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Experimental constraints on the textures and origin of obsidian pyroclasts

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1 Abstract

2 Obsidian pyroclasts are commonly preserved in the fall deposits of explosive silicic eruptions. 3 Recent work has suggested that they form by sintering of ash particles on the conduit walls above 4 the fragmentation depth, and are subsequently torn out and transported in the gas-particle 5 dispersion. Although the sintering hypothesis is consistent with the general vesicle textures and 6 dissolved volatiles in obsidian pyroclasts, previous sintering experiments do not capture all of the 7 textural complexities observed in the natural pyroclasts. Here, we design experiments in which unimodal and bimodal distributions of rhyolitic ash are sintered at temperatures and H₂O 8 9 pressures relevant to shallow volcanic conduits, and under variable cooling rates. The 10 experiments produce dense, welded obsidian that have a range of textures similar to those 11 observed in natural pyroclasts. We find that using a unimodal distribution of particles produces 12 obsidian with evenly distributed trapped vesicles, while a bimodal initial particle distribution 13 produces obsidian with domains of poorly vesicular glass among domains of more vesicle-rich 14 glass. We also find that slow cooling leads to resorption of trapped vesicles, producing fully dense 15 obsidian. These broad features match those found in obsidian pyroclasts from the North Mono 16 (California, USA) rhyolite eruption, providing strong support to the hypothesis that obsidian can 17 be produced by ash sintering above the fragmentation depth during explosive eruptions.

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19 Keywords: obsidian; ash; sinter; volatile; explosive eruption

21 Introduction

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23 Obsidian pyroclasts – pieces of glassy rhyolite found in tephra deposits – have played a central 24 role in the development of models for silicic magma degassing and volcanic eruptions. Found in 25 most rhyolitic tephras that are associated with domes or lavas (e.g. Rust et al., 2004; Castro et al. 26 2014; Watkins et al. 2017, Gardner et al. 2017), these pyroclasts record a wide range of dissolved 27 volatile contents, all of which are partially degassed relative to initial storage conditions (Taylor 28 et al., 1983; Newman et al., 1988; Taylor, 1991; Watkins et al., 2017). This observation, combined 29 with their low-to-zero vesicularity, was first explained by what became known as the "permeable 30 foam model", in which bubbly rhyolitic magma becomes permeable through partial bubble 31 coalescence, allowing gases to escape and the remnant magma to collapse to a bubble-poor melt 32 (Eichelberger et al., 1986; Jaupart and Allègre, 1991). This paradigm was linked to eruptive style, 33 such that the collapse of the foam was postulated to cause the magma to effuse as degassed lava 34 instead of erupting explosively (Eichelberger et al., 1986). More recent models have been 35 proposed to explain the high dissolved volatile contents that appear inconsistent with the 36 permeable foam model (e.g., Rust et al., 2004; Gonnermann and Manga, 2005a,b; Okumura et 37 al., 2009; Cabrera et al., 2011; Castro et al., 2014). Textural evidence for brittle behavior in 38 otherwise hot, viscous silicic magma has been used to implicate fracturing as the cause of open-39 system behavior (Tuffen et al., 2003; Tuffen and Dingwell, 2005; Cabrera et al., 2011), which in 40 turn appears consistent with the style and extent of degassing recorded by dissolved volatiles 41 (Rust et al., 2004; Castro et al., 2014). These models converge on the idea that brittleness 42 resulting from high local strain rates in the melt lead to fractures that open the system, and that

this fracturing occurs repetitively (Tuffen et al., 2003) and pervasively (Castro et al., 2012) during
magma ascent.

45 Gardner et al. (2017) re-examined obsidian pyroclasts from the North Mono eruption and found multiple lines of evidence that the obsidian is not collapsed foam. Instead, they proposed 46 47 a model in which volcanic ash travelling up the conduit, above the fragmentation level, sticks to 48 conduit walls and sinters to form low-porosity melt. Blebs of this melt are then reincorporated 49 into the eruption to be deposited as glassy pyroclasts in the mainly pumiceous tephra. In this 50 framework, variations in dissolved volatile concentrations represent differences in the depth at 51 which ash sinters, and in the residence time at any given depth before final expulsion (Watkins 52 et al., 2017; Gardner et al., 2017). An origin by in-conduit sintering explains the low porosity, the 53 extent of open-system degassing of volatiles (Watkins et al., 2017), and trapped lithic fragments 54 within obsidian pyroclasts (Rust et al., 2004; Gardner et al., 2017).

To explore the timescales involved in ash sintering, Gardner et al. (2018) experimentally subjected rhyolitic ash to temperatures and H₂O pressures approximating those in the conduit. They found that ash-sized particles can equilibrate their dissolved volatile contents and sinter to vesicle-poor melt in as little as a few minutes, demonstrating that dense melt can form on the timescale of an eruption and thus obsidian pyroclasts need not be pieces of quenched magma emplaced before the eruption (cf. Newman et al., 1988; Dunbar and Kyle, 1992; Rust and Cashman, 2007).

