Thermo-barometric constraints on the Mt. Etna 2015 eruptive event

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Abstract
The petrological study of volcanic products emitted during the paroxysmal events of December 2015 from the summit craters of Mount Etna allow us to constrain T-P-XH2O phase stability, crystallization conditions, and mixing processes along the main open-conduit feeding system. In this study, we discuss new geochemical, thermo-barometric data and related Rhyolite-MELTS modelling of the eruptive activity that involved the concomitant activation of all summit craters. The results, in comparison with the previous paroxysmal events of the 2011–2012, reinforce the model of a vertically extended feeding system and highlight that the activity at the New South-East Crater was fed by magma residing at a significantly shallower depth with respect to the Central Craters (CC) and North-East Crater (NEC), even if all conduits were fed by a common deep (P = 530–440 MPa) basic magmatic input. Plagioclase dissolution, resorption textures, and the Rhyolite-MELTS stability model corroborate its dependence on H2O content; thus, suggesting that further studies on the effect that flushing from fluids with different H2O/CO2 ratio are needed to understand the eruption-triggering mechanisms for high energetic strombolian paroxysmal episodes.

Keywords Open conduit volcanic feeding system · Olivine and clinopyroxene thermo-barometry · Chemical–physical crystallisation conditions · Chemical and textural zoning of phenocrysts · Rhyolite-MELTS modeling

Introduction
Mt. Etna’s persistent activity represents an almost unique opportunity to investigate the magmatic processes occurring prior and during the eruption of a complex stratovolcano, characterized by a multifaceted feeding system. Decades of petrological, geochemical, geophysical and tectonic studies on Mt. Etna have contributed to the characterization and quantification of the tectono-magmatic processes from the mantle source to eruptive activity.

The current Mt. Etna plumbing structure consists of three main feeding systems, resulting from the transtensional tectonic regional setting: (1) the central open-conduits (Corsaro and Pompilio 2004, 2009a, b; Giacomoni et al. 2018); (2) the lateral rift-related, namely the S Rift, the NE Rift and the W-Rift (Ferlito et al. 2009; Giacomoni et al. 2012), and (3) the eccentric (Rittmann 1965; Kieffer 1975; Armienti et al. 1988).

The central conduits culminate with five summit craters: Central Craters (CC) consisting of Bocca Nuova (BN) and Voragine (VOR); North-East Crater (NEC), South-East Crater (SEC) and New South-East Crater (NSEC). The height of the magma column inside the central conduits changes constantly across the H2O saturation depth, resulting in an efficient degassing of the magma, promoting the crystallization and fractionation of abundant plagioclase (Mollo et al. 2015; Giacomoni et al. 2014, 2018). Volcanic products commonly erupted at summit craters are thus highly porphyritic with a microcrystalline matrix, sometimes mingled with glassy, almost aphyric melts indicating episodes of basic magmatic inputs from the deeper portion of the plumbing system.

The activity of lateral rift-related systems is more discontinuous and strictly linked to the regional transtensive
Geological and volcanological background

Mt Etna is a 3340 m a.s.l. stratovolcano located on the Ionic coast of Eastern Sicily. The volcanic edifice has been built up on the thrust between the Appennine-Maghrebian chain and the undeformed Hyblean foreland (Lentini et al. 2006). The NNW–SSE transtensive tectonic structure of the Malta Escarpment separates the thick continental lithosphere of Eastern Sicily from the thinned Ionian oceanic crust, which represents the remainder of the Jurassic Tethys ocean (Hirn et al. 1997; Micallef et al. 2018). The entire volcanic edifice is crossed by normal faults striking NNW–SSE and NNE–SSW, with the four summit Central Craters currently located at their intersection. On the volcano’s flanks, the surface expressions of these tectonic structures are the N-S Rift and the NE Rift systems (Kieffer 1975; Monaco et al. 1997).

Magmatic activity started ~500 ka ago with the emission of olivine tholeiitic basalts, accompanied by occasional pigeonite-bearing tholeiitic basalts from fissural submarine and subaerial vents (Gillot et al. 1994; Tanguy et al. 1997). The early submarine volcanic events were followed by subaerial eruptions, together with regional uplift that occurred ~300 ka ago (Branca and Del Carlo 2004). The

regime that affects the eastern flank of the volcanic edifice (Monaco et al. 1997, 2005). The 2001 and 2002–2003 lateral eruptions attracted the attention of the scientific community, highlighting that, during an eruptive event, magma can migrate from central conduits towards the rift system (Andronico et al. 2005) or alternatively intrude independently into the lateral rift systems, eventually being stored in shallow, short-lived magmatic batches (Ferlito et al. 2009; Giacomoni et al. 2012, 2014). Mixing episodes between magma batches stored along the rift structures with incoming deeper basic melts can eventually trigger highly energetic eruptions (Viccaro et al. 2006).

The eccentric system is composed of several disperse monogenetic scoria cones and associated lava flows which are apparently not connected with the central or lateral feeding systems. A petrological study on the most recent eccentric event, the 1974 eruption of Mt. De Fiore (Corsaro et al. 2009b), highlighted that eccentric eruptions can be defined as “deep-dyke fed” (DDF), differing from central and lateral eruptions which are instead fed by magma residing at shallower depths and drained from the central conduits; thus, defined as “central conduit fed” (CCF). This feature, together with notably more rapid ascent rates, constrain the physical crystallization of magma, significantly reducing the stability field of plagioclase and resulting in the emission of mostly olivine-phyric products with scarce clinopyroxene (Metrich and Rutherford 1998).

