Magma diversity reflects recharge regime and thermal structure of the crust

Gregor Weber*, Guy Simpson & Luca Caricchi

The chemistry of magmas erupted by volcanoes is a message from deep within the Earth’s crust, which if decrypted, can provide essential information on magmatic processes occurring at inaccessible depths. While some volcanoes are prone to erupt magmas of a wide compositional variety, others sample rather monotonous chemistries through time. Whether such differences are a consequence of physical filtering or reflect intrinsic properties of different magmatic systems remains unclear. Here we show, using thermal and petrological modelling, that magma flux and the thermal structure of the crust modulate diversity and temporal evolution of magma chemistry in mid to deep crustal reservoirs. Our analysis shows that constant rates of magma input leads to extractable magma compositions that tend to evolve from felsic to more mafic in time. Low magma injection rates into hot or deep crust produces less chemical variability of extractable magma compared to the injection of large batches in colder or shallower crust. Our calculations predict a correlation between magma fluxes and compositional diversity that resembles trends observed in volcanic deposits. Our approach allows retrieval of quantitative information about magma input and the thermal architecture of magmatic systems from the chemical diversity and temporal evolution of volcanic products.

Understanding the variability and temporal evolution of erupted magma chemistry is critical to quantify magmatic and ore forming processes1–3, and to anticipate the potential future activity of volcanoes4. Detailed age-resolved geochemical records show that some volcanic centres erupt a wide variety of magma compositions, while others produce restricted chemical diversity throughout their lifetime5–12 (Fig. 1). Additionally, for systems that sample a large variability of magma types, the composition and variety of erupted magmas change over time (Fig. 1e, f; Fig. S3). Differences in compositional diversity and temporal trends between individual volcanoes have been attributed to different mechanisms. The evolving rheological properties of the crustal rocks hosting magma reservoirs can modify the capacity of magmas to rise to the surface and erupt or accumulate at depth13–16, which may impact on compositional diversity. The physico-chemical properties of the magma itself are a first order control in this respect. Density or viscosity barriers may prevent magmas of specific chemistry from erupting, leading to erupted magmas with a rather monotonous composition17,18. Such barriers might be overcome by mixing of mafic and silicic compositions10,19 or develop in parallel with the construction and destruction of large volcanic edifices20. In these models, volcanoes are depicted as physical property filters that sample only a part of the compositional spectrum present in their plumbing system. It is not clear, however, why such filters would be effective in volcanoes with restricted chemical variability and not be as effective in systems where erupted magmas exhibit a variety of compositions. Alternatively, differences in the range of erupted magma compositions could reflect contrasting recharge regimes and thermal states of crustal magma reservoirs feeding volcanoes21.

In this study we further test this hypothesis, exploring the thermochemical evolution of mid to deep crustal magmatic systems subjected to injection of hydrous basaltic magma.

Thermal and petrological modelling. To establish the link between magma recharge rates and the evolution of extractable magma chemistries in mid crustal reservoirs, we developed a numerical model that couples heat transfer and phase petrology. The design of our model is motivated by previous numerical studies22–25, petrological findings26–29 and geophysical evidence30–32 that mantle-derived hydrous basalt crystallization at mid to lower crustal depth plays a key role in generating intermediate to silicic subduction-related magma. We do not simulate magma extraction and eruption, but we trace the evolution in the chemistry of potentially extractable

Department of Earth Sciences, University of Geneva, Rue des Maraîchers 13, 1205 Geneva, Switzerland. *email: gregor.weber@unige.ch
magma (here defined as magma with < 50% crystallinity and all interstitial melt) over time, using this as a proxy for what is likely feeding shallow reservoirs of trans-crustal magma systems. Our calculations are thus quantifying the range and variability of chemical compositions available for recharge that can contribute and impact the compositional diversity sampled in eruptions fed from shallow reservoirs. To test the impact of magma and heat extraction on the thermal and chemical evolution of magma reservoirs, we performed a subset of numerical simulations in which we extract 20, 40 and 60% of magma during the assembly of the magma reservoir.