The experimental results of Gardner et al. (2018) validated the timescales for sintering ash to obsidian, but they did not address the vesicle textures of obsidian pyroclasts. In fact, many textures of the North Mono obsidian pyroclasts do not match those predicted by isothermal

65 sintering. For example, sintered samples reach an equilibrium porosity of ~3–5 vol.% 66 (Wadsworth et al., 2016; Gardner et al. 2018), which is thought to be the point at which 67 permeability approaches zero, and porosity becomes isolated (Wadsworth et al., 2017). 68 Although the value of 3 vol.% is in good agreement with predictions from percolation theory and 69 simulations (Kertész 1981; Elam et al. 1984; Vasseur and Wadsworth, 2017), the majority of 70 obsidian pyroclasts from the North Mono eruption have ≤1 vol.% vesicles, and some have no 71 vesicles. Other textural variations in the natural samples include vesicles that range in shape 72 from spherical to highly distorted, and in number densities (N_v) between 0 (i.e., no vesicles) and 73 ~10⁸ vesicles per cm³ (Gardner et al., 2017). Volatile contents are also heterogeneous, regardless 74 of vesicularity, and even very vesicle-poor obsidians have sub-domains with contrasting H_2O and CO₂ contents (Watkins et al., 2017). 75

76 Many textures seen in the North Mono obsidian pyroclasts were thus not captured by the 77 experiments presented in Gardner et al. (2018) or indeed in any other experimental work 78 designed to probe the origin of sintered rhyolitic deposits (Wadsworth et al., 2014). To address 79 this gap we carried out sintering experiments that complement those of Gardner et al. (2018), 80 but aim to replicate more accurately the natural scenario by using different particles sizes, mixed 81 particle-size distributions, and by imposing cooling on the sintering particle pack. Combined, 82 these results provide an improved framework in which to interpret the vesicle textures of 83 obsidian pyroclasts.

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85 Methods

87 All experiments used the same rhyolitic obsidian used by Gardner et al. (2018), the composition 88 of which is listed in Table 1. This obsidian has a starting H₂O concentration of 0.15 wt.% 89 (Gardner et al. 2018). The obsidian was crushed into pieces with a steel mortar and pestle, and 90 then those pieces were ground to a powder using an agate mortar and pestle. The powders 91 were sieved at different times during the grinding process to produce splits of different particle 92 sizes. The first split consisted of particles that remained on the no. 35 (500 µm) and no. 60 (250 93 μ m) mesh sieves. Because of their irregular shapes, the ash particles in this split have 94 dimensions that are smaller and larger than 250–500 µm. We measured 20 random particles 95 under the microscope and found that they had principal axes of $696\pm244 \mu m$, $429\pm73 \mu m$, and 212±81 µm; we refer to this sample as "coarse ash". The second split consisted of particles that 96 97 passed through the no. 325 (45 µm) mesh sieve. Under the microscope we found that these 98 are ≤ 2 to 45 µm in size; we refer to this sample as "fine ash".

Separately, a cylindrical core was drilled from the obsidian, and hydrated at 875° C and 40 MPa, following the methods of Gardner and Ketcham (2011). The resulting sample was clear rhyolitic glass with ~2.3 wt.% dissolved H₂O, as measured by FTIR (see below). The glass was crushed, ground, and sieved, and we collected particles that remained on the no. 80 (180 μ m) mesh sieve. Measurements of 20 random particles revealed that they are blockier than the coarse ash, with principal axes of 298±45 μ m, 209±33 μ m, and 159±25 μ m; we refer to this sample as "wet ash".

Each experiment used ~30–40 mg of powder (Table 1). In one set of experiments we used only one of the unimodal samples described above (fine, coarse, or wet). These experiments serve as a calibration of the sintering timescales for each population, before mixing

populations. In a second set of experiments, we used bimodal mixtures consisting of the fine
ash mixed with either ~25 wt.% or ~50 wt.% coarse ash or 50 wt.% wet ash.

111 Each sample was weighed into a gold capsule (3 mm O.D.) that was welded shut on one 112 end. The other end of the capsule was left open. Each capsule was placed into a sample holder 113 at the end of an Inconel rod, which was then inserted into a pressure vessel fitted with a rapid 114 quench extension, as described in Gardner (2007). The pressure vessel was connected to the 115 pressure line, and 22 MPa of H_2O pressure was applied. Because the capsule was open at one 116 end, the pressurized H₂O was in contact with the powder and thus the interstitial inter-117 connected pore space was at the same pressure as that in the pressure vessel. The capsule 118 thus exerted no stress on the particles within, and the only stress driving sintering arose from 119 interfacial tension between the molten particles and interstitial H_2O (Gardner et al., 2018). 120 An external magnet held the sample rod in place, which held the sample inside a water-121 cooled jacket while the pressure vessel heated to 750°C, as measured by K-type thermocouples 122 precise to ±5°C. When the vessel thermally equilibrated, pressure was set at ~2.0 MPa above 123 22 MPa, as measured by a pressure transducer precise to ±0.1 MPa. The external magnet was 124 then raised, lifting the sample into the hot zone of the pressure vessel in ~1 second; there was 125 an associated pressure drop of ~2.0 MPa. Pressure was then quickly adjusted to 22 MPa in ~15 126 seconds. The sample was held in place for 5 to 60 minutes (Table 1). During that time, 127 pressure varied by no more than 0.1 MPa and temperature varied by no more than 1°C. After 128 the target time was reached, the magnet was lowered, bringing the sample into the water-129 cooled jacket where it cooled at ~100°C per second (Gardner et al., 2018). When the sample

was lowered, pressure increased by ~2.0 MPa, but was quickly adjusted back to 22 MPa in ~15
seconds.