Recent geophysical, geochemical, and petrological studies (Di Grazia et al. 2006; Andronico et al. 2008; Aloisi et al. 2009; Ferlito et al. 2009; Corsaro et al. 2009a; Mollo et al. 2015; Putanè et al. 2006, 2013; Kahl et al. 2011, 2013) of eruptive events occurring from 2001 to 2014 agree that the geometry of feeding conduits and changes in the physical–chemical (P–T–O2–XH2O–XCO2) should be considered as the main factors governing the eruptive style rather than the magmas, which are characterized by fairly homogeneous chemical compositions.

Following the 2001 and 2002–2003 lateral events, eruptive activity since 2006 has been restricted to the five summit craters. The Voragine (VOR) is the oldest among the summit craters and has been represented in historical iconography since the fourteenth century (Cannata et al. 2018), while the cone shaped morphology of the South-East Crater (SEC) and Bocca Nuova (BN) formed in 1971 and 1969, respectively. The New South-East Crater (NSEC) is the youngest, having formed in 2006 from explosive activity originating in a pit crater, in turn created by flank collapse of the SEC in 2006 (Neri 2010; Ferlito et al. 2010).

Since magma can reside in central conduits long enough to extensively degas, most eruptions from the central craters or along fissures, which drain central conduits, are commonly effusive with weak explosive (i.e. Strombolian) activity. This type of activity, accompanied by sporadic lava fountaining, characterized the 2004–2010 period. The 2011–2013 eruptive period (Mollo et al. 2017; Giacomoni et al. 2018) represented an interesting and dramatic shift in eruptive style, whereby activity became significantly more energetic and was characterized by brief (few hours to days), strong Strombolian activity with episodes of intense lava fountaining from the NSEC, during which any remarkable seismicity was absent, defined as “paroxysmal episodes” (Behncke et al. 2014; De Beni et al. 2005).

Within this context of frequent and intense eruptive events from the NSEC, from 3 to 8 December 2015, volcanic activity affected all summit craters concomitantly: Central Craters (CC), North-East Crater (NEC) and New South-East crater (NSEC). This unusual event represented a drastic change from previous activity restricted to the NSEC, thus the December 2015 eruptive sequence provides an almost unique opportunity to study the intensive parameters affecting magmatic crystallization in the shallowest portion of the plumbing of Mt. Etna’s central conduit system and relationships with the summit craters.

This study focuses on selected scoria samples from the explosive activity in December 2015; a detailed petrological, textural, thermo-barometric, hygrometric characterization was carried out to highlight any differences in the crystallization conditions of the magmas erupted from the summit craters. Results are then discussed in a comprehensive model, including data from the 2011–2012 paroxysmal events.
composition of the erupted products shifted from tholeiitic to transitional to Na-alkaline (~200 ka; Tanguy et al. 1997; Casetta et al. 2019). The building of the central-conduit edifice started ~130 ka ago, resulting in a sequence of several distinct volcanic centers characterized by the alternation of effusive and explosive eruptions frequently associated with caldera collapses (De Beni et al. 2005). The “Recent Mongibello” activity began ~15 ka ago, being predominantly effusive with lava emissions and Strombolian eruptions from the summit craters and/or from eccentric parasitic cones forming on the volcano’s flanks (Branca and Del Carlo 2004).

The December 2015 eruptive activity

The temporal reconstruction of the eruptive event occurring between 3 and 8 December was possible thanks to the continuous monitoring and routine volcanological reports from the Istituto Nazionale di Geofisica e Vulcanologia-Osservatorio Etno (INGV-IE). On 3 December (h 2:32–3:08 UTC), mild Strombolian activity began in the Voragine central crater (VOR) but rapidly increased to a violent yet brief lava fountain reaching over 1000 m in height, in turn generating a sustained eruptive column up to 12 km with ash fallout dispersed 30 km southwest from the crater (Corsaro et al. 2017). Most of the activity remained limited to the Central Crater rim, enabling the onset of lava flows. This powerful paroxysmal event lasted only a few hours, but weak ash emissions from the VOR and NSEC continued for the entire day of 3 December.

On 4 December, explosive activity resumed with two short-lasting but intense lava fountaining events from the VOR crater at 9:07–10:06 UTC and at 20.35–21:14 UTC, respectively.

Both events started with mild Strombolian activity, which rapidly shifted to strong fountaining accompanied by an eruptive column 11 ± 2 km in height. Abundant ash emissions were observed at the NSEC, simultaneous with the 20:35 VOR event.

On 5 December (14.46–15.35 UTC), the intensity and frequency of explosive activity at the VOR increased and resulted in a fourth paroxysmal episode of lava fountaining.

On 6 December, the eruptive activity shifted from the VOR to the NSEC where ash emissions accompanied a scoriaceous lava flow directed towards the Valle Del Bove; at the same time, weak ash emissions were observed at the NEC crater. Later during the day, the explosive activity became strictly Strombolian while effusive activity continued to be fed by three distinct vents, flowing eastward and south-eastward into the Valle del Bove depression and reaching Mt. Centenari (Corsaro et al. 2017).