In our model, basaltic magma is successively under-accreted as cylindrical, sill-shaped bodies starting at a crustal depth of 20 or 25 km, which causes downward displacement of the floor, and construction of magmatic reservoirs in a depth range between 20 and 30 km (Fig. 2, "Methods" section). We also tested different emplacement modes, but find in agreement with previous studies that the geometry of magma injection is of secondary importance for the thermal evolution of magmatic systems (c.f. "Methods" section). Previous thermal modelling studies have shown that in order to accumulate magma bodies in the Earth crust, reservoirs have to be fed by recharge pulses in close spatial and temporal association. Such a scenario is consistent with rheological experiments and modelling, which show that the emplacement of a magma batch in the crust will attract further magma injection due to mechanical focussing of dykes and the rheological impediment exerted by the residing magma. We thus simulate repetitive injection of magma over a similar depth range. To constrain the impact of modelling variables on the thermal evolution of incrementally-built magma reservoirs.
is the total volume of extractable magma within a specified temperature range (i.e. specific chemistry) and \( V_{\text{ex}}^{T} \) is the total volume of extractable magma within the same time interval. Within this set of assumptions, we calculate the relative probability of different magma compositions to be extracted over time (Fig. 3b, d). Because in real systems magma might hybridize to various degrees by mixing or mingling processes \cite{48,49}, we also calculate the weighted average extractable magma composition (WAEMC) over the simulated time period (orange line in Fig. 3b, d). Our calculations are thus quantifying the temporal evolution of magmas that experience fractional crystallization, melt segregation and hybridization processes.

**Results and discussion**

**Temporal evolution of extractable magma chemistry.** Figure 3b shows the probability of occurrence \( P \) (Eq. 1) of chemical compositions in 20 ka steps over 200 ka for a simulation with magma flux of 0.011 km\(^3\) year\(^{-1}\) and intrusion depth of 25 km. Eruptions in the first 20 ka since the onset of magma injection in the crust have high probability of producing rhyolites, followed by basalt and minor dacite (or mixed compositions; WAEMC). The high probability of occurrence of chemically evolved magma during the early stages of evolution of a magma reservoir is driven by rapid magma cooling caused by the small system sizes and the steep tempera-
ture gradients between magma and colder crustal rocks. Such rapid cooling also causes intermediate magma composition to be present only for short timescales. With time, continued magma input into the system progressively heats up the crust. As a consequence, later during the evolution of a magmatic system, the probability of intermediate and mafic magmas to be extracted increases (Fig. 3b). Additionally, the variability of the W AEMC (Fig. 3b, d), drops progressively with time, showing that the thermal maturation of the system favours the extraction of progressively more chemically homogeneous magmas for systems that experience large-scale magma hybridization. Therefore, reservoirs assembled at the simulated conditions would initially produce recharge magmas of rather variable compositions (with W AEMC close to rhyolite), and progressively feed more mafic and less chemically variable reservoirs (Fig. 3b).

The results for a crustal magma reservoir built at 25 km depth by a volumetric magma flux of 0.0019 km³ year⁻¹ over 200 ka are presented in Fig. 3d. As in the case of the system built by relatively high magma input (Fig. 3b), in the early stages of evolution the WAEMC is mainly rhyolitic with some minor basalts. While small volumes...
Merapi (basaltic andesite; Fig. 1b) may be characterized by different magma fluxes. High magma flux regimes output basaltic andesites (Fig. 4a). The results indicate that volcanoes can erupt monotonous chemistry either in silicic-dominated magmas to intermediate chemistries, followed by a prolonged phase dominated by monotonous compositions through time, which can be attributed to progressive magma addition and heating of the crust. At these injection rates, the reservoir is mostly comprised of crystal mush for prolonged times rather than eruptible melt, which emerges transiently in association with magma recharge. Higher magma injection rates (e.g. 0.0126 km$^3$ year$^{-1}$ and 0.0180 km$^3$ year$^{-1}$), respectively. Dashed lines indicate numerical simulations run under identical conditions, but with heat and mass extraction at a rate equivalent to 20, 40 or 60% of the base model (solid lines). Simulations with heat and mass extraction (dashed curves) lead to a shift towards more silicic compositions in time.

