RESEARCH ARTICLE



Explosive style, magma degassing and evolution in the Chaimilla eruption, Villarrica volcano, Southern Andes

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Abstract Villarrica (Chile) is a basaltic stratovolcano, currently in an open-conduit condition. It now has relatively frequent Strombolian and effusive eruptions, but it had large explosive eruptions in prehistoric times. Among them, the most recent eruption was Chaimilla, which occurred about 3100 years ago and produced deposits that indicate complex, multiphase eruptive dynamics. Significant differences in mineralogy and glass compositions of the erupted scoria suggest the eruption was fed by two distinct magma batches with similar bulk compositions but distinct crystallization and degassing histories. The lower sequence scoria has a complex crystal assemblage with several crystal populations produced by mixing between a relatively degassed magma containing Fo75-79 olivine, normally or reversely zoned plagioclase (An₇₀₋₉₄) and augite (type 1 magma), and a subordinate volume of more-primitive and more volatile-rich magma rising from depth (type 2 magma) and carrying normally zoned plagioclase and higher-Mg (Fo₈₁₋₈₅) olivine crystals. Type 2 magma was the main component emitted during the larger and more explosive eruptive phase that deposited the upper

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sequence. The Chaimilla eruption occurred under closedvent conditions and was fed by water-rich magmas. When compared with the petrological features of the magma currently erupted at Villarrica, which has slightly more-evolved bulk compositions, lower crystal content and lower water content, these results suggest that the evolution in eruptive style of the volcano from highly explosive to a lava lake/Strombolian activity corresponds to significant changes in the shallow plumbing system (which is now at much shallower depths); these plumbing-system changes were not associated with significant changes in the parental magma compositions.

Keywords Basalt explosivity \cdot Melt inclusion \cdot Chilean volcanism

Introduction

Magmatic volatiles and degassing play important roles in explosive eruptions because they influence magma properties, fragmentation and eruptive styles (Wilson et al. 1980; Melnik 2000; Cervantes and Wallace 2003; Del Carlo and Pompilio 2004; Di Muro et al. 2004; Goepfert and Gardner 2010; Métrich et al. 2010). In particular, high initial volatile content and fast ascent rate favour explosivity (Cashman 2004). Degassing-induced crystallization of phenocrysts and microlites has been proposed as an additional process controlling pre-eruptive magma viscosity and, thus, explosivity (Blundy and Cashman 2005; Erlund et al. 2010).

Among basaltic systems, the role of volatiles in increasing explosivity is particularly relevant in volatile-rich magmas erupted in subduction settings. Several studies have recognized that basalts associated with highly explosive eruptions contain high initial water content (up to 6 wt.%, Roggensack et al. 1997; Newman et al. 2000; Pichavant et al. 2002; Grove et al. 2005: Gurenko et al. 2005: Spilliaert et al. 2006: Kamenetsky et al. 2007; Johnson et al. 2008; Métrich et al. 2010), suggesting that pre-eruptive volatile content, degassing dynamics and the amount of volatiles available in the magma at fragmentation strongly correlate with eruptive style. The importance of pre-eruptive degassing and outgassing is particularly evident at volcanoes that display a wide range in eruptive styles, from quiet lava emission to violent (subplinian to Plinian) explosions, but only limited variation in bulk chemistry of the erupted magma. Among them, Villarrica volcano (Southern Chile) is a striking example because of its current Hawaiian-Strombolian eruption style associated with a summit lava lake and its large explosive eruptions in the recent past (Clavero and Moreno 2004; Silva Parejas et al. 2010; Costantini et al. 2011).

We present a petrological and geochemical study of wellexposed deposits from a recent basaltic explosive eruption of Villarrica volcano, Chile, the 3.1-ka Chaimilla deposit, which has a volcanic explosivity index (VEI) of 4 (Costantini et al. 2011). Our aim is to determine pre-eruptive conditions and magma histories. The new dataset on explosive products is then compared with available data on magma erupted in prehistorical and recent activity (Vargas et al. 1989; Witter et al. 2004).

Geological setting and eruptive history

Villarrica is a stratovolcano located in the Southern Chilean Andes (Fig. 1). It is the most active volcano of this segment of the Andes in historical times (Lara 2004). The central cone (2847 m) lies on the NW rim of a 6.5×4.2 km caldera (Moreno et al. 1994), which formed during large-volume eruptions associated with emplacement of the Licán ignimbrite (~10 km³, 13.8 ka BP) and the Pucón ignimbrite (\sim 5 km³, 3.5 ka BP). Both ignimbrites have basaltic-andesite composition (Lohmar et al. 2012; Silva Parejas et al. 2010). Villarica's historical eruptive activity is associated with the central cone and has been mainly effusive. However, its postcaldera activity has also been mildly explosive (Van Daele et al. 2014), producing both large pyroclastic density currents (PDCs) and tephra fallout (not dated, Clavero and Moreno 1994; Lara 2004; Moreno and Clavero 2006). After the last eruption in 1984-1985, which produced both lava fountains and lava flows, Villarrica volcano maintained open-conduit conditions and an intermittent summit lava lake with very limited magma emission (Witter et al. 2004; Gurioli et al. 2008). In addition, partial melting of the summit glacier has generated lahars, which caused several fatalities during the 1948-1949, 1963-1964 and 1971-1972 eruptions (Rivera et al. 2008; Castruccio et al. 2010).