132 In a final set of experiments, samples were sintered isothermally for a given time and then 133 cooled slowly to a specified temperature before being quenched rapidly, to investigate the 134 effects of cooling on final textures (Table 2). In this set of experiments, most samples consisted 135 of fine ash - one held a 50:50 mix of fine and coarse ash - that were first held at 22 MPa and 136 750°C for 5, 10, or 30 minutes, mimicking the isothermal experiments described above; the 137 furnace was then switched off to allow the samples to cool. The rate of cooling changed with 138 time, because it was controlled by the heat loss from the insulated furnace: it took 49.6±5.7 139 seconds to cool from 750°C to 748°C, and then another 36.2±3.9 seconds to cool to 745°C. 140 After that, samples cooled at an approximately constant rate of 9.4±0.3°C per minute, and the 141 samples reached 700°C by 6.3±0.3 minutes, 650°C by 11.4±0.6 minutes, and 550°C by 23.5±1.2 142 minutes. Although cooling slowed below 650°C, the change was not significant. Throughout 143 cooling, pressure was maintained within 0.1 MPa of 22 MPa. All samples were quenched 144 rapidly as soon as they reached the final target temperature. 145 Experiments were run over a range of temperatures from 550° to 750°C at 22 MPa. 146 Assuming H_2O solubility (see below), those conditions equate to dissolved H_2O contents of 1.8 to 2.2 wt.% (Liu et al., 2005). We estimate that melt viscosity was thus $\sim 10^{7.1}$ to $10^{10.1}$ Pa s 147 148 (Hess and Dingwell, 1996), which is well below a viscosity of 10^{11.4} Pa s that is typically 149 considered for the glass transition (Gottsmann et al., 2002). Therefore, under all conditions of 150 our experiments, the samples were fluid and could behave as melts.

151 All samples were extracted from their gold capsules and, if coherent, sealed in epoxy and 152 thin sectioned to about 500 μ m thickness. A petrographic microscope was used to inspect 153 sample textures and measure vesicle numbers, shapes, and sizes. Vesicle sizes were measured 154 using a graduated ocular on the petrographic microscope, and are precise to $\pm 0.5 \ \mu m$. Number 155 density of vesicles (N_V) was measured by selecting different areas in a sample and counting all 156 vesicles that appear as the field of view is moved through it using the focusing knob of the 157 microscope; N_V is thus number density per unit total volume (melt plus vesicles). The depth 158 viewed was measured using a Heidenhain focus drive linear encoder. All vesicles, regardless of 159 shape, were counted, but only spherical vesicles were measured for size. Porosity (ϕ) was 160 measured by photographing samples in reflected light, and then using NIH Image J to make 161 binary images of the photographs to measure pore areas relative to the entire area of the 162 image, and converted directly to porosity (vol.%). Errors on ϕ are estimated at 10% of the 163 measured value.

164 Glasses in the hydrated core, G-1724, G-1728, and G-1762 (Table 1) were analysed for 165 dissolved H₂O contents by Fourier Transform Infrared (FTIR) spectroscopy in transmittance mode, 166 using a Thermo Electron Nicolet 6700 spectrometer and Continuµm IR microscope. Spectra were 167 collected using 60 scans at a resolution of 4 cm⁻¹, and measured either in near-IR (7000 to 3800 168 cm⁻¹) with white light and a CaF₂ beamsplitter or in mid-IR (4000 to 650 cm⁻¹) with a globar source 169 and KBr beamsplitter. Dissolved contents of molecular water (H_2O_m) and hydroxl water (OH) 170 were determined from absorbances at ~5250 cm⁻¹ and ~4500 cm⁻¹, respectively, using the model 171 of Zhang et al. (1997), or from the broad absorbance at ~3500 cm⁻¹, using an absorptivity of 75 L mol⁻¹ cm⁻¹ (Okumura et al., 2003). Sample thickness where each spectrum was collected was
determined using the method described above.

174 Area maps of OH, H₂O_m, and total H₂O concentrations were made for G–1783 (Table 2) using 175 a Thermo Nicolet Nexus 670 Fourier transform infrared (FTIR) spectrometer, following the 176 methods of Watkins et al. (2017). All measurements were made using a 15X objective, infrared 177 source, MCT-A detector, and KBr beamsplitter. Absorbances were converted to concentrations 178 using the model of Zhang et al. (1997). The thickness of the sample was measured in several 179 spots using a digital caliper with 0.001 mm precision. All spectra were collected at a resolution 180 of 4 cm⁻¹, 64 scans per spot, and 64 scans for the background, which was collected every 10 181 minutes. Two maps were made, one using a 100 μ m \times 100 μ m aperture and step size of 100 μ m 182 and the other a 40 μ m \times 40 μ m aperture and step size of 40 μ m. No significant differences were 183 detected between the two maps.