The results show that a constant rate of magma input into a mid-deep crustal reservoir tends to generate progressively less chemically evolved WAEMC with time (Fig. 4). For example, the composition of extractable magma from a reservoir assembled with a constant magma flux of 0.0045 km$^3$ year$^{-1}$ would be rhyolitic over the first 260 ka, become dacitic over the following 240 ka, and transition to andesitic about 500 ka after the onset of magma injection (Fig. 4a). Thus, a volcano erupting over 800 ka would sample a range of different magma compositions in time.

In a subset of simulations, we tested the effect of magma (i.e. heat) extraction (e.g., due to volcanic eruptions) on the compositional evolution of WAEMC in crustal reservoirs. Generally, similar temporal trends towards more mafic compositions are observed, but for the same rate of magma input, magma extraction increases the time required for the WAEMC to become progressively more mafic. For the same rate of magma input, this timescale increases with the percentage of extracted magma (Fig. 4b). The results show that for extraction efficiencies of 20% and 40%, the difference in the 2σ SiO$_2$ range of the compositional distributions with respect to calculations performed without extraction is always smaller than 4 wt% SiO$_2$. Increasing the magma extraction efficiency to 60% results in differences in the 2σ SiO$_2$ range of the WAEMC of <5.5 wt% for intrusions built over 800 ka relative to simulations without extraction. These results indicate that the impact on compositional variability of heat and mass extraction is second order when the magmatic systems' extraction efficiency is smaller than 20%.

Our calculations also show that for the same vertical sill accretion rate, both the size of magma pulses and the injection frequency affect the compositional diversity of extractable magmas (Fig. 5). Larger and more sporadic
**Figure 5.** Impact of injection frequency on magma variability. Total alkali versus silica diagrams of the WAEMC for intrusions built with an identical vertical sill accretion rate of 0.006 m year\(^{-1}\) but different size and frequency of sill injections. Total duration of injection was set to 400 ka and an initial intrusion depth of 25 km. Symbol size is proportional to the volume of extractable magma, while the colour is proportional to time. (a) Injection of 73.5 km\(^3\) sills every 40 ka with an initial geothermal gradient of 30 °C km\(^{-1}\) and magma flux of 0.0018 km\(^3\) year\(^{-1}\) results in a broad compositional distribution. (b) Injection of 0.14 km\(^3\) in a single pulse every 845 years, initial geotherm of 35 °C km\(^{-1}\) and magma flux of 0.0001 km\(^3\) year\(^{-1}\) shows limited compositional diversity for the same vertical sill accretion rate of 0.006 m year\(^{-1}\) as in (a). Different geotherms were used in the presented simulations in order to present contrasting cases. Note that the differences in the cumulative volume of extractable magma between the two cases result from respecting the natural scaling between sill thickness and length\(^{49}\). Injection of sills with radii equal to the ones in (a) also produces less geochemical diversity compared to intrusion of thicker sills (Fig. S2).

Magma injection events (Fig. 5a) produce larger compositional diversity compared to frequent injection of small batches of magma (Fig. 5b). Such a difference is caused by the greater amplitude of heating and cooling cycles of intrusions built by larger and less frequent magma injections\(^{50}\). Hence, while a constant vertical accretion rate will produce a similar long-term evolution of the mean magma chemistry, the compositional variance in a specific period of volcanic activity increases with the size of the pulses. The spread in extractable magma compositions is also a function of the thermal state of the crust, where magma is injected. Magmatic reservoirs built in hotter crust or at deeper levels produce WAEMC of more homogenous chemistry with respect to systems assembled in shallower or colder crust, as they are thermally buffered by the temperature of the surrounding wall-rocks.