Fig. 1 a Location of Villarrica volcano. b Stratigraphy and sampling beds at the type outcrop. The outcrop is located about 3.2 km N of the summit crater, along the dispersal axis of tephra beds of the units A, B, G and H

The Chaimilla eruption

The Chaimilla eruption (~3.1 ka) produced one of the bestpreserved and most widely dispersed pyroclastic deposits of Villarrica volcano. The deposit is now exposed over an area of 250 km^2 on the northern flank of the edifice and has a minimum volume of about 0.5 km³ (Costantini et al. 2011). The deposit comprises tephra from four different eruptive sequences (basal, lower, middle and upper, Fig. 1), which consist of several units (A–H) lying on top of the Pucón Ignimbrite, which was erupted 400 years earlier:

1. The basal sequence (BS) consists of two main ash and lapilli beds (units A, about 1.1×10^7 m³, and B, about 7.2×10^6 m³), produced during the opening phase of the eruption by repeated explosions that dispersed ash and

lapilli clasts up to several kilometres away from the vent, along with abundant lithic blocks and dense juvenile bombs in proximal areas.

- 2. The lower sequence (LS) consists of two fall deposits: C (about 1.4×10^8 m³) and D (about 1.3×10^7 m³). Unit C has wider dispersal than D and is made up of at least 20 different layers with distinct grain size (from coarse ash to bombs) and lithic contents. Unit D is a poorly dispersed, massive, moderately sorted layer, formed of glassy and highly vesicular scoria fragments. LS has been interpreted as the product of pulsatory activity associated with a series of discrete explosions of variable intensity or with a pulsating column (Costantini et al. 2011).
- 3. The middle sequence deposit consists of ash-rich, poorly sorted layers deposited by a PDC (unit E) and associated with a co-PDC ash layer (unit F), with a combined total volume of about 5×10^6 m³ (Costantini et al. 2011).
- 4. The upper sequence (US) consists of two units. Unit G is a structureless lapilli fall deposit, whereas unit H consists of a complex series of scoria lapilli fall beds and finely laminated PDC beds, which are inferred to result from partial to total collapse of the eruptive column. The total volume of the fall deposit is at least to 3.6×10^8 m³, and the PDC beds have a minimum volume one order of magnitude smaller.

Methods

Sampling and sample preparation

Samples were collected from lapilli fall layers of the Chaimilla deposit in a vertical section at a single outcrop located 3.2 km N of the vent (Fig. 1), along the downwind direction. In particular, we collected samples from units C and D in the LS and units G and H in the upper sequence. Whole-rock analyses were made on representative samples comprising 5–10 scoria lapilli collected from single stratigraphic layers.

Olivine crystals for melt inclusion studies were extracted from crushed scoria lapilli by hand picking from the size fraction 0.5–1.0 mm. Crystals were embedded in epoxy and studied to identify primary, unfractured, glass inclusions. Doubly polished wafers were prepared for FTIR (fourier transform infrared Spectroscopy) analysis, whereas one-sided polished wafers were prepared for micro-Raman and electron microprobe analyses. FTIR and Raman analyses were performed before EMPA analysis. Further analyses of phenocrysts, melt inclusions and groundmass compositions were performed from thin sections made from single lapilli.

Analytical methods

Major-element compositions were analysed with a Philips PW2400 XRF spectrometer at the University of Lausanne (Switzerland). Loss on ignition (LOI) was determined using the gravimetric method corrected for FeO oxidation during ignition. Trace elements were analysed on fused glass discs by laser ablation inductively coupled plasma mass-spectrometry (LA-ICPMS) at the University of Lausanne (Switzerland), using a Lamba Physik GeoLas 200 M 193 nm ArF excimer laser ablation system interfaced to an Perkin Elmer ELAN 6100 DRC quadrupole ICPMS. Major-element compositions of phenocryst phases, olivine and plagioclase-hosted melt inclusions and matrix glass were measured using a JEOL JXA 8200 Superprobe at the University of Lausanne (Switzerland). For the olivinehosted glass inclusion analyses, we tested the mineralliquid equilibrium based on Fe-Mg exchange coefficients $[K_{\rm D}({\rm Fe-Mg})^{\rm olv-liq} = ({\rm FeO/MgO})^{\rm olv}/({\rm FeO/MgO})^{\rm liq}]$ in order to correct for olivine post-entrapment crystallization according to Toplis (2004) and Putirka (2008). To evaluate the $K_{\rm D}$, we calculated the FeO/Fe₂O₃ ratio based on the composition of the selected (i.e. $Cr_2O_3 >$ 13 wt.%, $Al_2O_3 > 4$ wt.%; $FeO_{tot} = 13-22$ wt.% $TiO_2 <$ 2.5 wt.%) spinel crystals enclosed in olivines, following Maurel and Maurel (1982) and Danyushevsky and Sobolev (1996) and checking spinel reliability following Kamenetski et al. (Kamenetsky et al. 2001). Calculations produced an average value for Fe²/Fe³ of 2.5 and 3.9 for LS and US magmas, respectively. Crystallization of plagioclase-hosted melt inclusions was tested calculating K_D (Ca-Na)^{Pl-liq}=(CaO/NaO)^{Pl}/ (CaO/NaO)^{liq}, following Hamada and Fuji (2007). Plagioclase composition is expressed as mole per cent of anortite, An=100×Ca/(Ca+Na+K); olivine composition as mole per cent of forsterite, Fo=100×Mg/(Mg+ Fe). Augite composition was expressed as a function of wollastonite content, $Wo = 100 \times Ca/(Mg + Fe + Ca)$, ferrosilite, Fs=100×Fe/(Mg+Fe+Ca) and enstatite En= 100×Mg/(Mg+Fe+Ca). Bulk rock Mg# was calculated as Mg/(Mg+Fe_{tot}), whereas in glasses it was calculated as Mg/(Mg+Fe²⁺).

