Rapid differentiation in a sill-like magma reservoir: a case study from the Campi Flegrei caldera

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In recent decades, geophysical investigations have detected wide magma reservoirs beneath quiescent calderas. However, the discovery of partially melted horizons inside the crust is not sufficient to put constraints on capability of reservoirs to supply cataclysmic eruptions, which strictly depends on the chemical-physical properties of magmas (composition, viscosity, gas content etc.), and thus on their differentiation histories. In this study, by using geochemical, isotopic and textural records of rocks erupted from the high-risk Campi Flegrei caldera, we show that the alkaline magmas have evolved toward a critical state of explosive behaviour over a time span shorter than the repose time of most volcanic systems and that these magmas have risen rapidly toward the surface. Moreover, similar results on the depth and timescale of magma storage were previously obtained for the neighbouring Somma-Vesuvius volcano. This consistency suggests that there might be a unique long-lived magma pool beneath the whole Neapolitan area.

Large-volume silicic caldera systems have produced some of the largest explosive eruptions in the world. The generation of these large magmatic systems, their mechanisms of differentiation, and the timescale of overpressuring and opening are crucial issues for volcanic risk evaluation, particularly in densely populated areas.

The Campanian Volcanic Zone developed within the graben of the Campanian Plain, which formed during the Plio-Pleistocene epoch as a consequence of an extensional system likely related to the final stages of the opening of the southern Tyrrhenian Sea. Geothermal boreholes drilled north-west of the Campi Flegrei caldera revealed that widespread effusive volcanism characterised by calc-alkaline andesitic and basaltic compositions occurred within the Plain approximately 1.5 Ma ago. More recently, in the last 300 ka, widespread volcanic activity fed by alkaline magmas has taken place within the southern part of the Campanian Plain, especially in the last 50 ka on the Vesuvius central volcano and in the Campi Flegrei, Ischia and Procida volcanic fields. The composition of the eruptive products has been practically unchanged throughout this time, which suggests that a wide alkaline magmatic source has been active up to recent times. The largest part of the magma volume was erupted from central and fissure vents in a zone presently cantered in the Campi Flegrei caldera and its surroundings, the most active part of this long-lived plumbing system.

During its eruptive history, the high-risk Campi Flegrei volcanic district west of Naples emitted ca 400 km$^3$ of alkaline magmas in a variety of eruptions ranging from low Volcanic Explosivity Index (VEI < 3) events, generally supplied by less-evolved magmas, to super-eruptions involving chemically differentiated magmas. A wide low velocity layer, interpreted as an extended magmatic body, has been detected by seismic tomography at 8–10 km beneath the Campi Flegrei caldera and the neighbouring Vesuvius volcano$^{1,2}$. To investigate the likely state of this wide plumbing system, we have tracked the pre-eruptive and eruptive history of the magmas by studying the geochemical (matrix-glass compositions), isotopic (oxygen isotope compositions of separated minerals) and textural (crystal number density and size distributions) features of representative rock samples collected from volcanic deposits from past eruptions in Campi Flegrei. The data were compared to data that were previously acquired$^1$ for Somma-Vesuvius volcanic rocks in order to explore the possible relationship between the two feeding systems.

Results

Magma evolution in crustal reservoirs. The major element concentrations of matrix glasses in the samples range from mafic trachybasalt to felsic trachyte (Figure 1, Table 2). Felsic compositions are generally represented by pumice samples collected from deposits of highly explosive eruptions (VEI 4-7), with the exception of the scoria
samples produced by the strombolian Monte Nuovo eruption. Pumice samples are nearly aphyric (1–5% crystals), highly vesiculated (70%), with microlite-poor or microlite-free groundmass. In contrast, mafic compositions typically occur in scoria from strombolian eruptions (VEI 2–3). These are relatively dense, with 40–60% vesicularity, high (20–30%) phenocryst contents, and a groundmass generally containing abundant feldspar microlites (< 100 micron) (Figure 2, Table 3).