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185 Experimental results

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187 Experiments Using Unimodal Powders

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189 Post-experimental samples of sintered fine ash were all sufficiently coherent to section and

- analyze (Table 1). The highest measured porosity was ~46 vol.% for a sample that was
- 191 incipiently sintered, which suggests that initial, unsintered porosity was slightly higher. This is
- 192 consistent with the initial porosities measured by Gardner et al. (2018), who found that ash
- 193 poured into capsules has an initial porosity between the particles of 45–55 vol.%, with less

polydisperse ash having higher porosity. Over run durations from 5 to 60 minutes, porosity
decreased to the apparent equilibrium of ~3–5 vol.%, which was reached after 20 minutes
(Figure 1). After 20 minutes, the sample of sintered fine ash (G–1728) has 1.77±0.20 wt.% H₂O
dissolved in the glass, which agrees well with the H₂O content (1.81 wt.%) predicted by the
model of Liu et al. (2005). Gardner et al. (2018) also found that dry ash fully saturated within
20 minutes at 750°C and 40 MPa.

200 After 5 and 10 minutes, samples of sintered fine ash consist of completely open and 201 connected pore space around individual ash particles that are still visible, although their edges 202 have sintered to their closest neighbors, forming arcuate necks (Figure 2a). After 20 minutes, 203 most pores are isolated from neighbors, but about two thirds of these vesicles are still 204 distorted; the other third are spherical (Figure 2b,c). Between 30 and 60 minutes the 205 proportion of distorted vesicles decreases significantly (Figure 2d), and by 60 minutes 206 essentially all are spherical (Figure 2e). By 20 minutes, spherical vesicles are $<2-5 \mu m$ in 207 diameter but, later, larger ones become abundant. The larger vesicles did not grow from 208 smaller ones, but instead result from large contorted vesicles taking longer to relax in shape. 209 Values for N_V are ~10^{7.5-7.8} cm⁻³ (Table 1).

Gardner et al. (2018) proposed a classification for the textural evolution of sintering unimodal rhyolitic ash, which identifies four phases: 1) particles are loose and lack cohesion; 2) particles are sintered at their contacts but are interpenetrated in a continuous, tortuous pore space; 3) porosity is no longer fully connected, with small isolated spherical vesicles and networks of larger vesicles that remain multi-cuspate in shape; and 4) dense glass with fully isolated, spherical vesicles and maybe a few larger complex–shaped ones; duration of each

phase depends strongly on melt viscosity. We use this framework to present the results of
these experiments with unimodal samples, finding that the textures of the sintered fine ash
follow the same progression (Figure 3). Samples sintered for 5 and 10 minutes consist of
individual particles partially sintered in open pore space (Phase 2). Between 20 and 45 minutes,
samples have low porosities, but distorted vesicles make up 40 to 70% of the population (Phase
3). After sintering for 60 minutes the sample consists of dense glass with only spherical vesicles
(Phase 4).

223 The two experiments using only coarse ash were run for 30 and 60 minutes, but remained 224 unsintered and crumbled to loose powder on extraction from the capsules, implying that the 225 inter-particle porosity did not decrease much below the initial value and the samples remained 226 in Phase 1. The two experiments using only wet ash were coherent enough to section, and 227 both consist of individual ash particles that are only slightly sintered along their edges (Figure 228 2f). The particles are dispersed amongst an open and connected pore space that makes up 30-229 35 vol.% of the samples (Figure 1). The wet ash samples thus remained in Phase 2 of textural 230 evolution even after 60 minutes.

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232 Experiments Using Bimodal Mixed Ash Populations

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In all mixed-ash samples, even when sintering has progressed significantly – which normally
 removes evidence of individual particles – the coarse or wet ash component can still be
 recognized as large non-vesicular domains, and in the case of wet ash as microlite-free

domains. Overall, bimodal samples sinter more slowly, as measured by the evolution ofporosity, than samples that consisted of only fine ash (Figure 1).

239 In describing the final textures of the mixed-ash samples, we use *matrix* to refer to domains 240 that initially consisted of fine ash between areas that were initially either coarse or wet 241 particles. When coarse ash makes up half of the sample, the matrix after 20 minutes (G-1724)242 consists of small ash particles that are slightly sintered surrounded by ~33 vol.% of pore space 243 (Figure 4a). Pockets of large vesicles occur in the matrix that can be \geq 800 microns in size, with 244 most pockets found in the vicinity of relict large particles. FTIR measurements show that the 245 large relicts have 0.57−1.79 wt.% H₂O dissolved in them (measured in the interiors of large 246 particles, as far away from the edges as possible), which is below the solubility value of 1.81 247 wt.% (Liu et al., 2005). After 45 minutes, matrix porosity is still heterogeneous, and depends on 248 proximity to relict coarse particles (Figure 4b,c). Where coarse particles are closer together than 249 ~200 µm, the matrix is of the order of 26 vol.% porous and can preserve pockets of large 250 vesicles up to 600 µm long; where farther apart, the matrix is only about 4 vol.% porous. In 251 both regions, $N_V \sim 10^{7.2}$ cm⁻³ and the vast majority of vesicles are distorted and connected 252 together via narrow channels. After 60 minutes, the matrix is almost uniformly dense, and 253 large pockets of vesicles are rarer and $\leq 200 \,\mu\text{m}$ in size (Figure 4d). All vesicles are isolated, but 254 only about half are spherical. The dissolved H₂O contents in the interiors of the relict coarse 255 particles is 1.84 (\pm 0.14) wt.%, which is within error of the 1.81 wt.% saturation limit of H₂O (Liu 256 et al., 2005).

