 \pm 13\%$ of degassing,
- (4) multiple magma ponding zones in the last 30 km permit advanced melt differentiation and extensive degassing ($90 \pm 7\%$) and account for the weakness of CO_2 gas emissions and the emission of relatively evolved basalts in the central active area,
- (5) central melts erupted by the most active area are even more degassed and this might explain the very low $\delta^{13}\text{C}$ signature measured in surface gases (down to -8‰). We point out that this low isotopic signature could be common in volcanic gases, and reflect extensive CO_2 degassing during magma ascent towards the central part of the plumbing system (up to $94 \pm 5 \text{ wt}\%$ in this study).

We highlight that the main difference in early deep degassing between PdF and other oceanic islands like Kilauea appears to be related to the presence of multiple magma ponding zones starting already in the upper mantle ($<30 \text{ km}$ -depth) beneath PdF, leading to extensive differentiation and degassing. This model might be applied to numerous ocean basaltic volcanoes where similar plumbing systems are recognized. Consequently, important implications come for the current budgets estimations of CO_2 degassed from the Earth mantle, which could be largely underestimated. For instance, in this study we provide a new estimation of the carbon content in the mantle below PdF ($716 \pm 525 \text{ ppm C}$) that is considerably higher than the few previous calculations performed for Ocean Island Basalts (OIB) systems but is consistent with the recent reevaluation of the volcanic carbon budget made by the scientific community.

Finally, this study opens exciting perspectives regarding both volcanic volatile budget, CO_2 recycling in the mantle, past climate dynamics and monitoring strategy, especially for the Deep Carbon Observatory community. The temporal variability of $\delta^{13}\text{C}$ could bring strong information regarding magmatic dynamics occurring in the plumbing system.

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SUPPLEMENTARY MATERIAL

Supplementary data (Appendix A, B, C, D) associated with this article can be found, in the online version, at <https://doi.org/10.1016/j.gca.2018.06.004>.

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