Previous isotopic investigations (e.g.4 and references therein) have indicated that these co-genetic mafic and felsic rocks developed by crystal-liquid fractionation associated with a minor amount of crustal assimilation in a multi-level magma reservoir. Felsic magmas occupied an upper zone of crystal-poor buoyant melt located between 250–150 Mpa according to thermo-dynamical modelling and phase-equilibrium experiments5,6, and were supplied by a deeper crystal-rich mafic layer. In addition, a minor amount of limestone assimilation during magma differentiation has been inferred on the basis of isotopic studies on Campanian magmas2,7. Such crustal contamination processes, leading to the generation of CO2-rich fluid

| Table 1 | Studied Campi Flegrei volcanic formations |
|------------------|------------------|------------------|------------------|
| Volcanic Formation | VEI | Age (Ka) | Volcanological features | Composition |
| Campanian Ignimbrite | 7 | 39 | low aspect ratio ignimbrite | trachyte/phonolite |
| Neapolitan Yellow Tuff | 6 | 14 | wet pyroclastic surge | latite-trachyte |
| Agnano Monte Spina | 5 | 4.1 | pyroclastic flow | trachyte |
| Pomici Principali | 4 | 10.3 | plinian fallout deposit | latite/trachyte |
| Astroni | 4 | 3.8 | tuff ring | trachyte |
| Minopoli | 3 | 11 | strombolian deposit | trachybasalt |
| Baia | 3 | 8.5 | tuff cone | trachyte |
| Averno | 3.4 | 4.5 | tuff ring | trachyte |
| Montagna Spaccata | 3 | 9.7 | tuff ring | trachyte |
| Nisida | 2 | 10.6 | tuff cone | latite |
| S. Teresa | 2 | 10.6 | tuff cone | latite |
| Senga | 2 | 3.8 | cinder cone | latite |
| Monte Nuovo | 2 | 1538 AD | cinder cone | trachyte/phonolite |
| Monte Olibano | 0 | 3.9 | lava dome | latite |

Figure 1 | The Campi Flegrei volcano, showing the location of vents of the studied eruptions. Bold dashed lines: vent of Campanian Ignimbrite eruption presumed on the basis of geological evidence51, dotted line: Neapolitan Yellow Tuff caldera hypothesized on the basis of geophysical data52. Right: schematic chronogram of Campi Flegrei activity as recorded by stratigraphic successions. Bottom: plot of total alkalis vs. silica for Campi Flegrei studied rocks; symbols correspond to the vent locations on map.
## Table 2 | Electron microprobe analyses of matrix-glass of rocks of the Campi Flegrei formation

| Stratigraphic Height | Basal Fallout | Lower Flow | Middle Flow | Upper Flow | Lower Member | Upper Member | Member A |
|---------------------|--------------|------------|-------------|------------|--------------|--------------|---------|
| SiO₂                | 61.81        | 62.72      | 61.26       | 61.11      | 59.17        | 60.56        | 61.67   |
| TiO₂                | 0.42         | 0.42       | 0.44        | 0.42       | 0.50         | 0.38         | 0.49    |
| Al₂O₃               | 19.15        | 19.05      | 18.95       | 18.62      | 18.68        | 18.58        | 18.63   |
| FeO                 | 2.87         | 2.48       | 3.26        | 3.52       | 4.46         | 3.87         | 3.38    |
| MnO                 | 0.23         | 0.26       | 0.16        | 0.11       | 0.12         | 0.16         | 0.01    |
| MgO                 | 0.45         | 0.32       | 0.59        | 0.82       | 1.17         | 0.93         | 0.67    |
| CaO                 | 1.95         | 1.58       | 2.23        | 2.56       | 3.84         | 3.29         | 2.48    |
| Na₂O                | 5.03         | 5.52       | 4.65        | 2.96       | 3.64         | 3.58         | 4.13    |
| K₂O                 | 8.02         | 7.60       | 8.42        | 9.74       | 8.20         | 8.40         | 8.54    |
| H₂O                 | 0.08         | 0.06       | 0.11        | 0.15       | 0.23         | 0.17         | 0.00    |

