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Strain-dependent rheology of silicate melt foams: importance for outgassing of silicic lavas

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Key Points:

- High temperature deformation experiments on cores of FOAMGLAS® explore melt foam rheological and outgassing behaviors.
- Deforming melt foams show strain weakening, hardening and localization, but undergo little or no outgassing (i.e. are impermeable).
- Crystal-free bubble-rich silicic lavas outgas inefficiently, allowing for sustained low effective viscosities and efficient transport.

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Abstract

Outgassing of volcanic systems is a ubiquitous phenomenon. Yet, the mechanisms facilitating gas escape from vesiculating magmas and lavas remain poorly understood. Pervasive outgassing is thought to depend on the efficient and abundant formation of permeable pathways. Here, we present results from experiments designed to identify the conditions and mechanisms needed to form such permeable pathways. We use a foamed silicate melt (FOAMGLAS®) in our experiments as a proxy for natural vesicular melts. FOAMGLAS® cores are compressed under a range of temperatures and strain rates, and results are evaluated against the state of melt relaxation (parameterized using the Deborah number). We find foam microstructure and rheological and outgassing behaviors evolve with strain and as a function of melt relaxation state. Relaxed melt foams harden during deformation but remain impermeable. As foams become less relaxed at higher strain rate and/or melt viscosity, they show complex responses to deformation (strain weakening and hardening) yet remain impermeable. Strain localization and formation of high permeability bands occur only in highly strained, unrelaxed foams. However, these bands are thin and oriented perpendicular to the principal stress, resulting in limited outgassing. Permeable pathways do not readily form in foams; rather high porosity melts are "persistently impermeable". Our results imply that vesicular silicic lavas may not outgas as efficiently as previously thought. Instead sustained impermeability allows the lava to maintain low effective viscosities, and to flow extended distances. In addition, pore pressures may rise in deforming impermeable lavas, perhaps priming the lava for later explosive behavior.

1 Introduction

Large rhyolitic lava flows, such as those found at Newberry Volcano (Oregon, USA), Medicine Lake Volcano and Mono-Inyo Craters (California, USA), commonly feature interiors of dense obsidian (rhyolitic glass), encased by vesicular glassy carapaces (e.g., Castro et al., 2002a; Eichelberger et al., 1986; Fink & Manley, 1987; Fink et al., 1992). The process that explains, both, the large areal extents of these silicic lavas and the characteristic abundance of dense obsidian remains a topic of vigorous debate. There have been few modern opportunities to study the eruption and emplacement of rhyolite lavas (e.g., Chaitén Volcano (Chile), 2008-2009; Castro et al., 2012b; Puyehue Cordón-Caulle volcanic complex (Chile), 2011-2012; Schipper et al., 2013). Observations made on these recent effusive eruptions suggest that the interiors of the lavas are thermally insulated and remain close to eruption temperature (e.g., Farquharson et al., 2015; Tuffen et al., 2013). However, even at eruption temperatures (~700-900°C; Farquharson et al., 2015; Schipper et al., 2013; Yoshimura et al., 2019) silicic lavas have high viscosities (>10⁸ Pa s; Giordano et al., 2008) that make it challenging to produce areally-extensive lavas of dense rhyolitic obsidian (e.g., Eichelberger et al., 1986; Fink et al., 1992).

Vesiculation resulting in high bubble contents provides a means of lowering the effective viscosity of the rhyolitic magmas (Mader et al., 2013 and references therein). For example, vesiculation within the conduit (Eichelberger et al., 1986) or syn-eruption vesiculation at the surface (Fink et al., 1992; Magnall et al., 2018) has the potential to create a gas-filled, bubble-rich silicic lava having a substantially lower viscosity than the melt. A lower viscosity could support longer transport distances of thermally insulated lavas. The characteristic dense obsidian lavas we observe, however, require that the high vesicularity lavas have lost pore fluid and have reduced porosities after flowing extended distances. This

could be achieved by efficient outgassing during emplacement, leading to bubble collapse and subsequent deflation.

Eichelberger et al. (1986) proposed that obsidian flows originally erupt as high permeability bubbly melts (i.e. "permeable foams") having low effective viscosities that outgas continuously during emplacement (Figure 1a). Alternative models suggest vesicular lavas experience outgassing by the formation of ephemeral permeable pathways – interconnected bubble channels (e.g. Okumura et al., 2009; Yoshimura et al., 2019) and/or melt fractures (e.g., Cabrera et al., 2011; Castro et al., 2012b, 2014; Kushnir et al., 2017; Okumura et al., 2013; Saubin et al., 2016; Schipper et al., 2013; Shields et al., 2016; Tuffen et al., 2003) (Figure 1b). These models envisage periods of efficient outgassing followed by collapse and annealing leading to a loss of permeability; repeated formation of ephemeral pathways eventually drain the volatiles from the system to produce a degassed, dense, obsidian-dominated lava.

Existing models for the emplacement of silicic lavas, therefore, require that the initial bubbly lavas are permeable or rapidly develop permeable pathways during flow (Figure 1). High temperature experimental studies, often using hydrated natural glasses as the starting material, have sought to identify the processes, conditions and timescales required to form permeable pathways, and to facilitate pervasive outgassing. The results from these experimental studies are diverse – in some instances, the melt walls separating bubbles remain intact until very high porosities are attained, implying bubble-rich melts remain impermeable (Burgisser et al., 2017; Lindoo et al., 2016; Ryan et al., 2015; Takeuchi et al., 2009). Conversely, other studies show bubble coalescence to occur, suggesting a means of creating transitory permeable pathways (Bagdassarov et al., 1996; Burgisser & Gardner 2005; Burgisser et al., 2017; Castro et al., 2012a; Giachetti et al., 2019; Gonnermann et al., 2017; Lindoo et al., 2016; Martel and Iacono-Marziano 2015; Stevenson et al., 1997). Simple

shearing has been shown in some experiments to increase the efficacy of coalescence, even at low porosities (e.g., Caricchi et al., 2011; Okumura et al., 2009). In addition, fracturing of the melt joins bubbles in some experiments, resulting in localized outgassing (e.g., Kushnir et al., 2017; Okumura et al., 2010; Shields et al., 2014). Finally, other experiments record vesicle shrinkage due to resorption or diffusive loss of the H₂O-fluid within the bubbles (e.g., von Aulock et al., 2017; McIntosh et al., 2014; Westrich & Eichelberger, 1994; Yoshimura & Nakamura, 2008, 2010). This wide range in behavior observed in parallel experimental studies reflects the complexities of bubble growth and loss in natural melts.

Here we present a series of high temperature deformation experiments designed to explore the rheological behavior of bubbly silicate melts with a focus on processes that allow for gas retention or loss during flow. A key component of our study is a unique experimental material that allows to circumvent many of the issues presented by the use of natural materials: FOAMGLAS® is a crystal free soda lime glass foam that is texturally homogenous, bubble-rich and, because all bubble walls are intact, impermeable (Ryan et al., 2019).

We uniaxially compressed FOAMGLAS® at high temperatures (above the glass transition temperature) to high strain at two strain rates (2.5×10⁻³ and 2.5×10⁻⁵ s⁻¹). Our experimental data comprise mechanical (stress-strain curves) and acoustic emission (AE) data collected during deformation, analyses of sample microstructure post-deformation, and measurements of final porosities and permeabilities. These data show that the impermeable melt foam cores often remain impermeable, despite being deformed to high strain (60% axial strain). High permeability compaction bands form at extreme conditions (high strain, high deformation rates and melt viscosities), but result in only limited outgassing. Permeable pathways, therefore, do not form readily in FOAMGLAS® melts subjected to the investigated experimental conditions. Our experimental data suggest that silicic lavas are

"persistently impermeable". We explore the impact of inefficient outgassing on emplacement dynamics of rhyolitic lava flows.

2 Materials

FOAMGLAS® (HLB 2400) is a high-porosity, closed-cell glass insulation manufactured by Owens Corning Corporation. Here, we include a brief description of the material and readers are referred to Ryan et al. (2019) for full descriptions of its physical, thermal and transport properties.

FOAMGLAS® has a total fractional porosity of 0.91 and is impermeable. It can be cut, cored, handled and heated to high temperature without significant micro- or macroscopic fracturing, or other changes to its physical properties. When prepared as cylindrical 2.4 cm diameter × 5.2 cm length cores, as in this study, the isolated porosity of the cores is 0.85. The difference between the total and isolated porosities (0.06) is a measure of the bubbles that intersect the surface of the core, rather than bubbles that have become connected by FOAMGLAS® production or sample preparation processes.

The solid component of FOAMGLAS® is crystal free soda lime glass, synthesized from recycled glass and other natural materials, including sand, dolomite and lime (Ryan et al., 2019). We previously measured melt viscosities (η_0) by concentric cylinder and micropenetration viscometry, and used results to fit for the adjustable parameters in the Vogel-Fulcher-Tammann (VFT) equation ($\log_{10}\eta_0 = A + B/(T - C)$), where T is temperature in Kelvin; A = -2.14, B = 4238.96, C = 498.45; Ryan et al., 2019). We use our solution to model the temperature-dependent melt viscosity of FOAMGLAS®. The modeled rheological glass transition temperature (Tg), taken as the temperature at which melt viscosity is 10^{12} Pa s, is 525° C.

Bubbles in FOAMGLAS® contain gas (>99.5% CO₂), are homogenously distributed within the glass, and are bimodal in size: large bubbles (100-275 μ m in radius) are volumetrically dominant. These bubbles are sub-rounded to polygonal in shape, and separated from one another by thin, nearly straight or slightly curved glass films. Glass walls have an average thickness of ~25 \pm 15 μ m. Numerous small bubbles (<25 μ m in radius) sit within these glass films, or at triple junctions between large bubbles. Small bubbles are round to sub-rounded. Cores of FOAMGLAS® have been shown to retain the pore gas throughout deformation (Ryan et al., 2019).