To explore the impact of variable magma flux\(^{50}\) on compositional trends and variability, we ran simulations in which the rate of magma input is increased periodically by a factor 10 for different time periods (Fig. 6). The results presented in Fig. 6a–c show that compositional variability and trends for simulations including short-lasting (e.g. 1.7 ka) increase of magma input are very similar to those without the transient increase of input and similar average rates of magma injection. This similarity remains even at higher temporal resolution than the most comprehensive geochronological studies of Arc volcanoes\(^{6,7}\). Long-lasting (e.g. 15 ka) episodes of increased magma injection rates (Fig. 6d, e) can result in differences in the chemistry of the extractable magma that are important enough to be potentially recognizable in natural datasets (Fig. 6e). This is particularly true for systems assembled by generally high magma fluxes.

**Compositional variability and recharge regimes.** The compositional changes in the WAEMC calculated with our approach over 100 to 800 ka are comparable to geochronologically constrained eruptive histories of composite volcanoes in arc settings\(^{6,7}\) (Figs. 1, S3). Reconstruction of compositional trends for individual volcanoes requires detailed fieldwork in concert with dating and geochemical analysis, which to date is only available in a limited number of case studies\(^{4,8,11,12}\). Some of these studies find no obvious variations of chemistry with time\(^{3}\), however, trends towards more silicic compositions in time\(^{31}\), as well as the reverse sense of differentiation from silicic towards mafic\(^{6,7}\) have been documented in the geological record. Although our calculations cannot capture the details of chemically resolved eruption records, they provide a framework to invert the chemistry of volcanic products (average erupted magma chemistry over time and chemical variability of eruptive products) and obtain quantitative information on fundamental parameters such as the long-term average rate of magma input in subvolcanic reservoirs.

As an example, the timescales of long-term chemical change from rhyodacitic towards basaltic andesite are about 60 ka for the well-characterized Parinacota volcano in the Central Andes\(^{5}\) (Fig. S3) and about 200 ka for the first half of the eruptive history of the Mazama-Crater Lake system in the Oregon Cascades\(^{6}\) (Fig. 1e).
direction and temporal relations of compositional change compare favourably with our modelling results in which magma reservoirs experience a temperature increase with time at a relatively constant injection rate. Our results show that extractable magma accumulating under relatively constant average rate of magma input in magma reservoirs invariably evolves towards more mafic and more homogeneous compositions in time. These general trends are also observed in model scenarios with reduced extraction probability of 2% for rhyolite compositions from highly crystallised (i.e. > 75%) magma (Fig. S5) and for simulations with removal of 20, 40 or 60% of the hottest magma. Thus, temporal trends toward more felsic erupted magma chemistry suggests a decrease of the average rate of magma input (Fig. 4). Recharge rates from the mantle may change over timescales of hundreds of thousands of years52, and a decrease of the rate of magma input or a pause in the recharge of the deep portion of the plumbing system could increase the average silica content of the volcanic products as observed for the recent history of Crater Lake and Puyehue–Cordón Caulle8 (Southern Volcanic Zone, Chile; Fig. 1).