### Formations

| Stratigraphic Height | Member B | Member C | Member D | Member E | Member F | Member Alpha | Basal Fallout |
|---------------------|----------|----------|----------|----------|----------|--------------|---------------|
| SiO₂                | 61.91    | 61.75    | 60.91    | 61.04    | 60.52    | 57.83        | 60.74        |
| TiO₂                | 0.49     | 0.48     | 0.51     | 0.56     | 0.52     | 0.51         | 0.46         |
| Al₂O₃               | 18.89    | 18.34    | 18.63    | 18.61    | 18.64    | 19.33        | 18.33        |
| FeO                 | 3.48     | 3.47     | 3.74     | 3.87     | 3.77     | 4.40         | 3.55         |
| MnO                 | 0.02     | 0.01     | 0.01     | 0.01     | 0.02     | 0.15         | 0.14         |
| MgO                 | 0.71     | 0.62     | 0.77     | 0.91     | 0.91     | 1.12         | 0.65         |
| CaO                 | 2.14     | 2.50     | 2.71     | 2.99     | 2.89     | 3.74         | 2.43         |
| Na₂O                | 7.47     | 5.26     | 4.03     | 3.85     | 3.90     | 3.86         | 4.63         |
| K₂O                 | 4.88     | 7.27     | 8.69     | 8.77     | 8.82     | 8.88         | 8.95         |
| P₂O₅                | 0.02     | 0.01     | 0.00     | 0.00     | 0.18     | 0.11         | 0.11         |
| Cl                  | 0.46     | 0.61     | 0.57     | 0.56     | 0.56     | 0.58         | 0.93         |
| F                   | 0.38     | 0.25     | 0.07     | 0.05     | 0.06     | 0.22         | 0.32         |

### Formations

| Stratigraphic Height | Middle Fallout | Upper Fallout | Surge | Flow | Scoria | Basal Fallout | Scoria |
|---------------------|---------------|---------------|-------|------|--------|---------------|-------|
| SiO₂                | 59.88         | 59.82         | 60.24 | 59.99| 52.59  | 62.75         | 58.20 |
| TiO₂                | 0.52          | 0.47          | 0.48  | 0.49 | 0.96   | 0.40          | 0.34  |
| Al₂O₃               | 18.70         | 18.93         | 18.71 | 18.86| 18.00  | 18.26         | 17.83 |
| FeO                 | 3.68          | 3.44          | 3.45  | 3.49 | 7.37   | 2.99          | 2.90  |
| MnO                 | 0.12          | 0.14          | 0.14  | 0.15 | 0.15   | 0.26          | 0.15  |
| MgO                 | 0.75          | 0.61          | 0.61  | 0.65 | 4.61   | 0.56          | 0.57  |
| CaO                 | 2.41          | 2.61          | 2.47  | 2.48 | 10.01  | 1.79          | 2.28  |
| Na₂O                | 5.04          | 4.56          | 4.67  | 4.50 | 2.56   | 6.60          | 3.84  |
| K₂O                 | 8.75          | 9.31          | 9.13  | 9.30 | 3.78   | 6.69          | 7.06  |
| P₂O₅                | 0.14          | 0.11          | 0.10  | 0.11 | 0.52   | 0.01          | 0.10  |
| Cl                  | 0.59          | 0.87          | 0.95  | 0.89 | 0.20   | 0.86          | 0.70  |
| F                   | 0.27          | 0.32          | 0.38  | 0.32 | 0.17   | 0.45          | 0.19  |

### Formations

| Stratigraphic Height | S. Teresa | Senga | Monte Nuovo | Monte Nuovo | Monte Nuovo | Monte Nuovo | M. Olibano |
|---------------------|-----------|-------|-------------|-------------|-------------|-------------|-----------|
| SiO₂                | 57.98     | 58.62 | 61.21       | 60.96       | 60.13       | 61.73       | 55.98     |
| TiO₂                | 0.84      | 0.45  | 0.42        | 0.40        | 0.34        | 0.34        | 0.64      |
| Al₂O₃               | 16.38     | 18.92 | 20.32       | 20.46       | 20.22       | 20.47       | 17.82     |
| FeO                 | 7.08      | 3.56  | 2.86        | 2.82        | 2.54        | 2.51        | 5.88      |
| MnO                 | 0.22      | 0.11  | 0.22        | 0.23        | 0.24        | 0.20        | 0.15      |
| MgO                 | 1.57      | 0.71  | 0.23        | 0.20        | 0.29        | 0.19        | 3.35      |
| CaO                 | 7.88      | 2.58  | 1.76        | 2.06        | 1.72        | 1.78        | 7.02      |
| Na₂O                | 2.22      | 4.46  | 5.53        | 5.86        | 6.77        | 5.61        | 3.00      |
| K₂O                 | 5.35      | 9.47  | 7.39        | 6.95        | 7.72        | 7.15        | 5.72      |
| P₂O₅                | 1.05      | 0.11  | 0.04        | 0.03        | 0.04        | 0.02        | 0.42      |
Figure 2 | Back-scattered electron images of selected rocks from the Campi Flegrei caldera: (a) Campanian Ignimbrite super-eruption (39 ka), and (b) Astroni eruption (3.7 ka). These images were processed using dedicated programs to obtain the CSD curves by plotting population density (number of crystals per unit volume) versus crystal size (maximum length).
phases, could increase the gas content of the magmas and thus the eruption explosivity. Therefore, to quantify the role of limestone contamination in the evolution of the Campi Flegrei magmatic system, stable isotope studies were carried out on minerals that were separated from rocks representing eruptions with various explosivities.