Total dissolved water content of 37 olivine-hosted melt inclusions was also measured using confocal micro-Raman spectrometry at the Laboratory Pierre Süe of Saclay (France). Raman spectra were collected using a Renishaw inVia system connected to a Leica DMI microscope (×100 magnification) and equipped with a Peltier-cooled CCD detector. The concentrations of H- and C-bearing (dissolved as carbonate only in our samples) species were determined in nine olivine-hosted melt inclusions by FTIR at the same laboratory.

Results

Bulk rock chemistry

Chaimilla scoriae are basalts to basaltic andesites with SiO₂ ranging from 51 to 53 wt.% (Fig. 2). The compositional range is similar to some of the most mafic magmas erupted in prehistoric activity of Villarrica volcano (Vargas et al. 1989). It partially overlaps with the compositional range of lavas and scoria erupted in historical activity (Witter et al. 2004), but in general, it shows a wider variability towards more primitive compositions (Fig. 2). LS scoriae show slightly more-evolved compositions (average Mg# of 0.65) than upper sequence (US) scoriae (average Mg# of 0.68), but the most mafic sample (basalt with Mg# of 0.73) was collected from unit C (LS). The role of olivine, pyroxene and plagioclase crystallization (which are the main phenocrysts in all Villarrica volcanics) in the differentiation of the Villarrica magmas can be analysed by comparing their K₂O/Na₂O, CaO/Al₂O₃ and MgO contents. The magmas have very similar K₂O/Na₂O ratios (ranging from 0.25 to 0.18) and fit the general MgO-CaO/Al₂O₃ variation trend of Villarrica magmas (Fig. 3), suggesting that the magmas erupted at Villarrica were derived from the same parental magmas through different degrees of crystal fractionation of olivine followed by clinopyroxene and minor plagioclase. Trace-element concentrations confirm the subduction slab signature of these magmas and show similar patterns with respect to other magmas erupted from the Chilean Southern Volcanic Zone (Wehrmann et al. 2014), having MORBnormalized patterns with Pb, Ba U and Sr peaks, and troughs at Nb and Ta (Fig. 4). Chaimilla scoria has higher Ba/Nb (103-159) and Sr/Ce (25-35) ratios than do historical lava and scoria (59-100; 11-22, respectively) or those from older eruptions (62-106; 22-32) with similar compositions previously measured by Vargas et al. (1989) Witter et al. (2004) and Wehrmann et al. (2014). These data suggest a slightly larger slab fluid component in the Chaimilla magmas with respect to current erupted magma compositions (Fig. 5).

Petrography and mineralogy

All Chaimilla juvenile products are porphyritic, containing phenocrysts of plagioclase, olivine, clinopyroxene and rare chromian spinels: we infer a paragenesis similar to that for other major eruptions of Villarrica (Vargas et al. 1989; Lohmar et al. 2005, 2012). Glomero-porhyritic clusters are common in scoria from all units, and suggest the following phenocryst crystallization order: Mg-rich olivine, Mg-poorer olivine+An-rich plagioclase+pyroxene, An-poor plagioclase. Juvenile clasts from the US are characterized by lower phenocryst contents (11–18 vol.%, vesicle-free) than in LS clasts (20–25 vol.%, vesicle-free).

Plagioclase is the main phenocryst. It is more abundant in the lower sequence (LS) scoria (about 20 vol.%) than in the US scoria (about 12 vol.%). Crystals range from euhedral to subhedral in both LS and US (Fig. 6); they are typically 1.5 mm in size reaching a maximum size of 3 mm. The total compositional variability of plagioclase phenocrysts is large, ranging from An₆₇ to An₉₄. Crystals from units C and D display the largest compositional variability and are generally zoned; most crystals have An-rich (84 to 94 mol%) cores, and more-evolved compositions rimward (An₇₀₋₈₂). External rim compositions range from An₆₉ to An₇₃. These crystals are grouped as type 2 plagioclases (Figs. 6c and 7a). A subordinate fraction of crystals (type 1 plagioclase, Figs. 6d and 7a) is reversely zoned (An₇₃₋₈₂ cores and An₈₁₋₈₅ rims); some crystals from unit D show complex zoning (patchy or oscillatory) or resorbed rims. Phenocrysts from units G and H have more

Fig. 2 Variation diagrams showing SiO₂ compositions versus a MgO, b K₂O, c CaO and d Na2O of Chaimilla bulk scoria (open red squares LS samples; solid red squares US samples) compared with bulk rock composition from recent (solid black squares) and prehistoric (open black squares) activity (data from Witter et al. 2004; Vargas et al. 1989 respectively). Chaimilla whole-rock composition data from Costantini et al. (2011) have been also included. Crosses on the top right indicate the measurement error





Fig. 3 Variation diagrams showing compositional variations of **a** K₂O/Na₂O versus CaO/Al₂O₃ and **b** MgO versus CaO/Al₂O₃ for bulk rock compositions of magmas erupted by Villarrica volcano. *Symbols* and

uniform zoning patterns including An_{94-85} cores and An_{82-66} rims, similar to type 2 plagioclase compositions. In all units, plagioclase microlite compositions range from An_{69} to An_{61} ; similar compositions also characterize an external, up to 15-µm-thick rim of some phenocrysts (Fig.6c).