The Age-$SiO_2$ relations of Mount Mazama-Crater Lake6 and Puyehue-Cordón Caulle8 (Fig. 1e, f) also indicate that a large variety of compositions can be erupted in close temporal association at the later stages of volcano lifecycles. While the WAEMC converges towards more homogeneous compositions with time, an increasing variety of melt compositions may coexist in the later stages of a magmatic systems lifecycle (Fig. 3a, b), implying that a large variety of compositions may be feeding systems that lack large-scale magma mixing/mingling processes. Petrological data, which can either indicate a dominant role of magma hybridization10,29 or in other cases shows limited evidence for interaction of different melts41,53 may be used to evaluate, if erupted products of a particular eruptive centre are more consistent with a WAEMC or the diversity of coexisting melts. To compare our results with a larger set of volcanic systems, we complement the compilation of long-term volcanic fluxes from Ref.8 (Fig. 7a). We assume that time-averaged erupted volumes can be used as a proxy for the rate of magma input and the time integrated erupted volumes are proportional to the sizes of the magmatic plumbing systems. Estimates of time-averaged erupted volumes are affected by preservation issues and non-linearity in volumetric eruption rates. Variations of the volumetric eruption rate can be the result of changes in the rate of magma input21, but could also reflect higher eruption efficiency of already existing reservoirs due to crustal stress field changes resulting from mass redistribution processes (e.g. cone collapse or glacial unloading;20,54–56). Nevertheless, the compilation presented in Fig. 7a shows that larger volcanoes also have higher average volumetric output rates, which confirm that long-term eruption rates can serve as a proxy for crustal magma input. Importantly, long-term volcanic output rates are biased towards lower values with increasing age of the system56. Young systems with historic or Holocene records typically show higher eruptive fluxes compared to records built on long-term
geological reconstructions. It is therefore critical to state that a comparison between compositional diversity and fluxes in the model and natural data is most robust for natural systems with a relatively long eruptive history. The collected data show a clear tendency towards higher 2σ SiO₂ ranges with increasing long-term volcanic flux (Fig. 7a), both when considering the full dataset and a subset of volcanoes filtered for similar duration of the volcanic activity (i.e. 200–500 ka). Furthermore, this analysis shows that larger volcanic systems like Santorini (~ 300 km³—11.99 wt% 2σ SiO₂) are prone to erupt higher magma diversity compared to smaller size systems like El Chichón (2.82 wt% 2σ SiO₂—26 km³). Our thermo-petrological model predicts both of these effects as reservoirs assembled at high rates of magma input accumulate W AEMC of a wide variety of compositions, which contrasts with the chemically homogeneous nature of extractable magmas in reservoirs assembled at low rates (Figs. 3a, b, 7b). Additionally, our calculations suggest that magmatic systems that are built in deeper and/or hotter crust develop less compositional diversity compared to more shallow crustal or lower geothermal gradient cases (Fig. 7b). Estimates of mid-deep crustal reservoir depths and geophysical data that could be used to constrain temperature or melt fraction are scarce. However, our results are consistent with the relationship between shear wave velocity anomalies and compositional diversity of volcanic systems in the central Cascades.
Ranges (USA; Fig. 7c). The compositionally monotonous centres Mt. Hood (4.91 wt% 2σ SiO₂) and Mt Rainer (5.71 wt% 2σ SiO₂) are underlain by a shear wave velocity decrease of 7% at a crustal depth of 30 km[57]. Mt. St. Helens (12.27 wt% 2σ SiO₂) and Mt. Adams (10.74 wt% 2σ SiO₂), at the high end of compositional diversity, show close proximity to a shallow seismic anomaly at 22 km depth[34]. On the other hand, the collected data highlight a lack of correlation between compositional variability and the lifespan of volcanic systems (Fig. 7a).

The comparison of our results with age resolved geochemical records of arc volcanoes indicates that such an approach has potential to be used to obtain first order estimates of the magma input and thermal conditions of the crust when applied to eruptive records of individual systems (Fig. 7d). Generally, volcanoes characterised by erupted magma with limited chemical variability such as Nevado de Toluca[12], El Chichón[59] and Uturuncu[11] are associated with lower magma fluxes into relatively hot or deep crustal environments. Volcanoes erupting magmas with a wide range of compositions like Parinacota[7], Mt Adams[46], Mount St Helens[45] best compare with models in which the rate of magma input is high and the subvolcanic reservoirs are assembled into relatively cold (or shallower) crust. Magma mixing, segregation and density filters have been all proposed to control the compositional variability of magmas sampled by volcanoes[19,41]. Our calculations show that the recharge rate and thermal conditions of the surrounding crust are further important variables to explain compositional variability at volcanoes. This does not rule out the potential contribution from other processes, but suggests that the thermal evolution of magma reservoirs plays a fundamental role in modulating the chemistry of volcanic products. Additionally, our results show that transition toward more mafic chemistry of volcanic products does not necessarily imply an increase in the average rate of magma input into the system. How thermally evolving crustal rheology[35–37] modulates our results is discussed in the Supplementary information.