The activity at the NSEC ended on 8 December concomitantly with the onset of weak Strombolian activity at the NEC, which lasted for the next 48 h and gradually decreased and ceased on 11 December.

Sampling and analytical techniques

Six rock samples were collected from the explosive activity of the Voragine Central Crater (CC) of 5 December, and from the New South-East Crater (NSEC) and North-East Crater (NEC) emitted on 6 December.

Whole-rock geochemical analyses were carried out at the laboratory of the Department of Physics and Earth Sciences, University of Ferrara (Italy) using powdered aliquots of lava and tephra. Major and some trace element (Ba, Cr, Sc, V) concentrations were measured by X-ray fluorescence (Thermo ARL Advant XP) and corrected for matrix effects using the method of Lachance and Traill (1966). Loss on ignition (LOI) was determined by gravimetric method, assuming Fe₂O₃ as 15% FeO. Zr, Rb, Nd, Ce, Nb, Th, U, and REE concentrations measured by inductively coupled plasma mass spectrometry (ICP-MS) on a VG Elemental Plasma Quad 2Plus instrument.

The data reproducibility ranges between 0.9 and 7.9%; from a conservative point of view, an accuracy of 10% has been determined and a detection limit of 10 ppb for Th, U and REE.

Microchemical analyses and textural images were collected at the HP-HT Laboratory of Experimental Volcanology and Geophysics of the Istituto Nazionale di Geofisica e Vulcanologia (INGV) in Rome (Italy) by means of a JEOL-JXA8200 electron probe microanalyzer (EPMA) equipped with five spectrometers. Data were collected on core to rim profiles of phenocrysts, and coexisting glasses located a few microns next to crystal edges. Analytical conditions were set at an accelerating voltage of 15 kV and a beam current of 10 nA. Phenocrysts were analyzed with a beam size of 1 μm with a counting time of 20 and 10 s on peaks and background, respectively. For glasses, a slightly defocused electron beam with a size of 3 μm was used with a counting time of 5 s on background and 15 s on the peak. The following standards have been adopted for the various chemical elements: jadeite (Si and Na), corundum (Al), forsterite (Mg), andradite (Fe), rutile (Ti), orthoclase (K), barite (Ba), apatite (P), spessartine (Mn) and chromite (Cr). Sodium and potassium were analyzed first to mitigate alkali migration effects. The precision of the microprobe was measured through the analysis of well-characterized synthetic oxides and minerals. Data quality was ensured by analyzing the test materials as unknowns according to Iezzi et al. (2014). Based on counting statistics, analytical uncertainties relative to their reported concentrations indicate that the accuracy was better than 5%. Images were collected using the backscattered electron (BSE) mode of a field emission gun-scanning
electron microscope (FE-SEM) JEOL 6500F equipped with an energy dispersive spectrometer (EDS) detector.

Petrography and mineral chemistry

Petrographic features summarized in Table 1 show that all samples have a porphyritic index (PI), defined as the percentage of phenocrysts with respect to the whole thin section area, ranging from 15% at NEC to 40% NSEC. Despite samples showing similar mineral paragenesis consisting of olivine (Ol), clinopyroxene (Cpx), plagioclase (Plg) and Ti-magnetite (Ti-Mt), the relative percentage of phenocrysts and vesicles varies significantly.

Ol content ranges between 10 and 20% in samples from NSEC and 20% in CC and NEC. Cpx content ranges constantly between 20 and 35% in NSEC and from 30 to 50% in CC and NEC; Plg content varies between 25 and 35% in CC, up to 55% in NEC and varies from 50 to 55% in NSEC. Ti-Magnetite phenocrysts occur predominantly in the groundmass or enclosed in Ol and Cpx; its content is almost constant between samples (~ 5%). Glomerophyric assemblages of Ol (5%), Cpx (75–90%) and Plg (5–15%) are common in all studied samples.

Olivine phenocrysts are usually euhedral, but few larger crystals (> 4–5 mm) with dissolved irregular rims are present in all samples (Fig. 1a–c). Compositions range from Fo56.3 to Fo82.4 in NSEC, from Fo71.7 to Fo78.9 in NEC and from Fo70.0 to Fo81.6 in CC samples.

Clinopyroxene is mostly euhedral in shape and ranges in size from 50 μm to 4 mm (Fig. 1d–f); all analysed Cpx are diopсидic augites in composition (Morimoto 1988) with Mg# content varying from 79.1 to 67.7 in NSEC, 77.8–71.7 in NEC and 81.5–69.7 in CC samples. Most Cpx phenocrysts show direct oscillatory zoning characterized by repeated micrometric dissolution surfaces. Reverse zoned Cpx, characterized by a dissolved core (74–75 Mg#) enveloped by 76–78 Mg# rim occur in both NEC and CC samples (Fig. 1).

| Sample     | Porphyritic index (%) | Percentage of vesicles (%) | Relative percentage of phenocrysts (%) | Type of matrix |
|------------|-----------------------|----------------------------|----------------------------------------|----------------|
| CC_1       | 15%                   | 10%                        | 20% 50% 25% 5%                         | Hyalopilitic   |
| CC-2       | 20%                   | 15%                        | 20% 40% 35% 5%                         | Hyalopilitic   |
| NEC        | 15%                   | 8%                         | 20% 30% 45% 5%                         | Glassy        |
| NSEC_1     | 25%                   | 23%                        | 20% 20% 55% 5%                         | Hyalopilitic   |
| NSEC_2     | 40%                   | 20%                        | 10% 35% 50% 5%                         | Hyalopilitic   |
| NSEC_3     | 25%                   | 25%                        | 20% 20% 55% 5%                         | Hyalopilitic   |