Clinopyroxene and sanidine that are separated from rock samples have similar $\delta^{18}O$ values, ranging from 6.66–7.04‰ and 8.09–8.51‰, respectively. These values do not correlate with the explosivity indices of the eruptions (Figure 3; Table 4). Although the oxygen isotope fractionation factors have not been measured specifically for the alkaline Campi Flegrei rocks, various authors indicate that the clinopyroxene-sanidine fractionation varies, for silicic rocks, between 1.2 – 2.3‰ at magmatic temperatures. Therefore we consider that sanidine and clinopyroxene samples are in oxygen isotope equilibrium. Moreover, we can assume that the sanidine results closely approach magma values; in fact, the sanidine-melt fractionation would be the main cause of the explosive behaviour of the Campi Flegrei rocks, suggests that the limestone assimilation process cannot alone be the main cause of the explosive behaviour of the Campi Flegrei magmas. Indeed, the eruption dynamics would have been mostly caused by a combination of factors such as magma volume, degree of carbonate assimilation, and the mechanism and timing of degassing processes during the magma’s rise through the volcanic conduit. Therefore, the calculated low degree of contamination, common to eruptions with a very wide range of explosivity (as indicated by the similarity in $\delta^{18}O$ values measured in the selected Campi Flegrei rocks), suggests that the limestone assimilation process cannot alone be the main cause of the explosive behaviour of the Campi Flegrei magmas. Indeed, the eruption dynamics would have been mostly caused by a combination of factors such as magma volume, degree of carbonate assimilation, and the mechanism and timing of degassing processes during the magma’s rise through the volcanic conduit.

Magma ascent: degassing processes in volcanic conduit. To investigate the degassing process of these melts during ascent from the magma chamber toward the surface, the vesicularity of pumice and scoria specimens was measured as well as the residual water content in their matrix glasses, which represent the magma melt fraction quenched at the time of eruption (Table 2).

The measured values of vesicularity (60–70 wt%) of felsic pumices from highly explosive eruptions (VEI 4 to 7) are consistent with those calculated (following) in the case of a closed-system equilibrium.

### Table 3 | Textural parameters

| Formation          | units     | bulk vesicularity | n. of crystals | Intercept (mm$^{-1}$) | Slope (mm$^{-1}$) | Lmax (mm) | time$^1$ years | time$^2$ years |
|--------------------|-----------|-------------------|----------------|-----------------------|-------------------|-----------|----------------|----------------|
| Campanian Ignimbrite | basal fallout | 0.8              | $< 100$     | -                     | -                 | 1.1       | -              | 17–174        |
| Campanian Ignimbrite | upper flow  | 0.8              | 989          | 4.66                  | -2.79             | 1.5       | 11–114         | 24–238        |
| Neapolitan Yellow Tuff | lower member | 0.7              | $< 100$     | -                     | -                 | 1         | -              | 16–159        |
| Agnano Monte Spina | basal fallout | 0.8              | $< 100$     | -                     | -                 | 2         | -              | 32–317        |
| Pomici Principali | basal fallout | 0.7              | $< 100$     | -                     | -                 | 1         | -              | 16–159        |
| Astroni | basal fallout | 0.7              | 28           | 2.32                  | -0.7              | -         | -              | -             |
| Minopoli | scoria      | 0.7              | 184          | 4.02                  | -4.09             | 1.2       | -              | 8–78          |
| Baia | basal fallout | 0.8              | $< 100$     | -                     | -                 | 1         | -              | 16–159        |
| Nisida | scoria flow  | 0.5              | 251          | 0.12                  | -6.71             | 1.5       | 4–42           | 24–238        |
| S. Teresa | scoria flow | 0.7              | 79           | 5.26                  | -5.41             | 0.6       | 6–59           | 9–95          |
| Monte Nuovo | scoria flow | 0.5              | 318          | 8.42                  | -7.86             | 0.5       | 4–40           | 8–79          |
| Monte Olibano | lava dome    | 0.5              | 849          | 7.32                  | -3.71             | 3         | 8–85           | 48–476        |