The thermochemical model presented here permits a quantitative connection to be established between variables such as magma flux, frequency of magma injection and thermal state of the crust surrounding magmatic systems and data commonly collected for volcanic systems such as age of the eruption and major element analyses (Fig. 7d)[4–12]. This approach creates opportunities to constrain extensive variables such as magma flux, which are crucial to reconstruct the mechanisms governing the chemical and physical evolution of magmas in the crust.

Methods
We performed thermal modelling of pulsed magma injection into the mid to lower crust coupled with experimental phase relations[26], which allows us to track the temporal evolution of extractable magma chemistry. Our model design builds on the theory that silicic magmas originate at mid to deep crustal levels[22–32] and that magmatic bodies are built over protracted periods of time by incremental assembly[35,38,39].

Numerical modelling. We use the principles of heat conduction, derived by combining the conservation of energy and Fourier’s law, to describe the temporal and spatial evolution of temperature in a crust that experiences repeated magma injection. Our results were obtained by numerically solving the 2.5D (i.e., axisymmetric) formulation of the heat diffusion equation, which can be written as:

\[
\rho c \frac{dT}{dt} = \frac{1}{r} \frac{\partial}{\partial r} \left( r \frac{\partial T}{\partial r} \right) + \frac{\partial}{\partial z} \left( k \frac{\partial T}{\partial z} \right) + \rho L \frac{dT_c}{dt}
\]

where \( t \) is the time, \( T \) is the temperature, \( z \) is the vertical coordinate, \( r \) is the radial distance from the axis of rotational symmetry, \( k \) is the thermal conductivity, \( L \) is the latent heat of crystallization, \( \rho \) is the density, \( c \) is the specific heat and \( X_c \) is the fraction of crystals. Equation (2) was discretized and solved using an explicit finite difference method. Latent heating, due to the crystallization of magma, was implemented using the relation of temperature \( T \) and melt fraction \((1 - X_c)\), as derived from fractional crystallization experiments of a hydrous arc type basalt to rhyolite[26]. As \( X_c \) and \( T \) are nonlinearly dependent on each other, the governing equation was solved iteratively. The dependence of thermal conductivity \( (k) \) on temperature was implemented using the relations presented in Ref. 26 for average crust. In all simulations, zero heat flux was imposed in the perpendicular direction to all lateral boundaries of the modelled space, apart from the surface, where temperature was fixed at 8 °C. In order to test the reliability of our results, we benchmarked the model against an existing thermal code[63,64] that was solved with a different numerical method.

We model magma injection by modifying the temperature field at an initial crustal intrusion depth of 20 or 25 km, which is equivalent to increasing the initial geothermal gradient, and successive under-accretion of new basaltic sills at their liquidus temperature of 1,170 °C. In this setup, space is generated for freshly injected magma by downward advection of the host rocks. We also tested models with different emplacement geometry, in which magma was injected into the centre of the intrusion. Results obtained in this way differ from cases where under accretion is the emplacement style by higher average temperatures (Supplementary figure S1). Maximum temperature differences of about 25 °C between the different modelling geometries for a magma injection episode of 400 ka translate into 3 wt% SiO₂ difference for a case where the intrusion is at a temperature corresponding to the maximum difference in the SiO₂–T relation. This, however, does not alter the general conclusions drawn in this study.