When heated to temperatures up to 600°C, FOAMGLAS® is stable – the measured dimensions, and the total and isolated porosities of cores do no change significantly as a result of heating to temperatures above Tg. However exposure to high temperatures and air will change the color of FOAMGLAS® from black to green, as trace amounts of iron in the soda lime melt oxidize. The distribution of oxidized (green) regions in cooled cores can be used to qualitatively assess whether FOAMGLAS® remained impermeable while at high temperatures, and to determine the orientation, shape and extent of any permeable pathways formed during deformation (see Section 6).

3 Methods

3.1 Deformation experiments

We performed unconfined, uniaxial compression experiments to deform FOAMGLAS® at temperatures (T; °C) between 540 and 580°C, at strain rates of 2.5×10^{-3} and 2.5×10^{-5} s⁻¹ ($\dot{\varepsilon}$; s⁻¹), and to variable axial strain (ε ; fractional) (Table 1). These experiments were performed using the low-load Volcanic Deformation Rig (VDR) at the University of British Columbia (Canada) (Quane et al., 2004; Figure S1). Samples were placed between two ceramic pistons within a tube furnace and heated at 5°C/min to the

experimental T. The furnace was controlled by one K-type thermocouple and monitored by three K-type thermocouples positioned level with the bottom, middle and top of the samples. The thermal gradient in this region (~5.2 cm) was 1-5°C (<1% experimental temperature). Samples dwelled for 45 minutes at the experimental T to ensure they were relaxed and thermally homogenous prior to deformation. Sample cores were then deformed at constant $\dot{\varepsilon}$ of 2.5×10^{-3} or 2.5×10^{-5} s⁻¹, to ε of 0.10 to 0.60. A linear variable displacement transducer (LVDT) beneath the bottom moving plate measured transient sample shortening (displacement), and a load cell (max load: 1136 kg) at the top of the sample assembly measured load throughout deformation (Quane et al., 2004; Figure S1).

Following deformation, the furnace was cooled at 5°C/min. The time between the end of deformation and cooling below Tg (525°C) was 3-11 minutes depending on the experimental temperature (i.e. samples were 15-55°C above Tg). In most experiments samples were cooled while in contact with the upper piston. In some experiments (noted in Table 1) the samples were unloaded prior to cooling, and expanded vertically in the absence of a load (Ryan et al., 2019).

In three experiments a piezoelectric transducer, part of a MISTRAS PCI-2 AE system, was attached to the VDR load cell to record the output of AE throughout deformation (Figure S1). A threshold frequency of 45 dB was used to eliminate noise from the VDR's cooling system and step motor. These data are used qualitatively, in concert with mechanical data and microstructures, to understand processes occurring within bubble walls during deformation.

3.2 Post-deformation characterization

Final sample length (l; cm) and mass (m; g) were measured following deformation (Table 1). Sample diameter (d; cm) was measured ten times along the length of the cores to

account for bulging/barreling, and then averaged (Table 1). The final dimensional volumes (volume of glass + isolated bubbles + bubbles at sample surface; V_d ; cm³) were calculated from these measurements (Table 1). The skeletal volumes (volume of glass + isolated bubbles; V_s ; cm³) were measured using a Micromeritics AccuPyc II 1340 helium pycnometer. Total (ϕ_t ; fractional) and isolated (ϕ_t ; fractional) porosities of each core were calculated from V_d , V_s and the density of the glass component of FOAMGLAS® ($\rho_{glass} = 2.41$ g/cm³; Ryan et al., 2019) (Table 1). Uncertainties in all measurements are reported in Table 1.

The total volume of bubbles in each core, or the total pore volume of each sample (V_p ; cm³), was determined from measurements of V_s , m and ρ_{glass} (Table 1). Because FOAMGLAS® has a high gas:solid ratio, the differences between initial and final ϕ_t of highly strained samples may be small (Ryan et al., 2019). The normalized change in V_p (i.e., $\%\Delta V_p = \Delta V_p/V_{p_initial} \times 100\%$), however, has the same magnitude as ε . Therefore ΔV_p is a more sensitive metric to quantify the change in total pore volume following deformation.

The permeabilities of eight deformed cores were estimated using a benchtop nitrogen permeameter at the Institut de Physique du Globe de Strasbourg (IPGS, University of Strasbourg, France) (see Heap et al. (2017a) and Kushnir et al. (2018) for permeameter details, methods and supporting theory). Six measurements were made on small cylindrical subcores (2.0 cm diameter \times 2.4-4.6 cm length) drilled from samples deformed at 10^{-5} s⁻¹ to variable strain (0.10-0.60) (Table 1). Core lengths are oriented parallel to the compression direction (σ_1) (i.e. permeability was measured parallel to σ_1). Following our previous method (1) sample surfaces were coated in silicone sealant to minimize gas flow along the irregular surface of the core, (2) a confining pressure of 1 MPa was used, (3) a transient pulse decay method was used, with an initial upstream fluid pressure of 2000 mbar and an atmospheric downstream pressure, and (4) test times were <16 hours (Ryan et al., 2019).

Two additional permeability measurements were conducted on smaller cylindrical subcore (1.1 cm diameter \times 2.0-3.0 cm length) drilled from samples deformed at 10^{-3} s⁻¹ to high strain (0.60) (Table 1; Figure S2). One of these subcores, which was drilled parallel to σ_1 , was measured using the pulse decay method described above. The other, drilled perpendicular to σ_1 , was measured using the steady-state method. This measurement was performed in the same setup described above, also under a confining pressure of 1 MPa. Six steady-state volumetric flow rate measurements (in the range 10-40 ml/min) were taken for different pressure differentials, from 10 to 40 mbar. These data were then used to calculate permeability using Darcy's law (see Heap et al. (2017a) for more details). We note that, due to the high permeability of the sample, Forchheimer correction was required to correct for the influence of turbulent fluid flow (i.e. inertial effects; Rust and Cashman, 2004) on the calculated permeability (see Heap et al. (2017a)).

Following physical property measurements, deformed cores were cut to thin wafers oriented parallel to σ_1 , and imaged using scanning electron microscopy (SEM).

4 Results of deformation experiments

4.1 High strain, variable strain rate

Figure 2a,b show mechanical data (stress (σ ; MPa) – strain (ϵ) curves) for samples deformed to variable final strain (ϵ = 0.10-0.60) at two deformation (strain) rates ($\dot{\epsilon} \sim 10^{-5} \text{ s}^{-1}$; $\dot{\epsilon} \sim 10^{-3} \text{ s}^{-1}$) at 545-555°C (Table 1). At the same $\dot{\epsilon}$ -T conditions the mechanical behavior of samples is consistent – curves overlap. Small variations in the experimental temperature (Table 1) cause curves to shift slightly up or down.

In cores deformed at a deformation rate of 10^{-5} s⁻¹ (Figure 2a) σ reaches an initial peak value at $\epsilon \sim 0.02$ after which $(0.02 < \epsilon < 0.10)$ stress decreases slightly. This indicates minor strain weakening. Strain hardening – an increase in stress with strain – occurs as deformation

continues. Curves are smooth and show no sudden stress drops. In all experiments final σ values are larger (up to 3 times) than the initial peak σ values.

In cores deformed at a deformation rate of 10^{-3} s⁻¹ (Figure 2b) σ reaches an initial peak value after loading (ε ~0.04), then shows pronounced strain weakening (0.04< ε <0.30) before showing strain hardening behavior. Repetitive stress drops are recorded at high ε (0.45< ε <0.60). In all but two high strain experiments, final σ values are less than the initial peak σ values.

In summary, the melt foam shows a complex mechanical response to deformation, including strain weakening and hardening behaviors, as well as stress drops. Foams deformed at different strain rates show different patterns in their strain-dependent behavior. Increased $\dot{\varepsilon}$ causes (1) a rise in the maximum σ value, (2) higher magnitudes and longer durations of strain weakening behavior, (3) delays in the onset of strain hardening, and (4) stress drops at high values of ε , indicating strain localization. We also note that σ measured just after loading is often the maximum value in samples deformed at high $\dot{\varepsilon}$, whereas in samples deformed at low $\dot{\varepsilon}$ the σ value measured just after loading is near the minimum.

4.2 High strain rate, variable temperature

A second set of experiments was used to establish the sensitivity of strain-dependent mechanical behavior to changes in T (i.e. melt viscosity). Cores were deformed at 10^{-3} s⁻¹, to high strain ($\varepsilon = 0.60$) over a wider range of temperatures: T = 540-581°C (Table 1).

Figure 2c shows σ - ε curves grouped by T. Where T<550°C the mechanical behavior is punctuated by stress drops during loading (ε <0.05). Following loading σ - ε curves are smooth, showing strain weakening and hardening before further stress drops are observed. At 550°C<T<560°C sample σ changes smoothly with ε until stress drops are observed. Across

the dataset, as T increases, the onset of the stress drops occurs at greater ε . No stress drops are observed in experiments at >560°C.

Within these experiments the initial peak σ and the magnitude of strain weakening decrease as T increases. At T>555°C strain hardening causes the final σ values to exceed the initial peak σ .

4.3 Acoustic emissions during deformation

In a third set of experiments we recorded AE events during deformation at 10^{-3} s⁻¹ to high strain ($\varepsilon = 0.60$). Figure 3 shows the correlation in time between recorded AE events and mechanical data (stress-time curves) for experiments performed at 548° C, 555° C and 585° C. AE events with amplitudes >75 dB (red circles; "hi-AE") correspond to sudden large stress drops. AE events with amplitudes <75 dB (white circles; "low-AE") are infrequent during the experiment conducted at high T, indicating that, in all experiments, low-AE events are not caused by the noise of the VDR (Figure S1).

At 585°C the sample shows minor strain weakening and pronounced strain hardening behavior after 0.20 strain (Figure 3a). Throughout deformation few low-AE events were recorded.