| Microlites | number density (mm$^{-3}$) | n. of crystals | Intercept (mm$^{-1}$) | Slope (mm$^{-1}$) | Lmax (m) | time$^1$ hours | time$^2$ hours |
|------------|-----------------------------|----------------|-----------------------|-------------------|----------|----------------|----------------|
| Campanian Ignimbrite | basal fallout | -               | none                  | -                  | -        | -              | -              |
| Neapolitan Yellow Tuff | lower member | -               | none                  | -                  | -        | -              | -              |
| Agnano Monte Spina | basal fallout | -               | none                  | -                  | -        | -              | -              |
| Pomici Principali | basal fallout | -               | none                  | -                  | -        | -              | -              |
| Astroni | fallout | 1.1E±05          | 1055                  | 21.7              | -307         | 60          | 0.5–6         | 5–60          |
| Minopoli | scoria flow  | 9.9E±02          | 197                   | 20.69             | -152        | 38           | 1–13          | 3–38          |
| Baia | basal fallout | 9.4E±02          | 187                   | 9.7              | -23.3       | 120          | 7–85          | 10–120        |
| Nisida | scoria flow  | 1.2E±02          | 734                   | 15                | -27         | 80           | 6–74          | 7–79          |
| S. Teresa | scoria flow | 9.2E+01          | 257                   | 10.88             | -29.8       | 80           | 6–57          | 7–79          |
| Monte Nuovo | MN1     | 4.9E±03          | 970                   | 18.5              | -51.3       | 80           | 3–38          | 7–79          |
| Monte Nuovo | MN2     | 3.6E±03          | 723                   | 17.35             | -25.8       | 80           | 6–77          | 7–79          |
| Monte Nuovo | MN3     | 2.4E±03          | 472                   | 16.28             | -24.6       | 80           | 7–81          | 7–79          |
| Monte Nuovo | MN4     | 9.0E±03          | 1800                  | 18.29             | -35.5       | 130          | 5–56          | 10–130        |
| Monte Olibano | lava dome | 5.6E+03          | 1111                  | 10.49             | -28.9       | 120          | 6–69          | 10–120        |

$^1$ Lmax (mm) time 1 years, $^2$ Lmax (mm) time 2 years.
degassing, with the assumption that no further degassing occurred after fragmentation. This last assumption, although not always true (e.g.15,16), is generally accepted for high VEI eruptions, for which the time between fragmentation and quenching in the atmosphere is on the order of seconds. Moreover, in the case of Campi Flegrei magmas, experimental studies indicate that evolved alkaline liquids, due to their lower viscosity, usually degas at equilibrium conditions over the wide range of decompression rates that is typical of most explosive eruptions (e.g.17,18). Consequently, under the inferred equilibrium degassing conditions, the water content measured in matrix glass closely represents the residual dissolved water content in the melt at the time of fragmentation. Considering the relationship between pressure and water content obtained experimentally for trachytic melts19 at temperatures of 850–900 °C, the measured water content in matrix glass of pumice samples (1.4 to 1.8 wt%) corresponds to a fragmentation pressure of 25–40 Mpa (1100–1600 m).

In contrast, the low water content (0.8 to 1.1 wt%) of scorias from VEI 2–3 eruptions indicates a quenching pressure approximately 20 Mpa (800 m). These values, associated with low vesicularity (30–40 wt%), are direct evidence of open-system degassing, implying that gas loss occurred at shallow depth in the conduit, most likely due to low ascent rates. An exception is represented by the higher water content of sample MN2 from the phreato-magmatic units of the Monte Nuovo eruption, which is possibly related to the interaction of magma with ground water.

**Timescale of magma evolution and ascent processes.** To constrain the timescale of evolution of mafic magmas toward felsic compositions, and examine their eruptive ascent velocity, crystal size distribution theory (CSD)20,21 has been applied. A CSD is generally shown as a semi-logarithmic plot of population density (number of crystals per unit volume) versus crystal size (maximum length) with the slope equal to 1/(growth rate x residence time). Thus, if the growth rate is known, the residence time can be computed.