A similar progression of textures is observed when coarse ash makes up only ~25 wt.% of the sample, except that matrix porosity is lower after any given amount of time (Figure 1).

259 Relatively large porous pockets still exist, but they are generally smaller and few remain after 260 45 minutes. One experiment that contained fine ash and 50 wt.% wet ash was run for 20 261 minutes. Its matrix is similar in porosity to that of the coarse ash mix that sintered for more 262 than twice as long (Figure 1). It contains large pockets of vesicles up to 300 μm long and mainly 263 highly contorted, multi-cuspate vesicles; only about 16% of the vesicles are spherical. 264 Overall, the temporal evolution of textures in the mixed-ash populations is broadly similar 265 to the unimodal case for fine ash, such that porosity generally decreases with time and the end-266 state is isolated vesicles (Figure 1). The presence of large particles, however, slows the rate of 267 decrease in porosity, and causes individual vesicles to take longer to relax to spherical. After 60 268 minutes, for example, 13–20% of the matrix vesicles are still distorted in samples with 50% 269 coarse particles, whereas all were spherical in samples of fine ash only. Another important 270 difference is that bimodal samples produce textures that are considerably more 271 heterogeneous, with pockets of large vesicles. These pockets generally decrease in size with 272 time, and are less abundant when coarse particles are less abundant. 273 274 **Experiments Combining Sintering and Cooling** 275 276 Six samples of fine ash were cooled slowly after sintering at 750°C (Table 2). All have 277 significantly reduced porosities compared to samples that were not slowly cooled (Figure 5). 278 For example, sintering isothermally for 5 minutes produced only incipiently sintered particles 279 amongst ~45 vol.% connected pore space, whereas subsequent cooling to 550°C produced 280 dense glass with 7 vol.% isolated vesicles. Importantly, cooling is found to be able to produce

281	dense glass with no measurable vesicles (Figure 5), and reduced N_{ν} from 10 ^{7.66} cm ⁻³ to zero
282	vesicles (Table 2). FTIR Mapping across G–1783 (no vesicles present) found H_2O concentrations
283	between 1.88 and 2.16 wt.%, averaging 2.03±0.06 wt.%. That range falls between solubility
284	H_2O concentrations expected at 750° to 550°C and 22 MPa (Liu et al., 2005).
285	Although porosity is reduced during cooling, the preserved vesicles are relatively more
286	deformed than those found after similar durations at high temperature. For example, after 30
287	minutes of sintering at 750°C (G–1729) about half of the vesicles have relaxed to spherical
288	shape (Figure 6a). In contrast, G–1789, which sintered for 10 minutes at 750°C and then cooled
289	to 550°C for ~25 minutes (total time = 35 minutes), has the same porosity, but all of the vesicles
290	are distorted in shape (Figure 6b). Cooling can thus prolong the time it takes for distorted
291	vesicles to relax to spherical, while simultaneously reducing overall porosity.
292	One sample that consisted of equal proportions of fine and coarse ash was cooled to 650°C
293	after sintering for 30 minutes (G–1785; Table 2). Given the total amount of time spent at
294	elevated temperature (41 minutes), this sample would be expected to have a porosity of \sim 3
295	vol.% far from large particles and ~26 vol.% near relict particles (compared to G–1760; Table 1).
296	Instead, the corresponding values for the cooled sample are ~2 and ~16 vol.%. Furthermore,
297	cooling reduced N_V by an order of magnitude compared with the isothermal sample (Tables 1
298	and 2).
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300 Analysis of Experimental Results

302	The textural descriptions above highlight some of the complexities of the sintering process for
303	rhyolitic particles that have not been accounted for in existing sintering models (Wadsworth et
304	al., 2014; 2016). Here, we place these experiments in the context of existing theoretical
305	constraints to highlight these differences.
306	Gardner et al., (2018) defined two timescales relevant to sintering of unimodal rhyolite
307	particles that were also hydrating. The first timescale is for diffusive hydration (λ_d), during
308	which the dissolved H2O content increases from the initial value of 0.15 wt.% to the saturation
309	value, which depends on the sintering temperature and pressure. The second timescale is for
310	sintering (λ_s). For randomly packed, monodisperse spherical particles (droplets when molten),
311	the sintering timescale is given by
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313	$\lambda_s = \frac{\eta L}{\sigma} \tag{1}$
314	
315	where η is droplet viscosity (Pa s), σ is surface tension (N m ⁻¹), and L (m) is a characteristic
316	length scale, which can be approximated as the mean particle radius ($ar{R}$) in the case of sintering
317	angular particles (Gardner et al., 2018). The diffusive hydration timescale is given by
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319	$\lambda_d = \frac{\bar{R}^2}{D} \tag{2}$
320	
321	where D is the diffusivity of H ₂ O in rhyolite. Both η and D are material properties that must be

322 computed from constitutive models. We use the model of Hess and Dingwell (1996) to

323 compute η as a function of dissolved H₂O concentration (*C*) and temperature (*T*). In turn, we

use the model of Liu et al. (2005) to compute C at isothermal temperature (T_0) and isobaric

325 pressure (P_0). We use the model of Zhang and Ni (2010) to compute D, as a function of C, T_0 ,

and P_0 . We set $\sigma = 0.22$ N m⁻¹, following Gardner et al. (2018) for the same starting material.