At 555°C the sample shows greater strain weakening and stress drops at high strain (Figure 3b). At 548°C strain weakening is pronounced and the stress drops occur during loading and at high strain (Figure 3c). High-AE events correspond to stress drops in the mechanical data. However low-AE events recorded during deformation are not correlated with the mechanical response of the melt foam: the number and amplitude of low-AE events reach a minimum during strain weakening and a stable maximum (plateau) at ε -0.35, after the onset of strain hardening at ε -0.25. The onset of the stress drops in the mechanical

behavior is also not preceded by a change in the number or amplitude of low-AE events (Figure 3c).

4.4 Effective and relative viscosities

We calculate the effective and relative viscosities (η_e and η_r , respectively) as a means of quantitatively comparing deformational behavior between experiments. Model values of η_e approximate the tangent to the σ - $\dot{\varepsilon}$ curve at point i and are computed as:

$$\eta_e = \frac{\sigma_i}{\varepsilon_i} = \frac{(\text{mean}(F_{i-j}, F_{i+j})/A_0)}{\left(\frac{\text{mean}(x_{i+j}, x_{i-j})/l_0}{\text{mean}(t_{i+j}, t_{i-j})}\right)}$$
Eq. 1

where F is the measured load (N), A_0 is the original contact area (m²), l_0 is the initial length (m), x is the measured displacement (m), t is elapsed time (s), t is the index, t is half the prescribed interval.

The relative viscosity (η_r) is computed as:

$$\eta_r = \frac{\eta_e}{\eta_0}$$
 Eq. 2

where η_0 is the temperature-dependent melt viscosity, predicted by the VFT equation for the melt (Section 2) at the experimental temperature (Table 2).

Values of η_e and η_r are plotted against ε in Figures 4a,b. The shapes of curves reflect the presence and magnitude of strain weakening and strain hardening, as in Figure 2. In addition, at fixed ε , η_e is greatest at low $\dot{\varepsilon}$, and decreases as $\dot{\varepsilon}$ and T increase. As a result η_r is at a minimum in samples deformed quickly. Finally, the effective viscosity of the foam is 1-2.5 orders of magnitude less than the melt viscosity.

There have been few other studies of the effect of fluid-filled isolated bubbles on the bulk viscosity of silicic (rhyolitic) magmas. Bagdassarov and Dingwell (1992) measured the effective viscosities of foamed rhyolitic melts (total porosities = 0.25-0.65) and found the

differences between the magma and melt viscosities to be 1 order of magnitude. Vona et al. (2016) also measured foamed rhyolitic melts with similar porosities (total porosities = 0.40-0.65) and calculated effective viscosities 1.1-1.6 orders of magnitude less than the melt viscosity for the imposed experimental strain rates. Finally Sicola (2018) showed less porous rhyolitic melts (total porosities = 0.15-0.40) to have bulk viscosities 0.2-1 orders of magnitude less than the melt viscosity. In all three of these studies the range of modeled effective viscosities, which were calculated using modifications of the Gent perfect slip equation rather than an incremental treatment of the σ - $\dot{\varepsilon}$ data we have used here (Eq. 1), was attributed to the differing initial porosities of the samples (Bagdassarov and Dingwell, 1992; Vona et al., 2016; Sicola, 2018). In contrast to these studies, the differences in the relative viscosity of FOAMGLAS® reflect changes in the total axial strain (Figure 4), a feature that we were only able to investigate precisely because of the homogenous nature of the sample material.

5 Physical properties of deformed cores

As FOAMGLAS® cores deform, their geometry (diameter and volume) and porosity (total and isolated) change as functions of ε , $\dot{\varepsilon}$ and T (Figure 5a,b,c, Table 1).

Sample diameters increase (i.e. samples bulge) with ε . Strain rate and temperature influence the propensity for bulging (Figure 5a, top): in samples deformed quickly (10^{-3} s⁻¹), the maximum change in sample diameter is <10%. In contrast, samples deformed slowly (10^{-5} s⁻¹) show a greater increase in sample diameter, up to 30%. At a constant strain rate cores bulge more with increasing T.

Final dimensional volumes (V_d) reflect both sample shortening (change in l is equivalent to ε) and bulging. At the same strain, samples that show the greatest increase in d

therefore show the least volume reduction (Figure 5a, bottom). The maximum reduction in V_d (~60%) occurs in samples deformed quickly at low T.

Total porosity (ϕ_t) gives the fraction of samples occupied by all pore space (isolated + connected bubbles + bubbles at sample surfaces) (Figure 5b, top). ϕ_t values decrease as ε increases. The change in ϕ_t is greatest in samples deformed quickly and at low T. The maximum reduction in ϕ_t is small (0.10), as a consequence of the high gas:solid ratio of FOAMGLAS® (Section 3.2.1). Isolated porosity (ϕ_t) also decreases systematically with ε , $\dot{\varepsilon}$ and T (Figure 5b, bottom). It is at a minimum (~0.60) in samples that showed stress drops during deformation (×'s in Figure 5).

If gas is conserved throughout deformation, the change in core volume (V_d) is a consequence of the reduction in the total volume of gas-filled bubbles in the deformed foam. Figure 5c shows the normalized change in total pore volume $(\% \Delta V_p)$ against ε . Pore volume changes as a function of ε , $\dot{\varepsilon}$ and T. The greatest reduction in $\% \Delta V_p$ occurs in samples deformed quickly. Samples deformed at low T show a greater reduction in ΔV_p than samples deformed a high T. The magnitude of $\% \Delta V_p$ is similar to ε .

Lastly, we measured the permeability of eight samples (Figure 5d; Table 1). FOAMGLAS® cores deformed to variable strain ($\varepsilon = 0.10$ -0.60) at 10^{-5} s⁻¹ and moderate T have final permeabilities below the detection limit of our permeameter, indicating permeabilities << 10^{-18} m². Subcores taken from the sample deformed at 10^{-3} s⁻¹ to high strain (0.60) were cored in two orientations (parallel and perpendicular to the compression direction). These cores show microstructural evidence of strain localization perpendicular to the compression direction (discussed in detail in Section 6). The permeabilities of subcores were << 10^{-18} m² and 7.34×10^{-13} m² when the strain localization plane was perpendicular and parallel to flow, respectively (Figure 5d). Permeability is strongly anisotropic in samples showing stress drops and evidence of strain localization (Section 6).

Cores that did not show stress drops in the mechanical data, have unoxidized interiors and show no evidence of microfracturing (Section 6). Therefore, despite being deformed to high ε , at both variable $\dot{\varepsilon}$ and T, and showing marked changes in diameter, V_d , V_p (Table 1; Figure 5a,c), our closed-cell bubble-rich melts remained impermeable during deformation.

6 Microstructure analysis

Below we compare starting material (Figure 6a) and run products (Figure 6b-h) to develop a microstructural explanation for the rheological behavior.

6.1 Geometric effects

Our experimental products show different microstructural patterns, depending on strain rate:

- (1) In samples deformed at moderate temperatures and a low strain rate (10⁻⁵ s⁻¹), bubbles are flattened parallel to, and elongated perpendicular to, σ₁ (Figures 6b,c). With increasing ε intensity of flattening/elongation increases to produce ellipsoidal to rectangular bubbles. Bubble walls perpendicular to σ₁ are elongate and straight.
 Bubble walls oriented at an acute angle to σ₁ are also nearly straight. The few walls parallel to σ₁ are bent and buckled. Small bubbles in bubble walls are still primarily subrounded to rounded, in some cases flattened perpendicular to σ₁.
- (2) Samples deformed at the same temperature, but more rapidly (10^{-3} s^{-1}) contain contorted bubbles and folded bubble walls (Figures 6d,e). At moderate ε , bubbles do not show preferred flattening/elongation directions. Many bubble walls have buckled, and folded. With increasing ε , bubbles begin to flatten perpendicular to σ_1 , and bubble walls perpendicular to σ_1 are generally straight. The remainder of bubble walls are

intensely folded. Some small bubbles in the glass walls remain rounded, but others are folded or have irregular shapes.

The evolution of bubble wall geometry depends on temperature and strain rate. At constant temperatures, strain rate controls whether bubbles are (1) regular and ellipsoidal, with straight bubble walls (low $\dot{\varepsilon}$), or (2) irregular and contorted, with folded bubble walls (high $\dot{\varepsilon}$). Similarly, where strain rate is constant, increases or decreases in temperature causes bubbles to become (1) more regular in shape (Figure 6g) or (2) contorted (Figure 6f), respectively.

The development of the two geometries described above reflect the distribution of stresses in the samples, coupled with a dependence on $\dot{\varepsilon}$ and T. Walls oriented parallel to σ_1 are under compression, and are likely to buckle. In contrast walls oriented perpendicular to σ_1 are in tension, and likely to stretch. However, as $\dot{\varepsilon}$ and T increase and decrease, respectively, buckling of walls, even oriented perpendicular to σ_1 , is more prevalent, particularly at low ε .

The shift between dominantly stretching and dominantly buckling coincides with the mechanical data and the physical properties of samples. Samples that show no strain weakening, pronounced bulging and limited volume reduction contain ellipsoidal bubbles with straight bubble walls ((1) above) and were deformed at low $\dot{\varepsilon}$, or at high $\dot{\varepsilon}$ and high T. Conversely for samples where mechanical data includes pronounced strain weakening, as well as limited bulging and significant volume reduction, bubbles have complex shapes with folded bubble walls ((2) above). These samples were deformed at high $\dot{\varepsilon}$ and moderate to low T.

It is notable that in most FOAMGLAS® samples deformation is distributed homogenously; at low to moderate strain, where strain weakening is observed, we see no zones of localized deformation. Similarly, as strain increases and hardening occurs, strain localization is not apparent in the microstructure of the samples expect those that showed stress drops (see Section 6.2).

Regardless of whether cores show stretching-dominant and buckling-dominant geometries, bubble walls are free of microfractures. This suggests deformation, involving both strain weakening and hardening, results from viscous flow of the melt component and not a consequence of brittle micromechanisms.