Fig. 1 Back scattered (SEM) microphotographs of olivine (a–c), clinopyroxene (d–f) and plagioclase (g–i) in samples from Central Craters (CC), New South-East Crater (NSEC) and North-East Crater (NEC)
Plagioclase phenocrysts occur in two well-distinguished dimension populations. Large phenocrysts, often zoned and heavily texturized (Fig. 1g–i), can reach up to 3 mm in size along the C-axis. On the contrary, groundmass phenocrysts (20–30 µm) are usually euhedral and homogenous in composition. Larger phenocrysts present complex dissolution-regrowth textures at cores or at rim (Fig. 2) always associated with significant compositional changes in anorthite (An) content.

Following the classification of textures proposed by Giacomoni et al. (2014); three main textures have been recognized in studied samples: i) oscillatory zoned; ii) dissolved core; iii) resorbed dusty rim.

Oscillatory zoned plagioclases are rare in large phenocrysts. They are mainly found in NSEC samples and are characterized by a low frequency oscillation in An content varying from 70 to 82% (Fig. 2e, f). Plagioclase with rounded dissolved cores (Fig. 2b, c) usually show a high An content (An88-An70) enveloped by a oscillatory An50-60 regrowth. However, plagioclase with dusty resorbed rims (Fig. 2a, d) are commonly characterized by a low An oscillatory core (An60-70) enveloped by an An70-85 overgrowth made of recrystallized glass pockets and glassy channels. All three textural types are characterized by an outermost thin (up to 50 µm) overgrowth with An60-50 composition.

Opaque phenocrysts have Ti-magnetitic composition (Usp14–47 for CC, Usp29-42 for NSEC and Usp28-46 for NEC).

Fig. 2 Back scattered (SEM) microphotographs and anorthite content core-rim electron microprobe (EMP) profiles of studied plagioclases from Central Craters (CC; a, b); North-East Crater (NEC, c, d) and New South-East Crater (NSEC, e, f). The determined dissolved H$_2$O content in the equilibrium melt has been obtained by means of the Waters and Lange (2015) hygrometer. a dusty rim plagioclase associated with a An and H$_2$O increase; b rounded dissolved core followed by a An and H$_2$O decrease; c oscillatory zoned plagioclase; d dusty rim with a An and H$_2$O increase; e and f oscillatory zoned plagioclases from NSEC.
Whole-rock geochemistry

Whole-rock major and trace element compositions of the studied samples are summarized in Table 2. All samples are alkaline basalts (Fig. 3, Le Maitre 2002) with potassic affinity (trachybasalts). Their SiO$_2$ content varies from 47.77 to 49.66 wt%, at Mg# (calculated as MgO/[MgO + FeO] mol × 100, with Fe$_2$O$_3$/FeO ratio of 0.15) values ranging from 45.69 to 50.56, confirming that they lie in a restricted differentiation range, and do not significantly differ from other recently erupted products from the summit Craters (Giacomoni et al. 2018). Harker’s diagrams of SiO$_2$ versus CaO, TiO$_2$, FeO and CaO/Al$_2$O$_3$