327 Comparing timescales yields the dimensionless capillary Peclet number $(P_c) = \frac{\lambda_d}{\lambda_s}$

328 (Gardner et al., 2018). For $\lambda_d \ll \lambda_s$ (i.e., $Pc \ll 1$) diffusive hydration occurs rapidly compared 329 with sintering. Conversely, for $\lambda_d \gg \lambda_s$ (i.e., $Pc \gg 1$) sintering occurs rapidly compared with 330 diffusive hydration. Because both timescales depend on *C*, and *C* changes over time during the 331 experiments, we need to establish what value of *C* to use to compute the timescales. We 332 assume that the particles have equilibrium H₂O concentrations (*C_e*) for Pc \ll 1, and the

particles have their initial H₂O concentrations for $Pc \gg 1$. Gardner et al. (2018) found that

334 sintering can be assumed to be proceeding at equilibrium conditions as long as $Pc \lesssim 10$ (the

low Pc regime). All unimodal experiments of this study meet the condition Pc < 10, implying

336 that hydration was complete before sintering. This is confirmed by measurements of H₂O

concentration in G-1728 (measured for experimental conditions where time > λ_d), which is

338 within uncertainty of C_e . We thus assume C_e has been reached, and use that value to compute

the viscosity at C_e (which we term η_e) in (1) to predict λ_s .

340 The metric most commonly used to track the progression of sintering is the interstitial

porosity (ϕ) to the particles (Wadsworth et al., 2014; 2016). Wadsworth et al. (2016) found an

342 analytical approximation of a full sintering model that can be used to predict the porosity of a

343 random pack of particles as a function of time (*t*) following

$$\phi(t) = \phi_i \exp\left(-\frac{3t}{2\lambda_s}\right). \tag{3}$$

347	This expression has been validated for angular particles (Gardner et al., 2018; Wadsworth et al.,
348	2014). We find that the predicted $\phi(t)$ curve computed using (3), with λ_s computed from (1),
349	agrees well with the unimodal data using very different ash sizes as starting materials (Figure 7).
350	Note, initial porosity is taken as ϕ_i = 45 or 60 vol.%, and there are no adjustable parameters.
351	In the case of bimodal experiments, which contained two different $ar{R}$ values in different
352	proportions, the sintering process is complicated because the values of λ_d are different for each
353	constituent particle size class in the mixed population. This means that there can be conditions
354	for which $t < \lambda_d$ for large particles, while $t > \lambda_d$ for small particles. This is illustrated by the
355	cores of coarse particles in G–1724 having measured H_2O concentrations that is just over half of
356	C_e , whereas, after the same amount of time, unimodal fine ash in G–1728 is fully hydrated. The
357	result of this is that a system can be sintering particles with very different viscosities. Despite
358	such complexities, we found that the porosity of any bimodal mixture is intermediate, although
359	heterogeneous, between predicted values for $\phi(t)$ of the end-member unimodal $ar{R}$ classes
360	(Figure 7). In addition, the bimodal experiments that contain proportionally more fine ash
361	sintered faster, and hence approached the equilibrium porosity faster. While a full predictive
362	model for bimodal (and polymodal) sintering is beyond the scope of this study, we suggest that
363	$\phi(t)$ for bimodal sintering populations can be qualitatively estimated from those of the end-
364	member populations.

366 Resorption of Isolated Vesicles During Cooling

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368	Assuming that the cooled samples and isothermal samples initially had similar textures before
369	the start of the cooling ramp, our experiments demonstrate that slow cooling causes vesicles to
370	shrink and disappear (Figure 8). We hypothesize that this is caused by thermally-driven
371	resorption of H_2O ; i.e., the increase in H_2O solubility with decreasing T is sufficient to cause all
372	H_2O in the bubbles to resorb into the surrounding melt. McIntosh et al (2014) first proposed
373	that this mechanism could play a role in the formation of dense obsidian.
374	For bubbles to resorb completely, there must be sufficient time for all H_2O molecules within
375	the bubbles to diffuse into the surrounding melt. There must also be enough time for the melt
376	to flow viscously to allow the bubbles to shrink. To determine whether this is feasible in the
377	experiments, we compare three timescales: (1) the time available for diffusion and flow, λ_a ; (2)
378	the time that would be required for H2O to resorb diffusively, λ_{γ} ; and (3) the time that would
379	be required for the melt to flow, $\lambda_\eta.$ To test the resorption hypothesis, we determine these
380	three timescales for sample G–1783 (Table 2).