6.2 Causes of acoustic emissions

We saw two types of AE events recorded during deformation (Figure 2; Section 4.3). Here we identify their causes:

Low-AE events occurred throughout deformation at high $\dot{\epsilon}$ but did not correspond to strain weakening or hardening. As we observed no microfracturing of bubble walls, that is not the cause of low-AE events (Figure 6). Rather, low-AE events appear to result from the rupture of small bubbles within bubble walls (yellow outline, Figure 6h). In some samples thin films of melt ($<2~\mu m$) separating small bubbles within walls from large bubbles are distended or torn (white arrows, Figure 6h). However, the greater bubble wall is intact (Figure 6h), indicating rupturing does not result in an appreciable increase in bubble connectivity. This is confirmed by the low measured permeability of these samples (Figure 5d) To summarize, we posit low-AE events result from viscous tearing of thin films of melt.

In contrast, hi-AE events correspond well with the mechanical data (Figure 3), and evidence strain localization and fracturing of melt walls. Photographs and SEM images of samples show the orientation, size and position of strain localization bands (Figure 7a): the bands are thin and near-linear, cross-cut the sample, and are oriented perpendicular to σ_1 . In addition bands are oxidized (green). Thin "spurs" parallel to σ_1 terminate in the interior of the core. SEM images show bubble walls are disrupted in these ~500 μ m wide regions (Figure 7b). Angular fragments of glass fill the available pore space. Outside the band bubble walls remain intact (Figure 7b).

These regions of strain localization are compaction bands and comprise planar structures characterized by a lower porosity than the host rock that are oriented at high angles to the compression direction (Wong & Baud, 2012). Compaction bands are commonly observed in deformed porous materials (e.g., snow, polymers, metals, rocks), and have been shown to form as a result of the crushing of grains or the cataclastic collapse of pores/vesicles (e.g., Barraclough et al., 2017; Baud et al., 2004, 2006; Elliot et al., 2002; Fortin et al., 2006; Gibson & Ashby, 1999; Heap et al., 2015b, 2017b; Issen et al., 2005; Papka & Kyriakides, 1999; Wong et al., 1997, 2001; Wong & Baud 2012). The formation of compaction bands is often attended by stress drops and bursts of AE activity (Figures 2,3) (e.g., Baud et al., 2004, 2006; Heap et al., 2015a; Wong et al., 1997).

7 Discussion

Strain-dependent rheological behavior is relevant to the emplacement of silicic lavas – as lavas flow their response to deformation evolves and alters outgassing behavior (e.g., Kendrick et al., 2013; Shields et al., 2014). Below we explore the causes of strain-dependent rheological behaviors and their influences on outgassing mechanisms and efficiency.

7.1 Melt foam rheology

The mechanical behavior and microstructures of our experimental products change systematically with $\dot{\varepsilon}$ and T. During deformed at high strain rates bubble become irregularly shape, bubble walls fold, strain weakening occurs and the foam is "soft" (low η_r); when deformed at low strain rates bubbles remain ellipsoidal, bubble walls are straight to curvilinear, no strain weakening occurs and the composite is relatively "stiff" (high η_r); increasing temperature reduces the magnitude of melt wall buckling and strain weakening, which causes the relative viscosity of the composite to rise.

In bubble suspensions, including many vesicular magmas and lavas, rheological behavior can be understood as a function of the steadiness of flow and the relaxation state of the bubbles (Mader et al. (2013) and references therein). In these materials, melt viscosity, bubble radius and the interfacial tension between the fluid in bubbles and the melt (Γ , N/m) dictate the bubble relaxation time ($\lambda_b = (\eta_0 r) / \Gamma$; s). Where λ_b is less than the deformation time (related to the strain rate, $\dot{\varepsilon}$) interfacial forces act to reduce the surface energy and return bubbles to spherical shapes. At this condition, where interfacial forces dominate (i.e. where the capillary number ($Ca = \lambda_b \dot{\varepsilon}$) << 1) spherical bubbles hinder viscous flow of the melt (Manga et al., 1998; Llewellin et al., 2002; Rust and Manga, 2002, Mader et al., 2013). In contrast, where λ_b is greater than the deformation time (i.e. Ca >> 1), the interfacial forces are low, and the dominating viscous forces deform the bubble. The change in bubble shape accompanying deformation introduces slip surfaces that allow the liquid to flow more easily (Manga et al., 1998; Llewellin et al., 2002; Rust and Manga, 2002, Mader et al., 2013).

However, in melt foams ($\phi_t > 0.74$; Mader et al., 2013) where the total volume of gas present does not allow for the "packing" of spherical bubbles, the structure of the bubbly melt is different – "cells" of one fluid (here, CO_2 gas) are surrounded by thin, near planar films of a second fluid (here, silicate melt). This change in fluid structure, in turn, changes the rheological response (Mader et al., 2013). It is currently unknown how the relaxation state of the bubbles and steadiness of flow may influence foam rheology. Therefore we use studies of other cellular solids and foams to interpret our results.

Studies of the mechanics of elasto-plastic/viscoelastic cellular solids/foams (metals, polymers) have shown, where deformation is homogenously distributed, changes in the geometry of cell walls introduce strain weakening and hardening behaviors (e.g., Gibson & Ashby, 1999; Rashed et al., 2016). These studies identify three phases of compressive deformation: (1) elastic deformation during loading, (2) non-recoverable cell wall stretching

and/or buckling, and (3) densification at high strain. In the second phase, stretching vs. buckling impart distinct σ - ϵ patterns (e.g., Mu et al., 2010; Rashed et al., 2016; Scheffler & Colombo, 2005): where stretching is dominant, σ - ϵ curves reach a plateau following loading, but where bending/buckling dominate, strain weakening occurs as deformation proceeds. During densification, cell volumes decrease, bringing opposing or adjacent cell walls in to contact (e.g. Gibson & Ashby, 1999). During this final phase, increasing force is required for deformation to continue, resulting in strain hardening (Gibson & Ashby, 1999; Mu et al., 2010; Rashed et al., 2016; Scheffler & Colombo, 2005).

The relationship between FOAMGLAS® microstructure and rheological behavior therefore reflects the propensity for melt wall stretching or buckling. Where melt walls primarily stretch, a network of straight bubble walls form, producing a relatively stiff (high η_r) foam. Deformation proceeds at a steady-state until bubble walls are close enough to impinge on one another, at which point strain hardening begins. At conditions where melt walls primarily buckle rather than stretch, the foam is a network of folded bubble walls that is initially soft (low η_r). As it is compressed, bubble walls continue buckling, leading to strain weakening. Only when bubble walls interact does the foam harden.

The Deborah number (De) integrates the effects of T and $\dot{\varepsilon}$ into a single parameter and is used, here, to delineate regimes for stretching vs. buckling of bubbles. Because the strain-dependent rheological behavior appears to be dictated by the compliance of the melt rather than to the behavior of the bulk foam material, we define De as the ratio of the Maxwell relaxation time of the melt ($\tau = \eta_0/G$), where G is the elastic shear modulus ($\sim 10^{10}$ Pa; Dingwell & Webb, 1990) to the observation (or deformation) time (t) (e.g., Poole, 2012). Thus it is a measure of the competition between the capacity of a melt to dissipate stress viscously (relax) and the rate at which deformation occurs:

$$De = \frac{\tau}{t} = \frac{\dot{\varepsilon} \eta_0}{G}$$
. Eq. 3

Low De values correspond to effective dissipation of stress and melt relaxation. As η_0 and/or $\dot{\varepsilon}$ increase, viscous flow is hindered and stress accumulates in the melt. We adopt $De \sim 10^{-2}$ as the transition from relaxed to unrelaxed states (i.e., Dingwell, 1996; Gonnermann & Manga, 2003; Papale, 1999; Wadsworth et al., 2018).

In Figure 8 we plot η_r against modeled De. Representative mechanical data and microstructures are shown across our range of De values. There is a systematic relationship between De, bubble wall geometry, bubble shape, and η_r , indicating the propensity of bubble walls to stretch rather than buckle is controlled by the melt relaxation state: at low De numbers, the melt relaxes easily during deformation. At these conditions compliant bubble walls can stretch rather than buckle. In contrast, a reduced capacity to dissipate stress (i.e. higher De numbers) means bubble walls are less relaxed, less compliant, and buckle and fold rather than stretch.

The strain-dependent behavior of the foam also varies a function of De. In particular strain weakening behaviors begin when $De > 10^{-3}$ (Figure 8). This is particularly consequential because the onset of strain weakening has been posited to create a feedback in the deformational behavior (Caricchi et al., 2011; Okumura et al., 2010; Wright & Weinberg, 2009): as lava weakens during deformation, the same applied stress will promote higher local strain rates and greater strain (e.g. Vona et al., 2016). Localization and intensification of deformation can lead to a reduction in effective viscosity, facilitating further flow (Vona et al., 2016; Wright & Weinberg, 2009). Studies of experimental and natural materials have shown localized high strain rates can enhance bubble coalescence (Caricchi et al., 2011; Laumonier et al., 2011). Similarly, localization of deformation and concomitant increases in strain rate can force melt across the glass transition causing brittle disruption of the melt and allowing outgassing (e.g., Okumura et al., 2010, 2013; Wright & Weinberg, 2009).

In our study the onset of this weakening and localization behavior is evidenced by the high permeability (relative to the host material) compaction bands formed in highly strained, unrelaxed foams (Figure 8): the compaction bands result when strain rates localized at bubble walls exceed the relaxation rates, causing fracture of the bubble walls (e.g. Dingwell & Webb, 1990). Characteristically of compaction bands in many porous materials (including rocks, metals, snow, polymers; see Guillard et al. (2015) for extended discussion of compaction bands in brittle porous media), planar bands of localized strain are oriented perpendicular to σ₁, in this case as a result of the failure of melt walls, which propagated laterally (Figure 7b). This micromechanism is similar to the cataclastic pore collapse described for porous limestones (e.g., Zhu et al., 2010) and volcanic rocks (e.g., Heap et al., 2015b, 2017b; Zhu et al., 2011). It is notable that compaction band formation in porous rocks typically requires high effective pressures (e.g., Wong & Baud, 2012). The development of compaction bands deformed under uniaxial conditions, observed here and in other materials (e.g., polycarbonate honeycomb (Papka & Kyriakides, 1999), closed-cell aluminum alloy foams (Bastawros et al., 2000)), may be a peculiarity of having a high total porosity (>0.74).