The graphs of the distribution of crystal size in the samples show multiple slopes corresponding to phenocrysts and microlites (< 100 micron; Fig. 2). These, according to the results of previous experiments22, are evidence of different nucleation and under-cooling rates (Table 3) during magma crystallisation. In particular, the steep slopes (−23 to −307, Table 3) and high intercept values (9–21, Table 3) of the acicular microlites suggest that they underwent rapid nucleation because of under-cooling during magma ascent and syn-eruptive degassing in the volcanic conduit23. In contrast, the flat slopes (−2 to −7, Table 3) with low intercepts (2–12, Table 3) shown by the euhedral phenocrysts indicate that they grew under near-equilibrium conditions and modest cooling rates in crustal magma chambers (Figure 2).

Data on the average growth rates during the crystallisation of both phenocrysts and microlites were collected for the CSD Marsh model so that the crystal growth time could be estimated. Under conditions of slight under-cooling in magma chambers, experimental and CSD studies indicate that crystal growth rates are consistently between

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**Figure 3** | Oxygen isotope composition of clinopyroxene (squares) and sanidine (triangles) crystals from four main eruptive units of the Campi Flegrei caldera plotted versus (a) Volcanic Explosivity Index (VEI) and (b) CaO/Al2O3.

**Table 4** | δ18O measured in clinopyroxenes and sanidines of Campi Flegrei selected rocks

| Volcanic Formation | Samples | Weight (μg) | δ18Ocorr(‰) assumed % | %yield |
|--------------------|---------|-------------|----------------------|--------|
| **Cpx**            |         |             |                      |        |
| Averno             | V6 CPX  | 1753        | 7.04                 | 43     | 121    |
| Pomici Principali  | P1 CPX  | 1724        | 6.95                 | 43     | 87     |
| Monte Olibano      | OL CPX  | 1736        | 6.66                 | 43     | 103    |
| Averno             | AVI SAN | 1720        | 8.38                 | 47     | 75     |
| Averno             | AV6 fallout | 1751     | 8.32                 | 47     | 73     |
| Pomici Principali  | PP1 SAN | 1722        | 8.09                 | 47     | 67     |
| Pomici Principali  | P3 SAN  | 1751        | 8.11                 | 47     | 67     |
| Monte Olibano      | OL SAN  | 1724        | 8.16                 | 47     | 65     |
| Campanian Ignimbrite| TF SAN | 1722        | 8.51                 | 47     | 70     |
| **San**            |         |             |                      |        |
| Standards          | G JAG   | 1732        | 5.40                 |        |
|                    | G JAG   | 1726        | 5.45                 |        |

*G JAG - internal standard calibrated against the NBS-30 biotite international standard.*
Because decompression experiments indicate that the growth rate of microlites can be highly dependent on the decompression rate (e.g.\(^2\)), in our calculations we have used the entire range of growth rates (from \(1.39 \times 10^{-9}\) to \(1.65 \times 10^{-6}\) mm/s) available for sanidine in the Campi Flegrei rocks. These rates were obtained by experiments\(^{37}\) performed at different decompression rates (time intervals between 7200 and 50,400 seconds\(^{38}\)). By applying this range of growth rates to the CSD slopes of the microlite population and to the coarsest microlites, we calculate that magma ascent lasted from hours to days for low VEI events (Table 3). It is worth noting that decompression experiments\(^{37}\) have shown that the absence of microlites in the groundmass of pumices from highly explosive eruptions indicates that the magma decompression time was too short (in the order of a few hours) to generate microlites. The link between rapid magma ascent and the absence of microlites in large VEI eruptions is also confirmed by the growth rates obtained by decompression experiments on natural Campi Flegrei trachytic melts\(^{36}\). The absence of reaction rims on biotite crystals, which, as indicated by phase equilibrium experiments\(^{5}\) on Campi Flegri rocks, become unstable at pressures below 135 Mpa in a short time span, provides yet another line of evidence.