Olivine crystals makeup only 1-3 vol.% of the Chaimilla clasts and are usually euhedral to subhedral with sizes averaging between 1.0 and 1.5 mm in LS and ≤ 1 mm in the US (Figs. 6a, b and 7b). Larger crystals commonly contain glass inclusions. Most phenocrysts, microphenocrysts and groundmass crystals from unit C have homogeneous compositions ranging from Fo₇₂ to Fo₇₈, with the main mode at Fo₇₈. Only a small proportion of crystals display a magnesian core (Fo₈₀₋ ₈₂); a few phenocrysts, with resorbed rims, have Mg-rich compositions (Fo₈₁₋₈₃). All crystals measured from unit D scoria have homogeneous compositions (Fo75-78). Phenocrysts from units G and H are commonly normally zoned with cores ranging from Fo_{81} to Fo_{84} and 10–15-µm-thick rims with Fo_{75-79} compositions (Fig. 6a). Groundmass olivines have Fo-poor compositions (up to Fo_{69}). Clear bimodality in olivine compositions allows for the distinction of two olivine types: type 1 including crystals with Fo74-79 compositions and type 2 including crystals with Fo₈₀₋₈₅.

Clinopyroxene phenocrysts are less abundant than plagioclase and olivine (about 1 vol.%), with size ranging from 0.5 to 1.5 mm in both sequences. Generally, they have euhedral



sources as in Fig. 2. *Crosses* on the *right top* indicate the measurement error. The *shaded field* in \mathbf{b} underlines the main evolution trend

shapes (Fig. 6f). Pyroxenes only rarely contain glass inclusions, which have irregular shapes. Clinopyroxene phenocrysts display augitic composition with small heterogeneities within and among crystals (Wo₃₈₋₄₅, En₄₂₋₄₈, Fs₈₋₁₄, Fig.7c). As for olivine, unit D pyroxenes show the greatest uniformity with compositions of En₄₄₋₄₆. Crystals are generally weakly zoned, with variation in En and Wo molar units always <2 %.

Chromian spinels are rare and they usually appear enclosed within olivine crystals. Eleven chromian spinels embedded in olivine and two partially embedded within augite crystals were analysed in this study: they display variable compositions, with Cr₂O₃ ranging from 15 to 35 wt.%. The two spinels within augite crystals show high TiO₂ suggesting reequilibration with the melt. Spinels within olivine crystals have Cr# Cr/(Cr+Al) between 0.38 and 0.56. Crystals have similar Fe²⁺/Fe³⁺ ratios in lower units (C and D) and larger variability in upper units (G and H), which also appear correlated to variations of Cr#, Mg# and olivine composition. This suggests that spinels embedded in type 1 olivines in LS crystallized at homogeneous oxygen fugacity conditions (Maurel and Maurel 1982) and that the spinels from the US recorded distinct oxygen fugacity conditions experienced by the magma during rise and differentiation corresponding to the crystallization of the two olivine types.

The scoria groundmass is heterogeneous, varies from honey to light brown glass, is poorly to moderately crystallized



Fig. 4 N-MORB-normalized (after Sun and McDonough 1989) multielement variation diagram comparing Chaimilla scoria compositions with historical lavas and scoria (*yellow field*) and older deposits (*grey field*).

Data after Wehrmann et al. 2014; Witter et al. 2004 and Vargas et al. 1989 and this work



Fig. 5 Variation of Sr/Ce and Sm/Lu ratios in prehistoric, historic and Chaimilla scoria. Additional data from Wehrmann et al. (2014) and Vargas et al. (1989). *Symbols* as in Fig. 2. *Cross* on the *right top* indicates the measurement error

and is generally more crystalline in LS scoria than in US scoria. It contains microlites of plagioclase, clinopyroxene and oxides (Costantini et al. 2011). Enclaves of plagioclase-rich altered or fresh lava were also found in some scoria clasts and probably represent xenolith fragments embedded in magmas.

Textures of olivine and plagioclase-hosted glass inclusions

The shape of olivine-hosted glass inclusions ranges from sub-spherical to elliptical or sub-angular. Inclusions occur either dispersed in the core of crystals or concentrated along growth zones. Their largest axis varies in size from a few to 150 µm and they are generally more abundant in US than LS crystals. Most of the analysed glass inclusions are fully enclosed within the crystals, whereas only a few inclusions formed open embayments and only one was connected to the rim of the crystal by a thin neck (hourglass inclusion; Anderson 1991). Among the 47 analysed glass inclusions from both sequences, 15 contained small vesicles and two of them also contain an oxide crystal. The vapour bubbles are usually small, but they can represent a significant fraction of the melt inclusion, up to 30 vol.%.