With strain weakening beginning at $De>10^{-3}$, we can estimate that under a constant applied stress a feedback loop of weakening and effective viscosity reduction with increasing strain only needs to result in a 10-fold increase in local deformation rate to drive a melt, including a natural silicic lava, to an unrelaxed state. At this condition brittle failure of bubble walls is feasible, and the outgassing behavior changes.

7.2 Persistent impermeability and its consequences

Most of the experimental products remained impermeable despite (1) being deformed to high strain, (2) at both low and high strain rates, and (3) over melt viscosities spanning 1.5

orders of magnitude. These results speak to the surprising ability of melt foams to resist the formation of permeable pathways.

In FOAMGLAS®, permeability increase only occurs when strain weakening and localization produce compaction bands in unrelaxed foams at high strain (Figures 5d,7). However, bubble walls are intact outside compaction bands – the remainder of the sample does not communicate with the band, and is impermeable. In porous rocks, compaction bands typically form barriers to fluid flow (e.g., Baud et al., 2012; Heap et al., 2015b; Wong et al., 2001; Wong & Baud, 2012). For example, in porous sandstones, the compaction band is of lower permeability than the high permeability host rock (e.g. Baud et al., 2012). However, because FOAMGLAS® is initially impermeable, compaction bands in our melt foam create a high permeability channel and allow for fluid flow along their length. So, although compaction bands represent areas of lower porosity (Figure 7b), they are also zones of higher vesicle connectivity (connectivity is zero in the remainder of the material), and are conduits for rather than barriers to flow.

Our results support the model that ephemeral permeable pathways may outgas silicic lavas (Figure 1b). Previous work has shown that this can be accomplished in simple shear by the coalescence of bubbles to form interconnected bubble networks (Caricchi et al., 2011; Okumura et al., 2009). In domains within lavas that undergo greater compaction (pure shear) than simple shear (e.g., Befus et al., 2015; Castro et al., 2002b; Shields et al., 2016), compaction bands may also result and serve as permeable pathways. However the "extreme" flow conditions required for the formation of compaction bands (high strain, unrelaxed melt) makes them unlikely to form throughout flows. In addition, the permeability that has developed is anisotropic and is dictated by the orientation of σ_1 (permeability is still within error of zero perpendicular to the compaction band; Figure 5d). The efficacy of outgassing through compaction bands in a melt foam or lava will therefore depend on the orientation of

the principal stresses, as well as the size and lateral extent of these bands, and connectivity to other fracture systems or the greater atmosphere (e.g., Castro et al., 2012b; Heap & Kennedy, 2016; Kushnir et al., 2017; Okumura et al., 2010; Shields et al., 2016).

The restricted flow conditions required for strain localization to occur, and the propensity of outgassing pathways to form with a particular orientation to the principal stresses are not the only hindrances to pervasive outgassing: both melt fractures and interconnected bubble networks are susceptible to closure and rapid healing (e.g., Kendrick et al., 2016; Kolzenburg et al., 2012; Lindoo et al., 2016; Rust & Cashman, 2011; Shields et al., 2016; Tuffen et al., 2003; Yoshimura & Nakamura, 2010). This suggests other processes, such as diffusive dehydration and volatile resorption (e.g., von Aulock et al., 2017; McIntosh et al., 2014; Ryan et al., 2015; Westrich & Eichelberger, 1994; Yoshimura et al., 2019; Yoshimura & Nakamura, 2008) may be important mechanisms by which lavas, particularly silicic lavas, can outgas and densify (Figure 9a).

In the absence of efficient outgassing, fluids do not have a means of escaping the deforming melt foam. We see this in our experiments: during deformation, the CO_2 contained in bubbles in FOAMGLAS® can either leave (i.e. outgas), or remain within bubbles. While compaction bands allow for some gas escape, gas loss can only occur by diffusive processes for the remainder of FOAMGLAS® samples. However, we have observed the expansion of deformed samples unloaded while above Tg (Section 2; Ryan et al., 2019). This indicates a pore pressure greater than atmospheric pressure, indicating gas is conserved in our samples. Therefore when bubbles in our samples are compressed during deformation, their volume change results in an increase in pore pressure, making ΔV_p an indicator of pore pressure changes.

Figure 8 shows the relationship between the change in total pore volume (ΔV_p) and De, as well as representative core geometries. At low De conditions, where melt walls

primarily stretch in response to deformation, cores bulge. Because the foam is translated laterally, in addition to being deformed, the resulting reduction in pore volume is ~40%. In contrast, at higher De conditions, melt walls primarily buckle and cores are deformed with limited bulging. This core geometry produces a larger reduction in total pore volume, up to ~65%. Therefore, core geometry and ΔV_p (and pore pressure development) are governed by the state of melt wall

relaxation. In unrelaxed melt foams ($De>10^{-2}$) the reductions in ΔV_p with strain is more pronounced and the rise in pore pressure is greater as deformation continues.

Applying our experimental insights to Nature, suggests that highly vesicular silicic lavas and magmas will retain their bubbles for extended times and have the potential to become overpressurized as a result of continued deformation (Figure 9b). This impermeable state will be maintained unless: (1) specific flow conditions (high strain rates, low temperatures, high strain) are reached supporting formation of numerous permeable pathways, or (2) diffusive dehydration and resorption can moderate pore pressures.

7.3. Implications for outgassing and explosivity of rhyolitic lavas

We have shown that during compression at flow conditions expected for silicic lavas, outgassing of melt foams is inefficient – the foam does not transition from an impermeable to a permeable one, melt viscosities are not low enough to support coalescence (e.g., Lindoo et al., 2016; Castro et al., 2012a), and few permeable pathways form as a result of brittle behavior. Existing models for outgassing of silicic lavas (Figure 1) therefore appear to overestimate the outgassing potential of bubbly lavas.

The eruption of rhyolitic lava from Puyehue Cordón-Caulle (Chile) from 2011-2012 (Schipper et al., 2013; Tuffen et al., 2013) provides some evidence that lavas outgas very inefficiently. The lavas inflated during eruption, flow and post-eruption (Farquharson et al.,

2015). Inflation, driven by late-stage vesiculation (i.e. volatile exsolution, and bubble nucleation and growth occurring at Earth's surface), led to the formation of breakouts, and the progressive morphological change of the lava flow up to a year after the end of effusion (Farquharson et al., 2015; Magnall et al., 2018). The post-eruption growth of the lava flow and the long run-out distance are aided by the insulating properties of the flow surfaces (e.g., Farquharson et al., 2015; Magnall et al., 2018; Tuffen et al., 2013) but also could be facilitated by the long-term retention of bubbles: if lavas can flow for extended time without significant changes to their porosities, they maintain low effective viscosities orders of magnitude less than the viscosity of the melt. Sustained low effective viscosities promote efficient transport, allowing bubbly lavas to continue to deform and flow to produce extensive deposits.

Similarly, Holocene rhyolitic lavas preserve evidence of inefficient outgassing and the accumulation of trapped volatiles: for example, gas cavities and explosive craters mark the surfaces of rhyolitic flows hundreds of meters away from the vent (e.g. Big and Little Glass Mountains (Medicine Lake Volcano; California, USA), Big Obsidian Flow (Newberry Volcano; Oregon, USA)) (Castro et al., 2002a; Fink et al., 1992; Fink & Anderson, 2017; Fink & Manley, 1987). These features are interpreted as evidence of the retention and, in some cases, explosive release of pressurized gases trapped in the stalled and cooling lavas (Castro et al., 2002a; Fink et al., 1992; Fink & Manley, 1987). As such, they highlight the roles deformation and inefficient outgassing may play in causing hazardous behavior, including far-field explosivity. In addition, many of these Holocene flows contain both bands of dense obsidian, and thick layers of vesicular lavas (Fink & Manley, 1987; Fink & Anderson, 2017) (see photos in Figure 9). These vesicular units, which can comprise up to 40% of the total flow volume in some large obsidian flows (Fink & Manley, 1987), frequently contain isolated bubbles, attesting to the long-term retention of bubbles.

Finally our interpretation of outgassing dynamics in melt foams offers additional clarity to the observation of hybrid volcanic activity – the contemporaneous effusion of lava and explosions – seen at the recent rhyolitic eruptions at Chaitén (2008-2009; Castro et al., 2012b) and Puyehue Cordón Caulle (2011-2012; Schipper et al., 2013). Outgassing at these volcanoes is thought to be facilitated by the formation of fractures, which subsequently clog with pyroclastic debris and weld shut (e.g., Castro et al., 2012b; Saubin et al., 2016), rather than through a permeable foam. The "persistent impermeability" of high-porosity foams, described herein, may explain why volatiles are forced to outgas through fracturing as soon as the pore overpressure is sufficient to fracture the magma.

In conclusion, we see from our experiments that melt foams do not undergo pervasive outgassing at conditions relevant to the flow of silicic lavas, and posit that natural lava flows remain gas-charged for longer than expected. Consequences of inefficient outgassing include extended run-out distances for lavas and the potential for far-field explosive behavior.

8 Conclusions

We deformed a homogenous, impermeable analogue melt foam to explore the interplay between rheological behavior, microstructure and outgassing, as functions of strain, strain rate and temperature. We find that much of the deformational response of the closed-cell melt foam is dictated by the geometry of the melt walls, which is controlled by the capacity of the melt to dissipate stress and relax (i.e. *De*). The onset of strain weakening, driven by melt wall buckling, creates the potential for run-away deformation and can result in the formation of localized high permeability zones.