Discussion

Our data from volcanic rocks ejected from Campi Flegrei show that the alkaline magmas evolved relatively rapidly (over centuries) toward crystal-poor felsic liquids, which were stored at a shallow pressure of ca. 150–250 Mpa (6–10 km depth). These results on the depth and timescale of magma residence are very similar to those found for phonolites erupted from Somma-Vesuvius\(^{5}\), which suggests that since 50 ka, a wide common reservoir may have been active beneath the Neapolitan volcanic districts. Moreover, a unique shallower magmatic source for the two Neapolitan volcanic areas is also supported by the similarity in the Sr and Nd isotopic compositions in both silicic and mafic rocks of Campi Flegrei and Somma-Vesuvius (Figure 4A). In Figure 4B, we compare the crystallisation depth of different mineral phases of Campi Flegrei and Somma-Vesuvius magmas by using the Cl contents measured in matrix glasses (this study) and in melt inclusions\(^{34–37}\), and the values of the saturation pressure for different Cl contents based on the available solubility models\(^{39}\). In particular, the resulting trends indicate that the precipitation of diopside and sanidine occurred mostly at a maximum pressure of at least \(> 200–100\) Mpa for both volcanoes. Salite and leucite crystallised mostly at shallower pressures of \(100–25\) Mpa, most likely during a later crystallisation stage induced by magma degassing in the volcanic conduit, as their Cl contents match those measured in degassed matrix-glasses.

The geophysical anomalies detected by seismic tomography indicate the presence of a partial melting zone at 7–8 km depth beneath both Campi Flegrei and Somma-Vesuvius, while the heat flux distribution\(^{2}\) shows a continuous positive anomaly in the maximum values beneath the Campi Flegrei super-volcano, where the main volume of the feeding system would be concentrated (Figure 5). Furthermore, thermal numerical models\(^{39}\) estimate a high volume of magma (at least 1000 km\(^3\)) above 10 km beneath the caldera. On the other hand, tomographic studies\(^2\) indicate that the low velocity layer is only 1–2 km thick, which means that the large volume of magma has to be distributed as a wide sill-like layer, most likely extending beneath the whole Neapolitan volcanic area between 7–9 km depth, where there is an important lithological break at the transition between sedimentary and metamorphic rocks.

Our CSD data on feldspar phenocrysts indicate that this magmatic system can evolve toward less-dense felsic liquids in a time span comparable with the hundreds of years of volcano repose. This implies that a volatile-rich trachytic layer could be formed at the top of a wide magma reservoir, which could then erupt explosively. Although this scenario corresponds to the worst-case, and other less
severe scenarios are possible, the results of thermo-dynamical models and experiments indicate that the Campi Flegrei trachytic magmas were mainly water-saturated and were therefore capable of generating critical conditions of over-pressurisation in the chamber (5–25 Mpa, following), thus triggering eruption by wall rock rupture. Also, the capability of magmas to evolve toward explosive behaviour in the relatively short time suggested by our CSDs studies (because of their water-saturated nature), implies that the current magma reservoir could be activated at any time and does not necessarily need further magmatic evolution or any external influences (e.g., recharge, water/magma interaction, assimilation) to erupt.

Our data on oxygen isotopes reveal a subordinate (≤3 wt%) degree of limestone contamination, causing the liberation of low amounts of CO₂–rich gas phase, common to eruptions with a very wide range of VEI. These data allow us to exclude the possibility that the assimilation process alone could explain the explosivity of eruptions. Indeed, the eruption dynamics would have been mostly related to a combination of factors, such as magma volume and degree of differentiation as well as the mechanism and timing of the degassing during the magma’s rise through the volcanic conduit. Notably, as revealed by our microlite size distributions, the migration of magma from the reservoir toward the surface during past explosive eruptions at Campi Flegrei has been fast, which implies that in the case of similar behaviour in a future event, there could be very little forewarning before a major eruption.

The critical state of such rapidly evolving systems was demonstrated at Rabaul caldera, in Papua New Guinea, on September 18, 1994, where a VEI 4 event occurred after only 51 years of rest time, and was proceeded by less than a day of precursors. This is particularly crucial at the Campi Flegrei caldera, where the new uplift episode that lasted from November 2004 to October 2006 has been accompanied by the almost continuous increase of the magmatic component of the fumaroles and seismic activity since 2000, and where a new volcanic crisis could suddenly put the densely urbanised metropolitan area of Naples at risk.

Methods

Representative well-documented eruptions with different Volcanic Explosivity Index (VEI) have been selected for the present study, (Table 1) and samples collected at different heights along the stratigraphic sequence. For each sample, bulk vesicularity measurements have been carried out for clasts within –5 and 0 phi size range, as well as electron microprobe (major and volatile elements compositions) and textural (crystal number density and size distribution) analyses of matrix-glasses and minerals by using polished thin sections. Moreover doubly-polished wafers were prepared from selected samples in order to collect Fourier Transform Infrared (FTIR) spectra for evaluating water concentration in matrix glasses. Sanidine and clinopyroxene phenocrysts have been separated from selected samples and analyzed for stable isotope measurements.