| Sample | CC1 | CC2 | NEC | NSEC1 | NSEC2 | NSEC3 |
|--------|-----|-----|-----|-------|-------|-------|
| SiO$_2$ | 50.10 | 49.14 | 48.67 | 48.26 | 48.52 | 48.59 |
| TiO$_2$ | 1.58 | 1.72 | 1.78 | 1.66 | 1.65 | 1.63 |
| Al$_2$O$_3$ | 18.13 | 17.32 | 17.15 | 17.25 | 17.85 | 17.85 |
| FeO | 7.95 | 8.82 | 9.36 | 9.31 | 9.06 | 8.89 |
| Fe$_2$O$_3$ | 1.19 | 1.32 | 1.40 | 1.40 | 1.36 | 1.33 |
| MnO | 0.18 | 0.18 | 0.19 | 0.19 | 0.17 | 0.17 |
| MgO | 4.04 | 5.06 | 5.05 | 4.75 | 4.44 | 4.19 |
| CaO | 9.95 | 10.17 | 10.28 | 11.11 | 10.90 | 10.90 |
| Na$_2$O | 4.32 | 3.75 | 3.68 | 4.05 | 3.94 | 4.31 |
| K$_2$O | 2.34 | 2.05 | 2.07 | 2.03 | 2.06 | 2.08 |
| P$_2$O$_5$ | 0.21 | 0.46 | 0.37 | 0.00 | 0.06 | 0.03 |
| LOI | 0.13 | 0.32 | 0.25 | 0.12 | 0.18 | 0.13 |
| Total | 100 | 100 | 100 | 100 | 100 | 100 |
| Mg# | 47.50 | 50.56 | 49.04 | 47.64 | 46.61 | 45.69 |
| FeOt | 9.03 | 10.02 | 10.62 | 10.56 | 10.28 | 10.09 |
| Rb (ICP-MS) | 51.1 | 46.6 | 47.7 | 44.3 | 46.5 | 45.6 |
| V (XRF) | 263 | 276 | 300 | 314 | 299 | 303 |
| Sc (XRF) | 21.7 | 22.4 | 27.5 | 27.5 | 25.5 | 24.8 |
| Cr (XRF) | 29.9 | 34.4 | 27.1 | 46.5 | 38.8 | 40.7 |
| Sr (ICP-MS) | 956 | 831 | 988 | 994 | 1003 | 979 |
| Th (XRF) | 12.6 | 10.2 | 13.5 | 9.2 | 12.4 | 12.3 |
| U (ICP-MS) | 2.45 | 2.18 | 2.10 | 1.90 | 2.00 | 1.96 |
| Nb (ICP-MS) | 72.5 | 64.4 | 64.1 | 56.2 | 59.5 | 59.6 |
| Ba (ICP-MS) | 718 | 644 | 651 | 623 | 645 | 659 |
| Ta (ICP-MS) | 2.34 | 2.09 | 2.10 | 1.91 | 2.01 | 1.99 |
| La (ICP-MS) | 42.6 | 39.5 | 42.8 | 43.3 | 45.4 | 42.7 |
| Ce (ICP-MS) | 101 | 92.1 | 99.3 | 91.8 | 93.7 | 92.5 |
| Nd (ICP-MS) | 35.5 | 33.1 | 37.0 | 38.1 | 39.5 | 37.3 |
| Pr (ICP-MS) | 8.72 | 8.17 | 9.01 | 9.18 | 9.49 | 8.99 |
| Sm (ICP-MS) | 6.04 | 5.58 | 6.32 | 6.74 | 7.01 | 6.54 |
| Zr (ICP-MS) | 285 | 248 | 242 | 234 | 254 | 252 |
| Hf (ICP-MS) | 5.57 | 4.95 | 4.89 | 4.87 | 5.24 | 5.19 |
| Eu (ICP-MS) | 1.74 | 1.59 | 1.84 | 1.98 | 2.07 | 1.92 |
| Gd (ICP-MS) | 5.83 | 5.34 | 6.07 | 6.36 | 6.59 | 6.15 |
| Dy (ICP-MS) | 3.27 | 2.93 | 3.40 | 3.78 | 4.00 | 3.69 |
| Ho (ICP-MS) | 0.62 | 0.55 | 0.63 | 0.70 | 0.76 | 0.69 |
| Y (ICP-MS) | 17.3 | 14.3 | 16.7 | 19.4 | 22.0 | 19.9 |
| Er (ICP-MS) | 1.60 | 1.41 | 1.65 | 1.82 | 1.96 | 1.79 |
| Yb (ICP-MS) | 1.33 | 1.14 | 1.35 | 1.49 | 1.62 | 1.46 |
| Lu (ICP-MS) | 0.19 | 0.16 | 0.19 | 0.21 | 0.23 | 0.21 |

Loss in ignition (LOI) was determined by gravimetric methods assuming Fe$_2$O$_3$ = 0.15 FeO. Mg# = Mg/(Mg + Fe) mol%
Primordial mantle-normalized trace element diagrams (Fig. 4, McDonough and Sun 1995) show the typical intra-plate Etnean pattern (Peccerillo and Lustrino 2005; Ferlito et al. 2009; Giacomoni et al. 2018). All samples are characterized by a general enrichment in incompatible elements with negative anomalies in K, Rb, Th, Ta and Ti, and positive anomalies in La, and Nb.

Discussion

Mineral-melt equilibrium conditions

Equilibrium conditions between Ol, Cpx and Plg and melt have been tested before applying the thermo-barometric equations, to investigate the dynamics of crystallization and magma mingling in highly dynamic open-conduit volcanic systems (Mollo et al. 2015; Giacomoni et al. 2018, Lanza-fame et al. 2020).

According to previous studies on Mt. Etna’s plumbing system (Giacomoni et al. 2016, 2018), near-equilibrium crystallization is generally attained between phenocryst cores and their host liquids (i.e. whole-rock analyses). Rationally, phenocrysts cores formed during the early stage of magma crystallization when magmatic conditions (i.e. low undercooling degree) are more favourable to allow progressive chemical re-equilibrium between crystal and host-melt (Mollo et al. 2015; Giacomoni et al. 2018). In addition, the microcrystalline matrix of the samples rarely preserves analysable glass which is commonly too rich in SiO$_2$ and alkaline elements to be representative of the magma composition. In the highly dynamic Mt. Etna open-conduit feeding systems, frequent cryptic mixing involving magmas with similar composition and P–T–XH$_2$O conditions leads to the crystallization of outer crystal portions occurring under non-equilibrium conditions or, more realistically, in equilibrium with melt portions not represented by the whole-rock composition. For these reasons, we have initially tested equilibrium of Ol, Cpx and Plg with their host whole-rock composition. In case of disequilibrium, crystals have been tested with
more primitive or evolved melt compositions to obtain P–T information about crystallization at greater depth.