Plagioclase-hosted inclusions are more abundant, and more than 10 inclusions can occur within the same crystal. Some plagioclases display sieve structures with abundant irregular, elliptical or spherical closed glass inclusions. Sieve textures and skeletal shapes are common within plagioclase with compositions An_{87-82} . Larger inclusions have generally irregular shapes, and display large vesicles whose volume can occupy up to 80–90 % of the entire inclusion; smaller inclusions have sub-angular shapes and only rarely contain bubbles. All plagioclase-hosted inclusions are microlite-free.

Olivine-hosted inclusion compositions

Compositions of olivine-hosted glass inclusions and matrix glasses are shown in Figs. 8, 9 and 10. The inclusions from the US and LS follow the same covariation trends for major oxides (Fig. 8) but are distinct for volatile species (Fig. 10).

Glass inclusions have basaltic to basaltic-andesite compositions, which overlap most of the Villarrica volcanics (Figs. 2 and 8) and mirror the compositional variability of the hosting crystals. Inclusions trapped in US olivine are more primitive than bulk rock compositions.

Parallel variation of CaO/Al₂O₃ and K₂O/Na₂O ratios suggests that the most primitive inclusions from US (CaO/Al₂O₃ between 0.71 and 0.60, MgO between 8 and 5 wt.%, Fig. 9) differentiated first by crystal fractionation of olivine, while crystallization of plagioclase became important during crystallization of more-evolved LS olivines, whose inclusions define a liquid line of descent marked by a linear increase in K₂O/Na₂O with decreasing CaO/Al₂O₃.

The water contents of the olivine-hosted glass inclusions are shown in Fig. 10a. Covariation trends of H_2O and major elements and similar Raman and FTIR results suggest that dissolved water content in MI was not significantly affected by bubble formation. Water content as measured with the Raman technique varies from 1.7 to 3.1 wt.%. The nine inclusions analysed with FTIR techniques had water contents ranging from 1.5 to 2.7 wt.%, whereas CO_2 concentrations were above the detection limit (50 ppm) in only three inclusions, and ranged from 70 to 320 ppm. Analyses made on the same inclusions with the two techniques gave concentrations that differ by 10 to 30 %, with the exception of one sample where the difference is more significant.

Inclusions hosted in US type 2 olivine crystals have water contents ranging from 2.4 to 3.1 wt.%, roughly decreasing with decreasing Fo content (Fig. 10a). H_2O/K_2O ratios are higher and more variable in US scoria than in LS scoria. When compared with CaO/Al₂O₃ ratios, inclusions define a degassing trend which we infer is associated with olivine and pyroxene, and then mostly with plagioclase, crystallization (Fig. 11a).

Measured S and Cl contents are up to 1400 and 180 ppm, respectively. S content is positively correlated with Mg in olivine, whereas Cl content shows very minor variations (Fig. 10b, c). S/Cl ratios decrease with decreasing water and MgO contents, suggesting that it was not significantly affected by boundary-layer effects (Baker 2008) and can be correlated with water degassing (Fig. 11b, c).

Plagioclase-hosted inclusion compositions

Plagioclase-hosted melt inclusions show lower variability than do olivine-hosted inclusions, fitting in the basalticandesite and andesite compositional fields, partially



Fig. 6 SEM backscattered images of mineral phases of Chaimilla scoriae. **a** Euhedral type 2 olivine with Fo-poor rim, unit H scoria. **b** Skeletal, homogeneous, type 1 olivine, unit C scoria. **c** Skeletal, type 1 plagioclase, reversely zoned, unit C scoria. **d** Euhedral, normally zoned

type 2 plagioclase with multiple melt inclusions, unit H scoria. **e** Glomeroporphyre with clinopyroxene, type 1 olivine and An-poor plagioclase (type 1), unit D scoria. **f** Euhedral clinopyroxene, unit D scoria. *Scale bars* are 100 μ m long

overlapping groundmass compositions (Fig. 8). No evident growth rims were detected around glass inclusions in any of the analysed plagioclase crystals. Uncorrected compositions show Kd with the host plagioclase crystals ranging from 0.05 to 0.35, increasing with Ab content of the host crystal, along the same trend for most inclusions (Fig. 12). As Kd plagioclase-liquidus varies with T, P and water content (Hamada and Fuji 2007), it is expected to change during magma crystallization and degassing. The Chaimilla variability trend however, compared with the empirical relationship suggested by Hamada and Fuji (2007), suggests that Kd was mainly controlled by thermal effects, and that most of the inclusions retained pristine compositions and were not significantly affected by post-entrapment modification. For this



Fig. 7 Compositional variation of Chaimilla phenocrysts. **a** Plagioclase compositions; *dark grey, light grey* and *black rectangles* refer to rim, core and microlite compositions, respectively. **b** Olivine compositions (from both single grains of olivine crystals prepared for Raman and FTIR

analysis and from olivines in thin sections) and c clinopyroxene compositions. Norm. frequency (normalized frequency)=number of crystals in each class divided by the total number of analysed crystals

reason, the few inclusions that do not fit the main trend have been disregarded from further calculations. MI compositions follow the same trends of the olivine-hosted inclusions analysed in the same units (Figs. 9 and 11).