To ensure comparability between different numerical runs, all presented results were derived from intrusions that were built by under accretion. We assumed a linear initial geothermal gradient, which was varied between 20 and 35 °C km⁻¹ for individual numerical simulations. The duration of magma injection was systematically changed between 100 and 800 ka. To test the effect of different sill dimensions, we varied the thickness and radii of injected sills between 5 to 234 m and 3 to 20 km, respectively. As sill thickness and length are not independent of each other[65], we respected the natural scaling between these parameters in most simulations. However, we
also tested the influence of decreasing thickness at constant sill radius of 10 km. For a list of modelling variables, the reader is referred to Supplementary Table S1.

To test the impact of heat and mass removal on the thermal and compositional evolution during magma body construction, we implemented the possibility of magma extraction in the model. In a subset of simulations, magma was removed at a rate that is equivalent to 20, 40 or 60% of the final intrusion volume built by a particular magma flux, by imposing an appropriate extraction rate. Extraction sites were customized to match the previous injection site and dimension, which ensures removal of magma at the highest temperature in the reservoir, by advecting the temperature field upwards. This scenario was chosen under the assumption that the most mobile magma in the reservoir is represented by the highest temperatures.

**Petrologic calculations.** To quantify the temporal evolution of extractable magma chemistry in the built intrusions we performed petrological modelling. Magas are able to flow until they reach their rheological locking point, typically at about 50 vol% crystals\(^{18} \), while interstitial melt is considered to be able to leave the rigid crystal networks beyond the locking point by various segregation processes. Thus, we defined the extractable magma compositions as magma with < 50 vol% crystals and all interstitial melt. To calculate this composition, we tracked magma volumes within temperature intervals of 50 °C between the solidus \( T_s \) at 650 °C and 1,150 °C (intervals 1 to 9), with an additional 20 °C increment between 1,150 °C and the liquidus temperature \( T_l \) of 1,170 °C (interval 10) in the numerical model as functions of time. For each temperature interval we assigned average melt and bulk magma major element oxide compositions (wt%) and average melt fraction using the petrological experiments of Ref.\(^{26} \) (Supplementary table S2). These experiments were chosen because they represent the complete fractional crystallization sequence of a typical hydrous arc basalt to rhyolite at mid to lower crustal levels (0.7 GPa) that compares well to other experimental datasets in terms of evolutionary trends in major element components. Implementing a different temperature-chemistry relation in the model would shift the calculated values in compositional space depending on the particular relation used, but leads to more mafic chemistries and more homogeneous WAEMC with time (Fig. S6), which is equivalent to our findings and does therefore not alter the conclusions drawn in this study. Magma evolutions in this study do not account for crustal melting, which has a typical productivity of < 10% for most lithologies at the crustal depth considered here\(^{16} \).

We calculated the mass of melt \( m_m \) and crystals \( m_c \) in each temperature interval along the temporal sequence by:

\[
\begin{align*}
m^i_m &= X_m V_m \rho_m, \\
m^i_c &= X_c V_m \rho_c, \\
M_i &= \sum_{i=1}^{10} m^i_m + \sum_{i=7}^{10} m^i_c
\end{align*}
\]

The chemical composition of extractable magma (\( C_{\text{extract}} \)) in terms of the composition of its major element components (\( C \)) is calculated as:

\[
C_{\text{extract}} = \frac{\sum_{i=1}^6 \frac{m^i_m C^i}{100} + \sum_{i=7}^{10} \frac{(m^i_m + m^i_c)}{100} C^i_{\text{liq}}}{M_i},
\]

where \( C^i_{\text{liq}} \) is the composition of the specific major element component at the liquidus temperature and \( i \) specifies the index for each temperature interval. Finally, the weighted average extractable magma composition (WAEMC) of the built reservoir with time in weight percent (\( C_{\text{wt}} \)) for each major element component is constrained as:

\[
C_{\text{wt}} = 100 \frac{C_{\text{extract}}}{M_i}
\]