The equilibrium crystallization of olivine has been verified through Fe–Mg partitioning, which has a minor dependence on melt composition, \( T \), \( P \), and \( f_{O2} \), with an almost constant value of \( \text{ol–liq} K_{\text{Fe–Mg}} = 0.30 \pm 0.03 \) (Roeder and Emslie 1970; Mollo et al. 2015). The \( \text{Fe}^{2+}/\text{Fe}^{3+} \) ratio of the melt is assumed to be close to 0.15, as calculated from the equation of Kress and Carmichael (1991) by assuming an average oxygen fugacity between FMQ + 1 and FMQ + 2 (Giacomoni et al. 2014). Most olivine from NSEC plots in a Fo range of Fo 72.5–75.0 and is in equilibrium with the host rock, while several crystals with higher Fo (Fo 75–80) are not, and thus were reasonably formed in a more basic melt (Fig. 5a). NEC olivine presents a similar situation: equilibrium is obtained in the Fo 72.9–75.7 range, while olivine with Fo 76.5–78.9 are equilibrated with a more basic magma (Fig. 5a). Differently, only a small number of olivine crystals from CC with a restricted range of compositions (Fo 73.2–73.4) can be considered in equilibrium with the host magma. Most phenocrysts with Fo 77.6–81.4 formed in a more basic magma while at Fo 68.6–72.1 olivine would be equilibrated with a more differentiated melt (Fig. 5a).

Olivine crystals having \( \text{ol–liq} K_{\text{Fe–Mg}} < 0.27 \) have been traced back to equilibrium conditions using the composition of the products erupted during the 4/3/2012 eruptive
event (i.e. magma with Mg# 51.5; Giacomoni et al. 2018) to obtain further information about the crystallization conditions of the deeper portion of the feeding system (Fig. 5a). Despite some geochemical differences between magma erupted during the 2012 and the 2015 activity, the composition of the lava erupted during the 4/3/2012 paroxysmal event represents the least evolved and temporally closest melt.

Conversely, equilibrium of clinopyroxene has been tested assuming \( \text{cpx} - \text{melt} K_d \text{Fe-Mg} = 0.27 \pm 0.03 \), as indicated by Putirka et al. (2003). As with Ol, Cpx from NSEC and NEC contain several equilibrated phenocrysts in a range of Mg# varying from 72.9 to 77.5 and from 74 to 77.7, respectively (Fig. 5b). NSEC also contain a few crystals in equilibrium with both a more basic (Mg# 77.2–79.1) and more evolved (Mg# 67.7–72.5) melt, while phenocrysts not in equilibrium in NEC formed exclusively in a more evolved melt in a Mg# range of 70.7–73.4. CC Cpx find themselves in equilibrium in a Mg# range between 74.9 and 79.1, while some crystals (79.1–81.5) formed in a more primitive melt. However, the majority (69.7–74.8) were equilibrated with a more evolved magma (Fig. 5b).

**P–T crystallization conditions**

The crystallization temperature of olivine has been estimated by means of the Putirka et al. (2007) equation, which yields an error of \( \pm 25 \) °C; calculation was performed assuming 4 wt% of H\(_2\)O in the melt, in accordance with recent studies concerning Mt. Etna melt inclusions by Gennaro et al. (2018) and Giacomoni et al. (submitted). Results show that olivine phenocrysts in equilibrium with NSEC host rock form in a range of temperatures from 1092 to 1100 °C. Olivine from NEC and CC are equilibrated at a higher \( T \), ranging from 1100 to 1111 °C and from 1108 to 1111 °C respectively. Olivine crystals with \( \text{Ol-Liq} K_d \text{Fe-Mg} < 0.27 \), traced back to equilibrium with the 4/3/2012 melt composition yield crystallization \( T \) ranging from 1109 to 1117 °C in NSEC, from 1107 to 1117 °C NEC and from 1104 to 1113 °C CC.

Crystallization temperatures and pressure of clinopyroxene have been determined by means of the mineral-melt exchange reactions of diopside/hedenbergite-jadeite and hedenbergite/Ca-Tschermak as reappraised by Putirka (2008). The temperature has been estimated through geothermometer Eq. 33 (Putirka 2008) which has the advantage of being pressure independent with an estimated error of \( \pm 25 \) °C. Resulting temperature was then applied as input data to determine the crystallization pressure by means of the Putirka et al. (2003) equation with an error of \( \pm 170 \) MPa.

Results indicate that NSEC clinopyroxene formed in a range of pressure and temperatures from 640 MPa at 1134 °C to 180 MPa at 1088 °C. These data clearly differ from those obtained by NEC and CC phenocrysts which formed from 350 MPa at 1102 °C to 30 MPa at 1067 °C and from 650 MPa at 1122 °C to 4.7 MPa at 1104 °C (Fig. 6).
The December 2015 feeding system

Petrographic, geochemical and thermo-barometric constraints from the 2015 activity, in comparison with the previous paroxysmal events of 2011–2012 (Giacomoni et al. 2018), provide new insights on the physico-chemical conditions of magma and on the occurrence of pre-eruptive mixing and ascent dynamics.

The whole-rock geochemistry and petrographic features of products emitted during the December 2015 paroxysmal activity do not show significant differences from the highly porphyritic lavas commonly emitted during summit and sub-terminal activity of Mt. Etna. Samples erupted from the NSEC craters present a slightly more basic composition with respect to CC and NEC. However, both major variation diagrams and trace element concentration seems to indicate a common liquid line of descent, dominated by fractional crystallization at pressures higher than 600 MPa (< 22 km b.s.l.). Consequently, observed differences in composition must be ascribed to crystallization conditions and differentiation processes en-route to the surface, likely at a shallow crustal depth. Results from crystal-melt equilibrium and thermo-barometry of olivine and clinopyroxene allow us to delineate some distinct features of the plumbing system of the 2015 activity and to confirm some observations made by studying the previous 2011/2012 paroxysms.