S and Cl have maximum concentrations of 1200 and 900 ppm, respectively. Inclusions hosted in LS plagioclase follow similar trends to those of olivine-hosted inclusions (Fig. 8c). Most inclusions hosted in US, An-rich, plagioclase have very low S content and S/Cl ratios with respect to olivine- hosted inclusions from the same scoria (Fig. 11c). Plagioclase crystallization occurred during (and was likely promoted by) water degassing as suggested by the variability of S/Cl ratios, which is comparable with that in olivine-hosted melt inclusions.

Groundmass glasses

The groundmass of scoria clasts commonly contains numerous microlites. Analyses were limited to crystal-free glass areas, and only partially reflect real groundmass

Fig. 8 Variation diagrams showing compositional variations of SiO₂ versus a MgO, b K₂O, c CaO and d Na2O, for olivinehosted melt inclusions (black dots), plagioclase-hosted glass inclusions (green dots), bulk scoria (red squares) and matrix glasses (blue triangles). Open symbols LS samples, closed symbols US samples. The grey line on Fig. 6a show the liquid line of descent calculated with MELTS simulations. Crosses on the right top indicate the measurement error





Fig. 9 Variation of K_2O/Na_2O vs. CaO/Al₂O₃ in glass inclusions and glassy groundmass. *Symbols* as in Fig. 8. *Cross* at the *right top* indicate the measurement error

compositional variation. Compositions fit the liquid line of descent defined by the same covariation trend of glass inclusions and match the bulk compositions of the more-evolved magmas erupted at Villarrica in Holocene times. With respect to Villarrica historical magmas, Chamilla groundmass glasses differ in having lower CaO, Na₂O and Al₂O₃ content mostly





Fig. 11 a Variation diagram of H_2O vs. CaO/Al₂O₃. b S/Cl vs. water content. c S/Cl vs MgO content in olivine and plagioclase-hosted glass inclusions from Chaimilla scoria. *Symbols* as in fig. 8. Water data as measured with micro-Raman technique. *Crosses* at the *right top* indicate the measurement error



Fig. 10 Variation diagrams of a H_2O , b S and c Cl versus olivine composition in olivine-hosted glass inclusions. *Symbols* as in Fig. 8. Water data as measured with micro-Raman technique. *Crosses* on the *right top* indicate the measurement error

Fig. 12 Variation of Kd plagioclase-glass An content in plagioclasehosted glass inclusions. *Symbols* as in Fig. 8

in LS scoria, a depletion related to massive plagioclase crystallization in these samples (Fig. 8).

Interestingly, all groundmass compositions follow the same trend for CaO/Al₂O₃ and K₂O/Na₂O, excepting groundmass glasses from unit D; these show lower CaO/Al₂O₃ ratios that are compatible with their olivine-hosted glass inclusions (Fig. 9) and suggest distinct syneruptive crystallization conditions.

All groundmass glasses are volatile-poor, with maximum measured concentrations of 80 and 110 ppm for Cl and S, respectively, and concentrations below detection limits in about 10 % of the analyses.

Thermometry and barometry

The temperature range of magma crystallization was estimated based on olivine and glass inclusion compositions according to Putirka et al. (2007); we obtained temperature values that range between 1046 °C (Fo₇₇, 1.7 water wt.%) and 1113 °C (Fo₇₈, 2.73 water wt.%) in type 1 (LS) olivines and between 1099±10 (Fo₈₃, 2.29 water wt.%) and 1132±10 °C (Fo₈₅, 3.00 water wt.%) in type 2 (US) olivines. Analytical uncertainties led to an error of about 10 °C on the temperature estimations.

Minimum saturation pressures were estimated for the inclusions analysed with FTIR techniques by applying the Papale et al. (2006) model at the temperatures calculated as described above. Pressures obtained range from 37 to 91 MPa, with the highest pressures obtained from the LS melt inclusions that contain the highest CO_2 . The minimum pressures required to dissolve the water measured with the micro-Raman technique (and assuming complete exsolution of CO₂) vary from 28 to 50 MPa in LS and from 33 to 73 MPa in the US. Since H₂O solubility decreases with increasing dissolved CO₂, the saturation pressures as calculated from the micro-Raman data represent minimum values. The possible effect of CO_2 (and the uncertainty in pressure estimation) can be evaluated by comparing the results and the H₂O-CO₂ solubility curves calculated both for typical type 1 and type 2 inclusion glasses, as computed with the Papale et al. (2006) model (Fig. 13).

Discussion

Modeling magma evolution

We applied the software MELTS (Ghiorso and Sack 1995; Asimow and Ghiorso 1998) to reproduce the liquid line of descent and obtain further insights into the evolution conditions of the Chaimilla magmas. We have simulated decompression and cooling of magmas with compositions corresponding to the most primitive glass (originally melt) inclusion from the US (i.e. V112Bol2). We have simulated the effect of variable oxygen fugacities and total (lithostatic)