Pervasive outgassing did not occur in our experiments, though we deformed melt foam samples over a range of melt viscosities (Giordano et al., 2008) and strain rates applicable to natural lavas (Tuffen et al., 2003). Our results suggest permeable pathways may

not readily form in silicic lavas. Inefficient outgassing on the surface allows lavas to remain gas-charged for extended times, supporting longer flow distances, and perhaps feeding far-field explosivity. In the conduit, the inability to outgassing efficiently leaves the magma susceptible to pore pressurization and fragmentation in the subsurface.

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 Table 1. Experimental Conditions and Physical Properties of Experimental Products.

			_		Change ^a					Final ^b						
	#	T (°C)	ε	Δl (cm)	1σ	Δd (cm)	1σ	ΔV_d (cm ³)	p.u.	ϕ_t	p.u.	ϕ_i	p.u.	V_p (cm ³)	p.u.	$k (m^2)^c$
	strain ra	ate = 2.5	\times 10 ⁻⁵ s ⁻¹													_
	116 ^d	546	0.10	-0.52	0.00	0.04	0.03	-1.77	0.49	0.91	0.02	0.84	0.02	19.73	0.02	-
	110 ^d	552	0.10	-0.54	0.00	0.06	0.03	-1.53	0.52	0.91	0.02	0.84	0.02	19.76	0.03	<<10 ⁻¹⁸
	340	554	0.10	-0.53	0.00	0.06	0.03	-1.48	0.57	0.92	0.02	0.84	0.02	20.11	0.02	-
	121 ^d	545	0.20	-1.01	0.00	0.08	0.06	-3.49	0.94	0.90	0.04	0.85	0.04	18.52	0.02	-
	108 ^d	553	0.20	-0.98	0.00	0.11	0.07	-2.85	1.16	0.90	0.05	0.83	0.05	18.35	0.03	<<10 ⁻¹⁸
	343	555	0.20	-1.08	0.00	0.16	0.06	-2.55	1.00	0.91	0.04	0.84	0.04	18.93	0.02	-
	117 ^d	546	0.30	-1.42	0.00	0.15	0.06	-4.26	0.98	0.90	0.05	0.82	0.05	15.85	0.02	-
	106 ^d	549	0.30	-1.45	0.00	0.16	0.08	-4.65	1.22	0.89	0.05	0.84	0.05	17.37	0.04	<<10 ⁻¹⁸
	104 ^d	546	0.40	-1.92	0.00	0.18	0.12	-6.86	1.65	0.88	0.08	0.84	0.08	15.43	0.03	<<10 ⁻¹⁸
	118 ^d	548	0.40	-1.79	0.01	0.20	0.13	-5.83	1.84	0.88	0.08	0.85	0.08	16.36	0.02	-
	341	557	0.40	-2.10	0.00	0.39	0.19	-5.23	2.63	0.90	0.12	0.83	0.12	16.66	0.03	-
	115 ^d	543	0.50	-2.28	0.00	0.27	0.14	-7.39	1.70	0.88	0.09	0.83	0.09	13.95	0.02	<<10 ⁻¹⁸
	120 ^d	547	0.50	-2.30	0.00	0.34	0.19	-6.79	2.41	0.88	0.12	0.82	0.12	14.92	0.02	-
	112 ^d	545	0.60	-2.61	0.01	0.42	0.17	-7.80	2.08	0.87	0.11	0.79	0.11	13.60	0.04	<<10 ⁻¹⁸
	119 ^d	547	0.60	-2.55	0.01	0.38	0.16	-7.54	1.90	0.88	0.10	0.82	0.10	13.27	0.03	-
	337	554	0.60	-3.13	0.00	0.68	0.26	-8.96	2.71	0.86	0.14	0.77	0.14	12.57	0.03	-
$strain\ rate = 2.5 \times 10^{-3}\ s^{-1}$																
	316	551	0.10	-0.52	0.00	0.06	0.03	-1.36	0.50	0.90	0.03	0.83	0.03	19.59	0.02	-
	339	554	0.10	-0.51	0.00	0.06	0.03	-1.32	0.60	0.92	0.03	0.83	0.03	19.83	0.03	-
	322	554	0.20	-1.09	0.00	0.12	0.05	-3.32	0.78	0.88	0.03	0.81	0.03	17.82	0.02	-
	338	555	0.20	-1.07	0.00	0.11	0.04	-3.42	0.63	0.90	0.03	0.82	0.03	17.88	0.03	-
	335	552	0.30	-1.59	0.00	0.12	0.06	-5.95	0.83	0.90	0.04	0.80	0.04	15.11	0.03	-
	323	553	0.30	-1.59	0.00	0.11	0.03	-6.03	0.43	0.88	0.02	0.80	0.02	15.06	0.02	-
	333	554	0.40	-2.07	0.00	0.15	0.08	-7.29	1.04	0.88	0.06	0.78	0.06	11.92	0.02	-

328	554	0.40	-2.06	0.00	0.18	0.05	-6.96	0.67	0.88	0.04	0.76	0.04	12.25	0.01	-
329	554	0.50	-2.57	0.00	0.18	0.07	-9.65	0.71	0.85	0.04	0.74	0.05	9.86	0.02	-
332	554	0.50	-2.56	0.00	0.18	0.08	-9.58	0.84	0.85	0.05	0.75	0.05	9.99	0.02	-
348	551	0.60	-3.09	0.00	0.26	0.04	-12.39	0.39	0.81	0.02	0.61	0.02	7.48	0.03	-
330	553	0.60	-3.07	0.01	0.24	0.09	-11.65	0.78	0.82	0.06	0.67	0.06	7.63	0.03	-
308	554	0.60	-3.04	0.00	0.16	0.04	-13.31	0.40	0.81	0.02	0.70	0.02	8.15	0.02	-
336	554	0.60	-3.06	0.00	0.19	0.08	-13.34	0.71	0.84	0.05	0.68	0.05	8.14	0.02	-
361 ^e	553	0.60	-	-	-	-	-	-	-	-	-	-	-	-	$<<10^{-18}\mathrm{f}$
362 e	555	0.60	-	-	-	-	-	-	-	-	-	-	-	-	7.3×10 ⁻¹³
strain	rate = 2.5	$5 \times 10^{-3} s$	-1												
311 e	540	0.60	-	-	-	-	-	-	-	-	-	-	-	-	-
353 ^e	543	0.60	-	-	-	-	-	-	-	-	-	-	-	-	-
359	546	0.60	-3.08	0.00	0.19	0.11	-13.27	0.99	0.81	0.07	0.62	0.07	7.28	0.02	-
346	549	0.60	-3.09	0.00	0.20	0.16	-13.25	1.44	0.81	0.10	0.64	0.10	7.72	0.06	-
354	557	0.60	-3.08	0.00	0.25	0.06	-12.75	0.58	0.83	0.04	0.69	0.04	8.47	0.08	-
355	561	0.60	-3.08	0.00	0.23	0.05	-12.86	0.49	0.83	0.03	0.73	0.03	8.85	0.01	-
351	563	0.60	-3.07	0.00	0.31	0.05	-12.10	0.52	0.84	0.03	0.72	0.03	9.34	0.08	-
350	566	0.60	-3.09	0.00	0.35	0.07	-11.92	0.70	0.83	0.04	0.73	0.04	9.64	0.08	-
356	570	0.60	-3.08	0.00	0.32	0.07	-12.17	0.66	0.84	0.04	0.76	0.04	9.80	0.02	-
349	572	0.60	-3.10	0.00	0.41	0.11	-11.16	1.05	0.82	0.06	0.74	0.06	10.08	0.03	-
357	575	0.60	-3.09	0.00	0.33	0.12	-12.04	1.12	0.83	0.07	0.79	0.07	10.22	0.02	-
358	577	0.60	-3.10	0.00	0.42	0.19	-11.30	1.80	0.84	0.11	0.80	0.11	11.09	0.02	-
352	581	0.60	-3.10	0.00	0.56	0.09	-9.90	0.90	0.86	0.05	0.76	0.05	11.47	0.08	-

Note. Including temperature (T), axial strain (ϵ ; fractional) and strain rate. Physical properties of experimental products, including changes in length, diameter and dimensional volume (Δl , Δd and ΔV_d , respectively). Also final measured total and isolated porosities (ϕ_t and ϕ_i , respectively; fractional), cumulative pore volume (V_p), and permeability (k). Standard deviations (1 σ) and propagated uncertainties (σ) are included.

^aMeasured with digital calipers. Average initial l, d, and V_d are 5.19 ± 0.02 cm, 2.46 ± 0.04 cm and 24.73 ± 0.80 cm³, respectively. $^b\phi_t = 1 - 0.02$ cm, $^a\phi_t = 1.02$ cm, $^a\phi_t = 1$

 $(m/V_d \, \rho_{glass})$, where m is the mass (g), V_d is the dimensional volume measured by calipers, and ρ_{glass} is the density of the glass (2.41 g/cm³); $\phi_i = (V_s \, \rho_{glass} - m) / (V_d \, \rho_{glass})$, where V_s is the skeletal volume measured by helium pycnometry; $V_p = V_s - (m/\rho_{glass})$. Average initial ϕ_t , ϕ_t , and V_p are 0.91 ± 0.006 , 0.85 ± 0.003 and 21.03 ± 0.66 cm³, respectively. Detection limit of permeameter is 10^{-18} m². See text for details. Samples unloaded after deformation but before cooling ("rebound" samples). Samples not included in quantitative analyses. Permeability measured perpendicular and parallel to plane of strain localization.

Table 2. Modeled Viscosities.