Clinopyroxene thermo-barometry suggests that magma crystallized continuously during the ascent toward the surface from ~ 600 MPa in a range of temperature from 1140 to1104 °C (Fig. 6). These conditions are consistent with the cpx liquidus temperature on the cotectic with olivine in a hydrated (4 wt% H2O) melt as determined by Rhyolite-MELTS modelling, but far from the estimated ol liquidus conditions (1220 °C). This observation, together with the overall slightly evolved composition of whole rock, indicate that the mineral assemblage of studied lavas limits our ability to obtain information regarding magmatic crystallization at pressures higher than 600 MPa (< 22 km b.s.l.).

On the other hand, from this depth toward the surface, we can reasonably consider the feeding system as vertically extended without significant magma chambers or ponding zones (Giacomoni et al. 2014, 2018).

Since the cpx and ol should have formed along a cotectic line at the estimated T–P conditions, we have extrapolated the pressure of crystallization of olivine following the method described in Giacomoni et al. (2018). As shown in Fig. 6, olivine in equilibrium with the whole rock (M1 facies, Giacomoni et al. 2018) formed in a range of pressures between 500 MPa to 450 Kbar (CC) and between 500 and 400 MPa (NEC). Conversely, the NSEC M1 olivines equilibrated at significantly lower pressures, between 200 and 300 MPa. Olivine crystals re-equilibrated with the 4/3/2012 more primitive magma composition (M0) equilibrated between 480 and 500 MPa (NEC), 470–510 MPa (CC) and between 380 and 470 MPa (NSEC).

These results, together with data from the 2011–2012 eruptions, have been plotted in P–T space (Fig. 7) obtained by means of Rhyolite-MELTS software (version 1.2.0; Gualda et al. 2012); thermodynamic calculations were made starting from a mantle-equilibrated magma composition from the 2006 eruption (Giacomoni et al. 2014) at 4 wt% H2O at constant fO2 equal to FMQ + 1.5. Since plagioclase is not stable at these conditions above 800 °C, to visualize its stability field and to interpret the recognized textures and compositional zoning, plagioclase stability was determined starting from the NSEC trachybasaltic composition at decreasing H2O content from 3 to 1 wt% (Figs. 7, 8). Each calculation run was performed at equilibrium crystallization mode, starting from a constant pressure and decreasing the temperature from the Tliquidus to 1000 °C. The H2O solubility threshold was obtained in accordance with the model of Papale (1999) and the stratigraphic column after the seismic and geologic studies of Finetti (2005).

Rhyolite-MELTS modelling indicates that the mantle-equilibrated (mg#68) magma with 4 wt% of dissolved H2O reached the liquidus with olivine in equilibrium at Fo88 at ~ 1250 °C at lithospheric mantle depth. Olivine with comparable Fo contents are rarely found in recent products, but some crystals do occasionally occur such as those found in the 2011–2012 eruptive events (Giacomoni et al. 2018). Magma keeps crystallizing and fractionating olivine (~ 10%, Giacomoni et al. 2014 and 2018) during its route through the lithospheric mantle, reaching Ol-Cpx cotectic stability at ~ 1150 °C at 700 MPa, corresponding to the MOHO depth as stated by Finetti (2005). Cotectic Ol in these T–P conditions should vary in Fo content from 84 to 78 (M0 group of 2011–2012). From this depth, the P–T ascent path of magma inside is tracked by Cpx crystallization; this represents an ascent rate of 9.4 and 7.5 MPa/°C, respectively, along a liquidus line constrained by 3 wt% to 4.8 wt% of dissolved H2O for the 2011–2012 and 2015 magmas. However, if Cpx apparently tracks an almost continuous crystallization from 700 to 200 MPa, Ol compositions change discretely (Kahl et al. 2011; Giacomoni et al. 2018). A plausible explanation for this apparent discrepancy could reside in the faster interdiffusion of Fe–Mg in Ol which tend to erase most small microcumulative compositional zoning (Dohmen et al. 2007). Conversely, Cpx complex zoning is more easily preserved and thus tracks a more continuous magma ascent path. However, Ol compositional grouping provides some important information on the dynamics of magma ascent along central conduits, as well as eventual emission from different summit
craters. M1 Ol phenocrysts, in equilibrium with the erupted magma composition, erupted during 2011–2012 and 2015 from NSEC and equilibrated at a significantly lower pressure than those recognized in NEC and CC in 2015 samples. This could be interpreted as a significantly shallower depth of the magma column below the NSEC, thus suggesting that the final magma ascent path differs from those of NEC and CC.