Fig. 13 Solubility of water and carbon dioxide for the average composition of olivine-hosted glass inclusion in the Chaimilla scoriae compared with water content measured with micro-Raman techniques (*full circles* US olivines and *open circles* LS olivines) and water and carbon dioxide content of selected inclusion as measured with FTIR techniques (*open red circles*). *Isobaric grey lines* correspond to the solubility of the inclusions), and *isobaric black lines* correspond to the solubility of the inclusion within in type 1 olivines (average composition of LS inclusions)

pressures; both equilibrium and fractional crystallization processes were simulated using an initial water content of 3 wt.%. In all cases, MELTS could not simulate the crystallization of olivine unless pyroxene crystallization was blocked; this is because MELTS tends to overestimate the stability of pyroxene at the expense of olivine (Villiger et al. 2004). In any case, it was able to predict liquid lines of descent that overlap with the observed glass-inclusion compositions in both the LS and US. In all experiments, plagioclase was the most abundant mineral phase crystallizing from the magma. The liquid line of descent for inclusions was best reproduced by crystallization of olivine and plagioclase through cooling from liquidus compositions (around 1200 °C, depending on water content) down to 1020 °C and depressurization from 120 to 62 MPa with oxygen fugacity constrained by the OFM+1 curve (Fig. 8a). These results are in accordance with the oxygen fugacity estimated based on the olivine-spinel compositions, but they predict significantly higher pressures (and lower water solubility in the magmas) compared with the maximum solubility pressures obtained applying the Papale et al. (2006) model.

Compositions of olivine crystals obtained from the simulation, moreover, match well the real data for dissolved water content of 2 wt.% or lower. For higher contents, it predicts compositions Fo-poorer than the measured ones, probably due to the low water solubility allowed in these magmas by the MELTS libraries. For the same conditions, the model also predicts crystallization of An_{79-84} plagioclase during olivine crystallization and with more-evolved compositions (up to An_{75}) in the final stages. When compared with the compositional variability of the melt inclusion data and plagioclase prenocrysts are compatible with crystallization from a magma with compositions corresponding to the less-evolved and water-richer (>2 wt.%) glass inclusions hosted within Fo_{82-83} olivines.

Chaimilla magmas: degassing and evolution

The petrologic features of the analysed scoriae show systematic variations among the magmas erupted during phases C and D (LS) and G and H (US) of the Chaimilla eruption, suggesting that they were fed by distinct magma batches.

Both plagioclase and olivine phenocrysts show a definite bimodality. Two components occur in different proportions in scoria from each unit: unit C has both type 1 and 2 olivines (85 and 15 % of our analyses, respectively) and type 1 and 2 plagioclases (20–80 % of our analyses, respectively). Unit D contains only type 1 olivines and both plagioclase types (occurring in equal proportions in our analyses). Units G and H have both type 1 and 2 olivine crystals as microphenocrysts and phenocrysts, respectively (occurring in equal proportions in our analyses) and only type 2 plagioclase crystals.

Melt inclusions hosted in type 2 (Mg-rich) olivine suggest that the crystals formed in a magma evolving by fractional crystallization of pyroxene and only minor olivine, associated with gradual magma degassing (from 3.1 to 2.4 wt.% of water, 1400 to 300 ppm of S and 2000 to 200 ppm of Cl). Inclusions hosted in type 1 (Mg-poorer) olivines show a compositional variability recording more extensive plagioclase and pyroxene (and minor olivine) crystallization and associated with more advanced degassing (from 2.5 to 1.0 wt.% of water, 1000-250 and 1800-200 ppm of S and Cl, respectively). Inclusions hosted in An-rich cores of type 2 plagioclases have compositions matching the variability of olivine-hosted melt inclusions and higher K₂O/Na₂O ratios in rims, whereas inclusions in type 1 plagioclase (reversely zoned) show more homogenous compositions, with almost constant CaO/Al2O3 ratios of 0.5 and K₂O/Na₂O ratios between 0.2 to 0.3.

Geochemical trends recorded by Chaimilla glass inclusions and matrices can be explained by multiphase magma differentiation (through crystallization, cooling and degassing) associated with magma mixing. Two magma end members can be identified: first is a type 1 magma, which evolved from a melt with the same composition as the bulk Chaimilla scoria and was stored in a shallow reservoir (at around 2 km depth) where it was subjected to degassing and An-poor (type 1 plagioclase cores) plagioclase crystallization, associated with minor pyroxene and olivine (type 1) crystallization.

The second end member (type 2 magma) evolved from a basalt with compositions corresponding to the more primitive inclusions hosted in type 2 olivines. This magma mostly evolved through fractional crystallization of olivine and pyroxene accompanied by S and water (from 3 to 2 wt.%) degassing during rise. A minor amount of An-rich (An₉₄₋₈₄)

plagioclase crystallized, probably under undersaturated conditions.

The mineral assemblage in LS scoria, including type 1 and 2 olivines and both normally and reversely zoned (type 1 and 2) plagioclase, suggests mixing between the two magmas before eruption, probably due to the entrance of a small batch of crystal-poorer, water-richer, type 2 magma into the shallow reservoir where type 1 magma was crystallizing. Magma mixing caused a temperature increase in the reservoir with partial dissolution of An-poor plagioclases and further magma crystallization with formation of the An-rich rims in type 1 plagioclases. The deposits of the basal sequence reflect activation of the shallow system, causing vent opening and outgassing followed by vulcanian activity. These events further destabilized the reservoir and triggered the main explosive phase depositing the lower sequence, with almost complete emptying of the reservoir during the phase of eruption that deposited the middle sequence. Finally, a larger batch of type 2 magma fed the most explosive phase of the eruption, depositing the upper sequence and culminating with the repeated column collapses generating the PDC deposits of unit H.