**Data compilation.** In order to compare our modelling results to natural observations we compiled geochemical and age data for various volcanic systems in arc settings for which detailed studies of their eruption histories are available (e.g. Parinacota\(^{11} \), Mazama\(^{12} \), El Chichón\(^{13} \), Nevado de Toluca\(^{14} \), St. Helens\(^{15} \), Puyehue Cordón Caulle\(^{16} \), Uturuncu\(^{17} \), Supplementary table 4). Care was taken to exclude prominent eruptions that have been intrusions we performed petrological modelling. Magmas are able to flow until they reach their rheological locking point, typically at about 50 vol% crystals\(^{18} \), while interstitial melt is considered to be able to leave the rigid crystal networks beyond the locking point by various segregation processes. Thus, we defined the extractable magma compositions as magma with < 50 vol% crystals and all interstitial melt. To calculate this composition, we tracked magma volumes within temperature intervals of 50 °C between the solidus \( T_s \) at 650 °C and 1,150 °C (intervals 1 to 9), with an additional 20 °C increment between 1,150 °C and the liquidus temperature \( T_l \) of 1,170 °C (interval 10) in the numerical model as functions of time. For each temperature interval we assigned average melt and bulk magma major element oxide compositions (wt%) and average melt fraction using the petrological experiments of Ref.\(^{26} \) (Supplementary table S2). These experiments were chosen because they represent the complete fractional crystallization sequence of a typical hydrous arc basalt to rhyolite at mid to lower crustal levels (0.7 GPa) that compares well to other experimental datasets in terms of evolutionary trends in major element components. Implementing a different temperature-chemistry relation in the model would shift the calculated values in compositional space depending on the particular relation used, but leads to more mafic chemistries and more homogeneous WAEMC with time (Fig. S6), which is equivalent to our findings and does therefore not alter the conclusions drawn in this study. Magma evolutions in this study do not account for crustal melting, which has a typical productivity of < 10% for most lithologies at the crustal depth considered here\(^{16} \).

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\]

where \( C^i_{\text{liq}} \) is the composition of the specific major element component at the liquidus temperature and \( i \) specifies the index for each temperature interval. Finally, the weighted average extractable magma composition (WAEMC) of the built reservoir with time in weight percent (\( C_{\text{wt}} \)) for each major element component is constrained as:

\[
C_{\text{wt}} = 100 \frac{C_{\text{extract}}}{M_i}
\]
Whole rock analysis. To extend the database we present geochemical analysis of 40 new bulk-rock samples from Nevado de Toluca volcano in Central Mexico. The analysed units span the entire eruptive history of the volcano and are either stratigraphically constrained or have been dated previously by radiocarbon or 40Ar/39Ar geochronology. All samples were cleaned, washed and soaked in de-ionized water over night before they were dried in an oven at 50 °C. The dried rocks were then crushed and reduced to powders using an agate mill. The powdered material was mixed with Li-tetraborate and fused at 950 °C. Analyses for major elements were carried out at the X-ray fluorescence (XRF) laboratory at the University of Lausanne using a PANALYTICAL Axiosmax. The composition of several international reference materials (SY-2, TS-2, BHVO, NIM-N, NIM-G, BE-N) has been determined before and after sample analyses in the same analytical session.

Data availability
Data that supports the findings of this study are available within the paper and supplementary information.

Code availability
The numerical code that was used in this study is available from the corresponding author upon request.

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discussing the data and the writing of the final article. G.W. and L.C. conceived the study. G.W. wrote the thermal code with contribution from G.S. Petrological model-

Author contributions

G.W. and L.C. conceived the study. G.W. wrote the thermal code with contribution from G.S. Petrological modeling was performed by G.W. with input from L.C. G.W. wrote the manuscript draft. All authors contributed to discussing the data and the writing of the final article.

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G.W. and L.C. conceived the study. G.W. wrote the thermal code with contribution from G.S. Petrological modeling was performed by G.W. with input from L.C. G.W. wrote the manuscript draft. All authors contributed to discussing the data and the writing of the final article.

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Correspondence and requests for materials should be addressed to G.W.
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