#	T (°C)	ε	$log_{10}\eta_0^a$	$\log_{10} \eta_e^{\ b}$	$\log_{10} \eta_r^{\ c}$				
$strain\ rate = 2.5 \times 10^{-5}\ s^{-1}$									
116	546	0.10	11.1	9.2	-1.9				
110	552	0.10	10.8	9.2	-1.7				
340	554	0.10	10.8	9.0	-1.7				
121	545	0.20	11.1	9.3	-1.8				
108	553	0.20	10.8	9.2	-1.6				
343	555	0.20	10.7	9.1	-1.7				
117	546	0.30	11.1	9.4	-1.7				
106	549	0.30	11.0	9.4	-1.6				
104	546	0.40	11.1	9.5	-1.6				
118	548	0.40	11.0	9.5	-1.5				
341	557	0.40	10.6	9.3	-1.3				
115	543	0.50	11.2	9.6	-1.6				
120	547	0.50	11.0	9.6	-1.5				
112	545	0.60	11.1	9.8	-1.4				
119	547	0.60	11.0	9.7	-1.3				
337	554	0.60	10.8	9.7	-1.1				
strain rate =	$= 2.5 \times 10^{-3}$	s^{-1}							
316	551	0.10	10.9	8.8	-2.0				
339	554	0.10	10.8	8.6	-2.2				
322	554	0.20	10.8	8.7	-2.0				
338	555	0.20	10.7	8.5	-2.2				
335	552	0.30	10.8	8.5	-2.3				
323	553	0.30	10.8	8.6	-2.2				
333	554	0.40	10.8	8.6	-2.2				
328	554	0.40	10.8	8.6	-2.1				
329	554	0.50	10.8	8.8	-2.0				
332	554	0.50	10.8	8.7	-2.1				
348 ^d	551	0.60	10.9	9.0	-1.9				
330 ^d	553	0.60	10.8	8.7	-2.1				
308 ^d	554	0.60	10.8	8.8	-2.0				
336 ^d	554	0.60	10.8	8.8	-2.0				
	$= 2.5 \times 10^{-3}$.								
354 ^d	557	0.60	10.6	8.6	-2.0				
355	561	0.60	10.5	8.6	-1.8				
351	563	0.60	10.4	8.6	-1.8				
350	566	0.60	10.3	8.6	-1.7				
356	570	0.60	10.2	8.4	-1.7				
349	572	0.60	10.1	8.5	-1.5				
357	575	0.60	10.0	8.3	-1.7				
358	577	0.60	9.9	8.3	-1.6				
352	581	0.60	9.8	8.2	-1.6				

Note. Melt viscosity (η_{θ}) and final effective and relative viscosities $(\eta_{e}$ and

 $\eta_{\it r}$). Samples that fractured during loading are not included.

 a log₁₀ η_0 = -2.14 + 4238.96 / (T(K) - 498.45) (Ryan et al., 2019). b At final ϵ , or just prior to first stress drop. $^c\eta_r = \eta_e$ / η_0 . d Samples show stress drops at high ϵ .

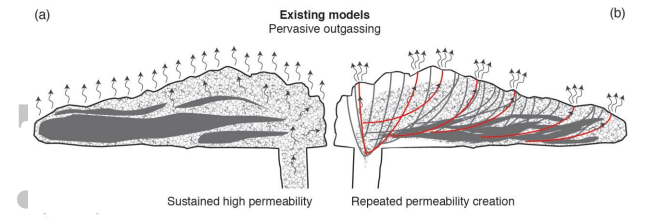


Figure 1. Existing models for outgassing of silicic lavas. (a) "Permeable foam" model put forward by Eichelberger et al. (1986): lava vesiculates in the subsurface to form a "permeable foam". Sustained high permeability supports continuous outgassing (arrows) as the lava exits the conduit and flows across the surface. Eventually sufficient outgassing reduces lava porosity to create dense, non-vesicular regions in the lava (obsidian; grey regions). (b) Formation of ephemeral permeable pathways: lava vesiculates near or at the surface. In contrast to (a), the foam is not highly permeable. Permeable pathways (e.g. interconnected bubble channels, melt fractures) form (red lines) and collapse (grey lines), allowing for cyclic outgassing (arrows) (e.g., Castro et al., 2012b; Yoshimura et al., 2019). Volatiles are drained from the lava to produce non-vesicular zones.

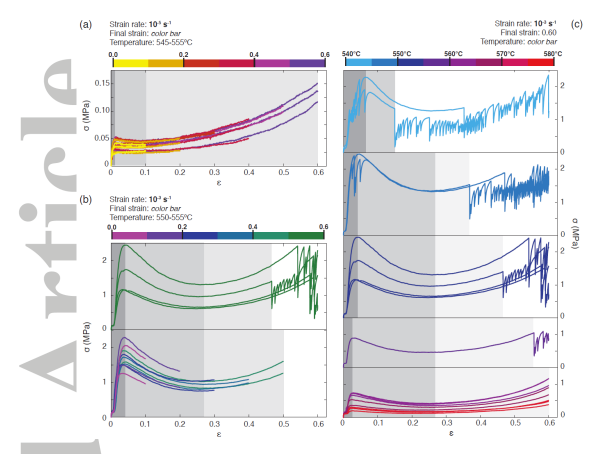


Figure 2. Mechanical data from deformation experiments plotted as load stress (σ) vs. instrumental strain (ϵ). Grey panels demarcate periods of loading (dark grey), strain weakening (medium grey) and strain hardening (light grey). (a) High strain experiments conducted at 545-555°C at 2.5×10^{-5} s⁻¹ to variable final strain (color bar). The initial ($\varepsilon < 0.02$) steep rise in stress is an experimental artifact resulting from sample loading. The load stress curve has an initial peak value, then decreases slightly (i.e. strain weakening) before rising markedly (i.e. strain hardening). Small temperature differences between shift plateaus of curves up or down. (b) Experiments conducted at 550-555°C at 2.5×10⁻³ s⁻¹ to variable final strain (color bar). Samples show pronounced strain weakening followed by strain hardening. At high strains $(0.45 < \varepsilon < 0.6)$ sharp repetitive stress drops occur (curves from samples compressed to $\varepsilon = 0.6$ are offset within figure for clarity). (c) Experiments conducted at 2.5×10⁻³ s⁻¹ to 0.60 strain at temperatures from 540°C (light blue) to 581°C (red) (color bar). Curves that show stress drops are grouped by temperature range and offset for clarity. Below 550°C samples fracture during loading. Below 560°C samples deform viscously, followed by stress drops. As temperature increases stress drops occur at higher strains. No stress drops occur in samples deformed above 560°C. Initial peak σ values and the magnitude of strain weakening (0.04<ε<0.30) decrease as temperature increases. At temperatures above 555°C strain hardening (ε >0.30) causes the final σ values to exceed the initial peak σ .

Figure 3. Acoustic emissions (AE) during deformation. Solid curves are stress (σ; MPa) against time (s) and strain (ε; fractional) data for samples deformed at 2.5×10⁻³ s⁻¹ to 0.60 strain. Symbols show the amplitude of AE events against time (i.e. strain) for events recorded after sample loading. AE events >75 dB (high-AE; red circles) correlate with stress drops in mechanical data. Grey panels demarcate periods of loading, strain weakening, strain hardening, and the onset of stress drops. (a) Experiment at 585°C. No stress drops occur. Few AE events <75 dB (low-AE; white circles) occur. (b) Experiment at 555°C. The number and amplitude of low-AE events reach a minimum at ε~0.10 (middle of strain weakening), and a stable maximum at ε~0.35 (middle of strain hardening). (c) Experiment at 548°C. Sample fractures during loading. Pattern of low-AE events the same as in (b). Following large stress drops, and high-AE events, the number of low-AE events sharply decreases, then sharply increases back to the plateau level.

0.6

90 Amplitude (dB) 50

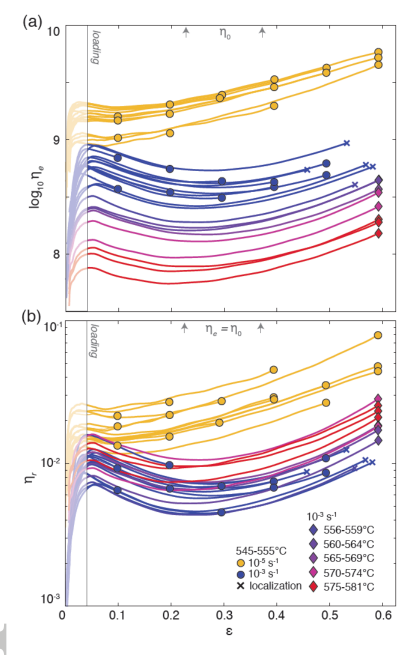


Figure 4. Transient effective (η_e) and relative (η_r) viscosities. Curves show (a) η_e and (b) η_r against strain (ε; fractional) for all experiments (legend). Symbols show final ε- η_e values (Table 2). For samples that showed stress drops and strain localization crosses mark the maximum ε- η_e values prior to the first stress drop. At fixed ε, η_e is greatest for samples deformed slowly, and lowest for samples deformed quickly and at high *T*. Curve shapes reflect the presence and magnitude of strain weakening and strain hardening (Figure 2). After accounting for the effect of temperature on deformational response, samples deformed slowly show the greatest values of η_r at fixed ε.