A soon as the magma column reaches the H₂O saturation depth, degassing and undercooling promote the stability and crystallization of plagioclase (Mollo et al. 2015; Giacomoni et al. 2014). In accordance with the estimated P–T ascent path, this occurs at ~280 MPa inside the foreland carbonate sequence (Finetti 2005). The stratigraphic positioning of plagioclase appearance is also in agreement with the depth of “High Velocity Layer” confirmed by several geophysical studies (DeGori et al. 2005; Patanè et al. 2006) and could be interpreted as a rigid cumulitic gabbroic assemblage. The Rhyolite-MELTS thermodynamic plagioclase stability model allows us to interpret the recognized dissolution-regrowth texture (Fig. 8). Plagioclase stability is strongly constrained by H₂O content and An content increases with T, H₂O and decreases with P (Tsuihyama 1985; Humphreys et al. 2006; Viccaro et al. 2010; Giacomoni et al. 2014). Rhyolite-MELTS modelling (Fig. 8) suggest that an An₈₈ plagioclase core could have formed at 1040–1030 °C at 290–200 MPa in a magma with 2.2 wt% of dissolved H₂O (Fig. 8a crystal zone 1), as confirmed by the Waters and Lange (2015) hygrometer. Since Rhyolite-MELTS modelling does not simulate disequilibrium crystallization during degassing, these estimated T should be interpreted as crystallization T under strong undercooling process (and not as the T of the melt). The de-hydration of the melt is indispensable to allow plagioclase crystallization (Humphreys et al. 2006; Giacomoni et al. 2014), as the relatively fast and steady ascent ratio of magma does not allow a significant T drop due to the slow heat diffusion towards the surrounding rocks. Two main processes could promote volatile exsolution and consequent plagioclase crystallization: decompression and CO₂ flushing. Decompression is a well-studied process that can induce magmatic crystallization especially in a volatile-saturated melt (Hammer and Rutherford 2002; Mollard et al. 2012). Alternatively, a CO₂-rich volatile fluid phase exsolved in a deeper portion of the plumbing system and flushing the shallower part of the magmatic column could promote H₂O exsolution and crystallization without significant magmatic volume transfer (Riker et al. 2015; Caricchi et al. 2018).

Early formed An-rich cores often present a rounded edge followed by a less calcic (An₇₀₋₅₂) overgrowth (Fig. 8a, crystal zone 2). The process responsible for this plagioclase destabilization is a “simple dissolution” response occurring when plagioclase reacts with a melt that is undersaturated in the plagioclase phase (Tsuihyama 1985; Giacomoni et al. 2014). The most common
A magmatic process that could cause such disequilibrium is the increasing in H₂O content in the melt due to mixing/mingling with a basic incoming magma with its pristine 4 wt% of dissolved H₂O or, as recently hypothesised, by flushing of volatile H₂O-rich fluids (Ferlito et al. 2010, 2014; Ferlito 2017). In fact, the H₂O/CO₂ ratio of the exsolved fluid increases as CO₂-rich fluids are progressively extracted and migrate upward. Since the magma in the flushed shallower magma has been gradually enriched in CO₂ and depleted in H₂O, it re-equilibrates by exsolving CO₂ and incorporating H₂O. Although this concept still requires further investigation and numerical modelling, it is comparable with that described by Caricchi et al. (2018) and hypothesized by Ferlito et al. (2014). Rhyolite-MELTS modelling suggests that after dissolution, plagioclase growth resumes at shallower depth due to decompression and further magma volatilization, between 130 and 50 MPa.

The second type of plagioclase is instead characterized by a core with low calcic composition (An70-52; Fig. 8b zone 2), analogous to the outer zone of the first type, surrounded by an anorthitic dusty rim (An88-80; Fig. 8 zone 3). Dusty rims have been experimentally reproduced after a “partial dissolution” reaction between the pre-existing low calcic phenocryst with a melt in which a more anorthitic plagioclase is stable (hotter or more H₂O-rich Tsuchiyama 1985). Since Rhyolite-MELTS, hygrometers and volatile solubility models indicate that An70-52 cores are stable in a magma with 1.8 wt% of dissolved H₂O, they should have formed at significant shallower depth with respect to the previously described An88-80 plagioclases. In accordance with the solubility model of Ghiorso and Gualda (2015), feasible
stability conditions are reached between 130 and 50 MPa, suggesting that both magma mingling and H2O-flushing could have been responsible for triggering an eruption.

Concluding remarks

The petrologic and thermo-barometric study of the December 2015 eruptive event has contributed to improving our knowledge of the vertically continuous feeding system of Mount Etna, which had been already illuminated thanks to the paroxysmal activity of 2011–2012 (Giacomoni et al. 2018).

Thermo-barometric estimates and Rhyolite-MELTS modelling indicate that (1) primary volatile-rich magma crosses its liquidus T (~1250 °C) at ~1GPa, (2) formed Ol phenocrysts (Fo88) keep re-equilibrating during magma ascent, reaching iii) the Ol-Cpx cotectic T (~1150 °C) at ~700 MPa. Ol crystal equilibrium conditions of the NSEC crater differ significantly from those of the CC and the NEC, indicating a shallower depth of the magmatic column which fed the eruption at NSEC. However, Ol phenocrysts suggest that each crater was subject to a deep magmatic recharge between 530 and 440 MPa.

Finally, Plg stability is strongly constrained by the H2O/CO2 of the melt. According to Rhyolite-MELTS simulations, the great majority of Ar88-80 plagioclases are in equilibrium with a magma with 2.2 wt% of dissolved H2O, at 290–200 MPa. However, CO2 flushing could be an efficient process to promote H2O exsolution and consequent Plg crystallization at greater depth. Similarly, the increase of the H2O/CO2 ratio of the flushing fluid in an almost steady-state vertical conduit regime could have a deep effect on the stability of crystallized phenocrysts, inducing crystal dissolution and resorption without significant magma volume transfer and therefore worthy of further, more focused studies.

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