Comparison with recent activity of Villarrica volcano

At present, Villarrica is an open-conduit volcano that is persistently outgassing. The recent activity is characterized by a narrower compositional range and more uniform phenocryst composition (and content) than that which produced the Chaimilla products (Witter et al. 2004 and Gurioli et al. 2008). Olivine crystals are smaller in scoriae of the recent activity (reaching a maximum of 0.5 mm) and Fo content corresponds to type 2 (Fo-poor) olivines in Chaimilla scoria (Fo₇₆₋₇₉). Chaimilla plagioclase crystals are larger and significantly more Ca-rich than the ones currently erupted (An₅₈₋₆₀ in year 2000 scoria), but have compositions similar to the most An-rich plagioclase (An₈₀₋₈₅) embedded in a lava flow emitted in 1984. No pyroxenes have been found in the recent products (Witter et al. 2004). Olivine-hosted and plagioclase-hosted glass inclusions from reticulites of the year 2000 explosions have similar Mg# as do some of the inclusions hosted in type 2 (Fo₇₆₋₇₇) olivines in the Chaimilla scoria, but with significantly less water (no more than 0.28 wt.%) than in the Chaimilla magmas. Interestingly, the maximum water content of glass inclusions hosted in type 2 (Fo-poor) olivines is compatible with the initial water content predicted by Witter et al. (2004) for the recent magmas (i.e. 2 wt.%), based on very similar olivine compositions.

Our data suggest very different crystallization and degassing conditions for the current activity of the volcano compared to that represented by products of the Chaimilla eruption, but similar parental magmas. Recent magmas, in fact, experience equilibrium crystallization and degassing in a very shallow reservoir (<1 km), at much lower P_{H2O} (17 MPa, as calculated by Witter et al. 2004), and a stable magmatic system is maintained for long periods by convection driven by outgassing (Witter et al. 2004). This was also suggested by geophysical and geochemical monitoring observations (Palma et al. 2008; Ripepe et al. 2010), by analogy with other open-conduit volcanoes (i.e. Stromboli: Landi et al. 2006; Mt Erebus: Caldwell and Kyle 1994). The Chaimilla magmas, instead, experienced crystallization during rise from greater depth (at pressures up to 120 MPa) and with higher dissolved water concentrations. Scoriae do not preserve direct information on the amount of volatiles still dissolved in the magma just before eruption, because groundmass glass is usually completely degassed. Crystal, glass and scoria composition are, however, reliable petrologic indicators for degassing processes. MELTS simulations show that the Na-richer plagioclase phenocryst rims are in equilibrium with magmas containing about 1.8 wt.% water and imply that a high fraction of the original water was still within the system just before syneruptive crystallization of microlites in the groundmass and possibly during fragmentation of the microlite-poor US magma. We suggest that these differences in magma ascent and degassing history played a primary role in causing the high explosivity of the Chaimilla eruption, and confirm the primary role of volatile content and magma ascent rate in controlling eruptive dynamics. Finally, we note that slightly different Ba/Nb, Sr/Ce ratios in the Chaimilla magma could be indicative of a larger initial water content in the source magmas with respect to other Villarrica magmas. This hypothesis should be confirmed or rejected based on more-systematic trace-element studies on historical lava and tephra.

Conclusions

The Chaimilla eruption was the largest magnitude event produced by the Villarrica summit cone after the caldera-forming Pucón ignimbrite eruption (3.7 ka) and was fed by inhomogeneous basaltic andesite magmas. The eruption dynamics were complex and produced four main sequences of events having different intensities. Our data show that the main changes in the eruption dynamics were associated with changes in the composition and petrology of the erupted magmas. The magmas erupted during the main phases (C-D and G-H) are characterized by petrological differences related to slightly different initial compositions, different degrees of crystallization, equilibrium conditions and magma ascent rates. The eruption was probably triggered by a mafic and volatile-rich input (with water up to 3.1 wt.%) within a shallow magmatic reservoir (at about 1 km below the volcano) containing a more crystalline and degassed magma (about 1 wt.% water), which fed almost all of the (weaker) eruptive phases that produced the lower and middle sequences. The magma erupted during the more violent late phase that deposited the upper sequence, instead, experienced minor crystallization at shallow levels and less-efficient degassing, which are indicative of a rapid ascent of a larger magma batch from depths of about 3 km below the volcano. This hypothesis explains the increased explosivity of this phase of the eruption, characterized by higher intensity and repeated column collapses.

The Chaimilla scoriae also have bulk compositions similar to that of magma currently erupted at Villarrica, but there are marked differences in crystallinity, storage pressures and volatile contents (Witter et al. 2004). The current magmas have a much lower dissolved volatile content, indicating very shallow and stable crystallization and outgassing in open-vent conditions. We suggest that open-conduit conditions and low feeding rates are key factors in limiting the explosivity of the volcano in its current state, because they allow efficient outgassing and circulation of magma, favouring equilibrium crystallization and homogeneous properties of the magma body. On the other hand, the greater explosivity of the Chaimilla eruption was probably controlled by the storage of a significant volume of magma with intermediate volatile contents in a well-developed shallow reservoir and triggered by deep, hot and volatile-rich magma recharges.

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