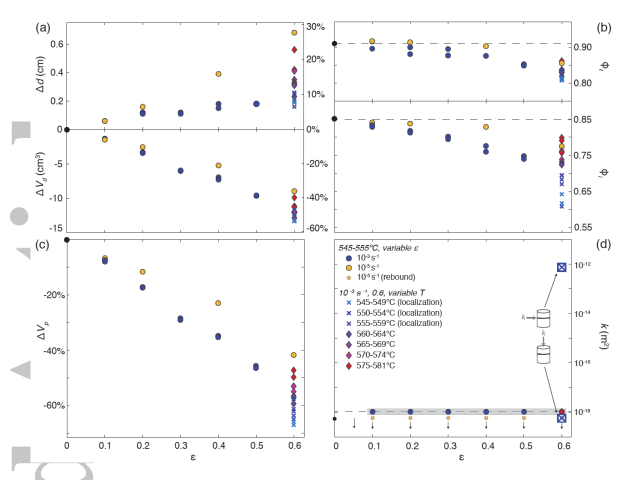


Figure 5. Physical properties. (**a; upper**) Change in diameter (Δ*d*; cm and %) and (**lower**) dimensional volume (Δ*V_d*; cm³ and %), against strain (ε; fractional) (Table 1). (**b; top**) Measured final total (ϕ_t) and (**bottom**) isolated porosities (ϕ_t) against ε (Table 1). Dashed lines show initial total and isolated porosities. (**c**) The normalized change in total pore volume (Δ*V_p*; %) against ε. (**d**) Measured permeability (k; m²) against strain. For the initial material (black circle) and rebound samples (small yellow circles), the estimated k is below the apparatus detection limit (dashed line at 10^{-18} m²). Most cores are estimated or inferred (grey box) to have permeabilities << 10^{-18} m². Large ×'s show permeability of samples showing strain localization, measured perpendicular (k << 10^{-18} m²) and parallel to the plane of localization (k = 7.3×10^{-13} m²) (cartoons). Legend in (d).

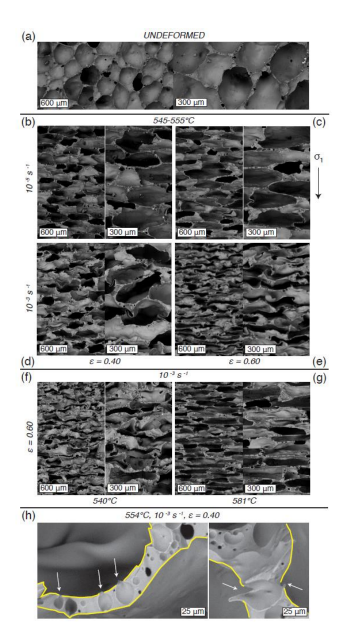


Figure 6. Bubble and bubble wall shapes. SEM images of each sample at two magnifications (scale bars in microns). Deformation conditions listed above image pairs. σ_1 shows the compression direction. (a) Undeformed FOAMGLAS®. Large bubbles are polygonal to circular and near-equant. Bubble walls are straight to curved. (b) Large bubbles are slightly elongated perpendicular to σ_1 , and shortened parallel to σ_1 . Bubble walls are stretched and curvilinear. (c) At higher strain bubbles are more elongate perpendicular to σ_1 , and shorter parallel to σ_1 . Large bubbles are rectangular or elliptical. Most bubble walls are straight. Few are buckled. (d) At higher strain rates, large bubbles have complex shapes resulting from folding and buckling of bubble walls, both parallel and perpendicular to σ_1 . (e) With increasing strain, bubble shapes become increasingly complex and generally elongate perpendicular to σ_1 . Bubble walls oriented perpendicular to σ_1 are straight. Walls oriented parallel to σ_1 are intensely buckled and folded. (f) At the same strain rate and strain, and lower temperatures, bubble and bubble wall geometries are complex. (g) At higher temperatures bubble shapes are more regular and rectangular. Bubble walls are folded where oriented parallel to σ_1 , but straight when oriented perpendicular to σ_1 . (h) White arrows show torn melt films between small bubbles within the bubble wall (outlined in yellow) and adjacent large bubble(s). Sample is 328, shown in (d).



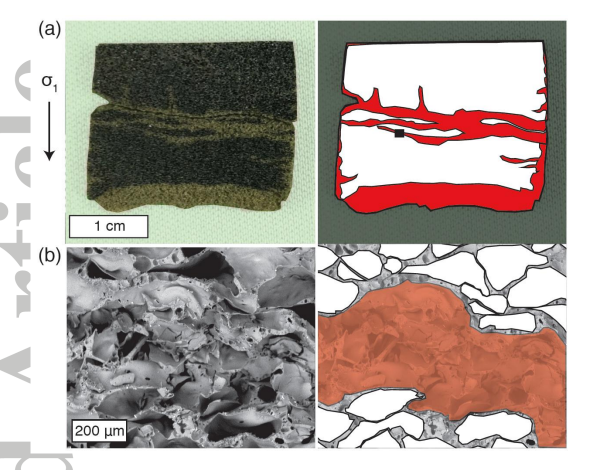


Figure 7. Compaction bands. (a) Photograph (left) of sample 311 (Table 1), showing green (oxidized) fractured regions, which form perpendicular to σ₁ and cross-cut the sample. "Spurs" oriented parallel to σ₁ are only observed in this sample. Schematic (right) shows the band in red and the impermeable remainder of the core in white. Black box shows the position of (b). (b) SEM image (left) of a compaction band. Bubbles throughout sample are highly deformed and have complex shapes (Figure 6f). The compaction band is evidenced by disrupted, fractured bubble walls. Schematic (right) shows bubbles that appear intact as white shapes, and the area occupied by disrupted bubbles (compaction band) as a red zone. The width of the band is ~500 μm.

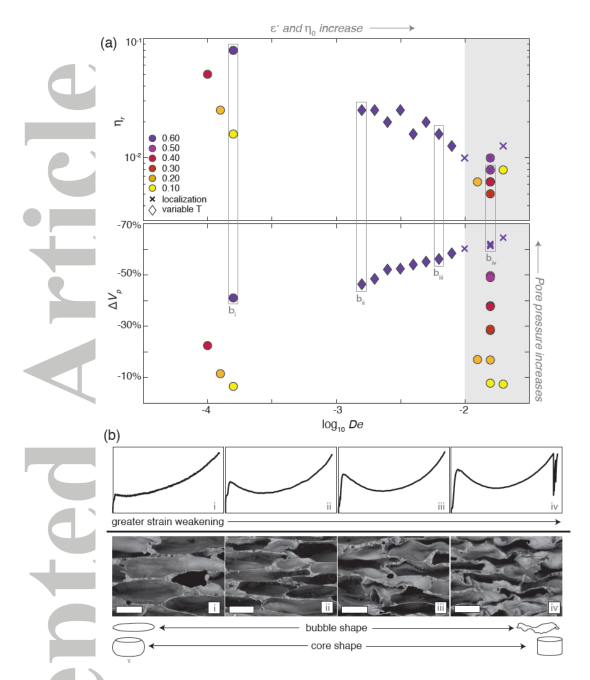


Figure 8. Microstructure, rheology and outgassing as a function of Deborah number (De). At high De, the melt is unrelaxed ($De > 10^{-2}$; grey vertical zone). At low De the melt easily dissipates stress (relaxes). (**a; top**) Modeled relative viscosities ($η_r$) against De. Colors show strain, and crosses show samples with compaction bands (legend). At the same strain $η_r$ decreases with increasing De. (**a; bottom**) Modeled changes in cumulative pore volume ($ΔV_p$; %) against De. The reduction in pore volume increases with increasing strain, and as melt viscosity and/or strain rate increase (higher De). (**b; top**) Representative mechanical data, corresponding to boxes in (a). As melt viscosities and strain rates increase (increasing De), samples show greater strain weakening. (**b; bottom**) Cartoons and SEM images showing the change in core and bubble shape, and microstructure with increasing De (boxes in (a)). Scale bars on SEM images are 200 μm. With increasing De bubble walls buckle, bubble shapes become contorted, and cores undergo less bulging.

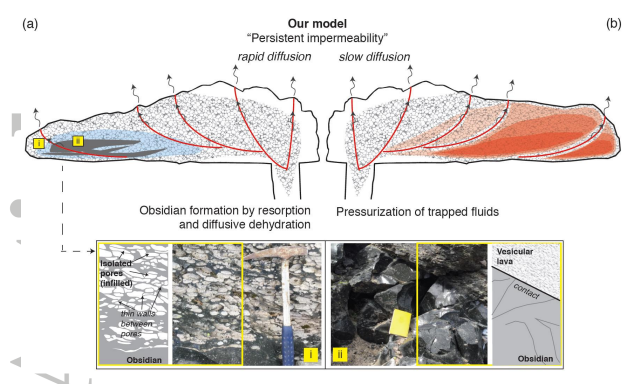


Figure 9. Consequences of "persistent impermeability" of silicic lavas. Schematics show a lava flow populated with isolated bubbles formed during syn-eruptive vesiculation. The lava is initially impermeable. In regions where strain rates and melt viscosity are high, the foam is unrelaxed and strain localization causes permeable pathways, such as compaction bands, to form (red lines). These pathways support localized outgassing (arrows). However permeable pathways do not form in the remainder of the flow, and it remains impermeable. (a) Where diffusive processes (i.e. diffusive dehydration, volatiles resorption) are rapid, they may allow for outgassing and densification to occur. Observed dense, non-vesicular lavas (obsidian; grey regions) may be formed by diffusive processes operating in the flow interior (blue region) rather than by outgassing through numerous permeable pathways (Figure 1b). Boxes mark locations in hypothetical flow where textures observed in nature (at flow front of Little Glass Mountain, Medicine Lake Volcano, California, USA) could be expected. These textures include: (i) large, flattened, closely packed bubbles that appear to be isolated from one another (hammer for scale; see schematic at left). Bubble interiors are filled with precipitated/alteration materials (beige color infilling bubbles) that have uniform thickness along the interior surfaces, perhaps indicating they accumulated on the inner surfaces while the bubbles remained isolated from one another and the atmosphere; (ii) thick band of dense obsidian containing few bubbles, bounded on its upper surface by vesicular lava containing many deformed vesicles (field book for scale; see schematic at right). The apparent connectivity between bubbles is not zero (as in (i)), but is low. Obsidian formation may result from diffusive dehydration, or by localized increases in bubble connectivity leading to formation of ephemeral permeable pathways. (b) Where diffusive processes are slow, the core of the lava still contains compressible fluid-filled bubbles. As the lava flows and bubbles are compressed, pore pressures rise (red regions). Sudden unloading of this region can trigger far-field explosive activity.