381 For this analysis, the time available for the bubble to resorb, λ_a is taken simply as the 382 duration of the cooling ramp. We derive order-of-magnitude estimates for the time required for diffusion, λ_{γ} , and viscous flow, λ_n , in Appendix A. We find that the time required for 383 384 diffusion to resorb all of the water in the bubbles depends strongly on the initial size of the 385 bubble, while the time required for viscous flow to allow the bubble to collapse depends only 386 weakly on the initial bubble size. In Figure 8, λ_{γ} and λ_{η} are plotted (solid and dash line respectively) as functions of initial bubble radius, given P_o = 22 MPa and T_o = 750°C. The time 387 388 available for diffusion and flow (λ_a) is also plotted for both the cooling ramp (red line), and for

rapid quench (blue line). We find that, over the range of initial bubble sizes in our experiments (indicated by the grey box), the time available for resorption during slow cooling is longer than the diffusion and flow timescales, hence there is sufficient time for resorption to go to completion. In contrast, the time available for resorption during rapid cooling is shorter than both the diffusion and flow timescales, hence there is insufficient time for resorption. This supports our hypothesis that thermally driven resorption is sufficient to cause the complete removal of the bubbles during slow cooling (Figure 5).

To first order, therefore, we find that slow cooling of porous rhyolitic melt can cause H₂O pores to resorb and produce porosities lower than the equilibrium sintering value of 3–5 vol.%. We note, however, that our approach (Appendix A) to determine order-of-magnitude values for λ_{γ} and λ_{η} is highly simplified. We thus caution against using this approach to make quantitative predictions about the conditions under which resorption may occur in natural systems. While such predictions are possible, it would require a numerical approach that is beyond the scope of this work.

403

404 **Discussion**

405

406 Our sintering experiments, combined with those of Gardner et al. (2018), place constraints on 407 the formation of vesicle textures generated during sintering. Comparing those results to 408 textures of natural obsidian pyroclasts provides a framework for understanding how the 409 pyroclasts form. All experiments show that, as ash sinters, the volume of open pore space 410 decreases and is sealed off, generating isolated vesicles (Figure 3). Those vesicles start off

411 highly distorted and convolute, but eventually relax to spherical shapes (Figure 2). That 412 sequence occurs in the different sets of experiments, but the timescales for sintering and 413 vesicle relaxation depend strongly on viscosity and initial particle size (Wadsworth et al., 2014; 414 Gardner et al., 2018; this study). The general progression from distorted to spherical shapes 415 observed in the experiments supports the overall premise of the model proposed by Gardner et 416 al. (2017) for the formation of obsidian pyroclasts in the North Mono eruption. They suggested 417 that the main mechanism for the formation of these pyroclasts is sintering of volcanic ash in the 418 conduit, speculating that vesicles start out distorted in shape but relax to spherical with time. 419 Furthermore, the experiments demonstrate that distorted and spherical vesicles may coexist 420 (i.e., Phases 2–3) during the sintering process, as observed in North Mono obsidian pyroclasts 421 (Figure 9a).

422 Our experiments show that sintering unimodal ash evolves texturally towards 423 homogeneously distributed spherical vesicles (Figure 2e). About 10–15% of the North Mono 424 obsidian pyroclasts studied by Gardner et al. (2017) contain only spherical vesicles (Figure 9b; 425 see also Figure 5a in Gardner et al., 2017). We infer that those obsidian pyroclasts preserve the 426 end products of sintering unimodal distributions of ash. Experimental sintering of unimodal ash 427 also shows that N_{ν} is related to the size distribution of particles sintered. Sintering of fine ash (\leq 45 µm; this study) produced N_{ν} = 10^{7.51±0.19} cm⁻³ (Table 1). Sintering of ash that ranged in size 428 by 1–1600 μ m (\overline{R} = 89 μ m) resulted in N_v = 10^{6.55\pm0.23} cm⁻³ (Gardner et al., 2018). Sintering of 429 coarser but less polydispersive ash of ~63–400 μ m in size (\overline{R} = 185 μ m) produced even fewer 430 vesicles, $N_{\nu} = 10^{5.62 \pm 0.19}$ cm⁻³ (Gardner et al., 2018). The differences in N_{ν} are significant, 431 432 representing one to two orders of magnitude variation in numbers of vesicles within a given

433	volume. Vesicles in North Mono obsidian pyroclasts occur in N_{ν} values from 0 (no vesicles) to
434	$10^{8.1}$ cm ⁻³ , but more than half have $N_{\nu} > 10^{6.8}$ cm ⁻³ (Gardner et al., 2017).

We propose that North Mono obsidian pyroclasts with abundant vesicles formed by sintering fine-grained ash. Fine-grained ash in fact makes up a large fraction of tephra produced in explosive eruptions (Walker, 1981; Kaminski and Jaupart, 1998; Bonnadonna and Houghton, 2005; Alfano et al., 2016). If the fine ash is approximately unimodal, we show that $\phi(t)$ can be predicted, and Gardner et al. (2018) provide a full framework for estimating it at conduit conditions.