Electron microprobe analysis (EMPA). Major and volatile elements analyses of matrix glasses were performed at the CNR - Istituto di Geologia Ambientale eGeoingegneria (CNR-IGAG), Rome, using a Cameca SX-50 electron microprobe and at Istituto Nazionale di Geofisica e Vulcanologia in Rome with a JEOL-JXA-8200
Fourier Transform Infrared spectroscopy analysis (FTIR). To determine the water contents of glasses, FTIR analysis has been performed at the laboratory of the Dipartimento Scienze della Terra Università “Roma Tre”, Rome. The IR spectra were collected using a Nicolet Magna 760 FTIR spectrometer with a KBr beam splitter and a MCT-A detector. A 30-μm diameter, doubly-polished area was examined under a Nic-Plan analytical microscope to avoid bubble and surface imperfections. Each spectrum was obtained with 4 cm⁻¹ resolution and 128 scans, and the background was measured during each analysis. Analytical error on any measured water content was around 10% due to uncertainty in density and glass thickness measurements and error introduced during the measurement of absorption. Results are reported in Table 2.

Textural analysis. Textural analysis of crystal number density and size distribution was carried out on polished thin sections of epoxy-impregnated grains using a standard polarized light microscope for preliminary rock characterization followed by a Field Emission Scanning Electron Microscope (FE-SEM) JOEL JSM-650F (Istituto Nazionale di Geofisica e Vulcanologia, Roma, Italy) for quantitative measurements. Each backscattered electron (BSE) images were processed using Adobe Photoshop and NIH Image 1.60 software to measure the number, shape, size, and volume of microlites and phenocrysts. Stereological methods have been used to convert the area percentages determined by studying two- to three-dimensional textural values. Specifically, crystal number densities and size distributions have been obtained with the CSD Corrections 1.3 program****+ which includes corrections for both intersection probability and cut section effects. Emerging three-dimensional textural analysis methods††††+ that use serial sectioning or X-ray tomography techniques (e.g.) were not appropriate for this study because those methods require a large density contrast between crystals and matrix glass in order to achieve resolutions as small as 40 μm. Moreover, Mock and Jerram have shown that CSDs determined with two-dimensional textural data compare reasonably well with three-dimensional CSDs for sample sizes greater than 200 individual crystals and regular crystal shapes. Results are reported in Table 3.

Bulk vesicularity measurement. Bulk vesicularities have been obtained by comparing the density of juvenile vesicular clasts with the density of non-vesicular clasts for the composition of interest (as in†††††). In order to account for possible density variations, sets with clasts of 2 to 0.5 phi size range were used for density measurements. Sets of 30 clasts for each granulometric class between 0 and 5 phi were weighted and coated with a thin film of paraffin wax, and their average density was determined using a pycnometer. In order to evaluate a standard deviation for clast density within the same sample, the density of single clasts in the range of 2 to 5 phi were determined individually. The volume of the paraffin wax film has been considered negligible because its density was about equal to that of water (1000 kg/m³). The average density for each sample has been calculated. Results are reported in Table 3.

Stable isotope analysis. Oxygen isotope ratios were measured at Royal Holloway, University of London (RHUL), using a Laser Fluorination Prep system with a Synrad CO₂ Laser and BfSf reagent on line to a GV Instrument Optima dual inlet IRMS, described in Mattey. Optical cleaning of sandine and clinoxyroxene separated minerals were prepared for analysis, with sample weights for each analysis around 1.7 mg. The grains were rinsed with water and ethanol, but no acid leaching was applied. Values are reported relative to V-SMOW. Two working standards, an olivine SC OLII at 0.10 ± 0.02 ‰ and a garnet G AIG at + 5.23‰ and the international biotite NBS 30 at + 5.06‰ were measured during the sample analytical run. Overall precision on standards and for samples is better than ± 0.1‰. Results are reported in Table 4.

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Author contributions

LP defined the project strategy and performed geochemical, textural and isotopic analyses. LP and GM discussed the results and wrote the manuscript.

Additional information

Competing financial interests: The authors declare no competing financial interests.
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