441 The majority of North Mono obsidian pyroclasts have <1 vol.% porosity; furthermore, 3 of 442 81 samples reported by Gardner et al. (2017) were entirely vesicle-free. Experimental sintering 443 of unimodal ash, in contrast, typically produces an equilibrium texture of 3–5 vol.% vesicles 444 (Figure 1; Wadsworth et al., 2014; Gardner et al., 2018). The cause of this discrepancy has 445 remained enigmatic until now. Our experiments suggest that the relatively low porosities of 446 the natural obsidians result from resorption induced by cooling during sintering, and the 447 associated increase in H₂O solubility in the melt (Figure 8). We posit that many of the North 448 Mono obsidian pyroclasts thus preserve a record of cooling during sintering. In support of this, 449 our results show that non-spherical vesicle shapes can be preserved during cooling, and may 450 remain abundant well below the equilibrium porosity of 3–5 vol.% (Figure 6). Very poorly 451 vesicular North Mono obsidian pyroclasts, in fact, contain vesicles of all shapes, including highly 452 distorted ones (Figure 9c,d). Only a small minority of North Mono obsidian pyroclasts are free 453 of vesicles, which suggests that the time available for cooling was relatively short. Indeed, the

454 overall timescale for formation of obsidian pyroclasts as deduced from diffusion modelling of
455 volatile gradients is less than a few hours (Watkins et al., 2017).

456 It thus appears that some vesicle textures in natural obsidian pyroclasts can be explained by 457 sintering of unimodal ash, and that variations in textures among such pyroclasts are the result 458 of variations in starting particle sizes and in the cooling rate. There are other textural features 459 common to obsidian pyroclasts that are not replicated by experimental sintering of unimodal 460 ash, such as significant spatial heterogeneities in vesicles and volatile contents within a single 461 pyroclast (Rust et al., 2004; Rust and Cashman, 2007; Castro et al., 2014; Watkins et al., 2017; 462 Gardner et al., 2017). These include large pockets of vesicles dispersed amongst poorly 463 vesicular, dense glass (Figure 9d) and regions of poorly vesicular glass separated by glass with 464 numerous vesicles (Figure 9c,e,f). Many of these vesicular heterogeneities coincide with 465 heterogeneous dissolved volatile concentrations (Watkins et al., 2017). 466 We suggest that the spatial heterogeneities in vesicles and volatiles within obsidian 467 pyroclasts result from sintering of particles of differing size, based on our results of sintering of 468 mixed-ash populations. We found that large scale heterogeneities in vesicles can occur when 469 samples consist of mixed ash particles (Figure 4). For example, sintering of coarse and fine ash 470 produced regions poor in vesicles (large relict particles) separated by porous bands of matrix 471 (Figure 4e). Sintering of mixed ash samples also resulted in sub-domains of matrix with large 472 pockets of vesicles (Figure 4b,f), which are reminiscent of isolated clusters of large vesicles in 473 the natural obsidian pyroclasts (Figure 9e).

The degree of heterogeneity is likely related to the contrast in sizes between sintering particles. In this study, bimodal populations resulted in highly vesicular bands separating

476 relatively large, non-vesicular regions (Figure 4c,e). When the sintering pack includes a wide 477 range of particle sizes, then heterogeneities in vesicular textures range in size reflecting the 478 particle sizes (Gardner et al., 2018). Our results additionally suggest that heterogeneities in 479 volatile contents will result when the residence time at a given T and P is sufficient to 480 homogenize small particles ($t < \lambda_d$) but not sufficient to homogenize large ones ($t > \lambda_d$). 481 Commonly, the large, poorly vesicular domains in the North Mono obsidian pyroclasts are H₂O-482 rich relative to the vesicular domains (Watkins et al., 2017). We suggest that they are pieces of 483 glass formed by sintering and cooling of ash that was then ripped off of the conduit walls, only 484 to be re-plastered onto the walls and sinter with adjoining ash. 485 Finally, a prominent texture of North Mono obsidian pyroclasts not replicated in any sintering 486 experiment so far is stretched/elongated vesicles (see Figures 5c,d and 6d,f in Gardner et al., 487 2017). Such stretched vesicles usually align with their neighbors, implying that they formed by 488 shearing of distorted or spherical vesicles (Gardner et al., 2017). No shear occurred in our 489 experiments because the applied pressure is isotropic. Over 70% of the North Mono obsidians 490 preserve stretched/elongated vesicles, indicating that most, if not all, pyroclasts underwent 491 shearing during their formation. It is not possible to determine the cause of shearing based on 492 available evidence, but we speculate that it could be associated with gravitational slumping of 493 the sintering mass at the conduit wall or shearing along the margins induced by the ascent of the 494 adjacent erupting gas-particle dispersion, either before or during the disruption of the obsidian 495 to form pyroclasts.

496

497 **Conclusions**

498

499 Rhyolitic ash particles were experimentally sintered to examine the evolution of the vesicle 500 textures that result. Results show that, regardless of particle size, porosity decreases with time 501 and the interstitial pores quickly isolate to form vesicles amongst the sintering particles. With 502 time, isolated vesicles that are initially highly distorted relax to spherical shapes. The rates of 503 sintering and relaxation depend on melt viscosity and initial sizes of the particles. All else being 504 equal, smaller particles sinter more quickly than coarser ones. We also found that high vesicle 505 number densities ($N_{\nu} \ge 10^7$ cm⁻³) and generally small vesicle sizes result when the sintering 506 particles are unimodal fine-grained ash. Cooling of the sintering pack causes thermally-driven 507 resorption of volatiles, which reduces vesicle abundances below the equilibrium abundance of 508 3–5 vol.%, and can ultimately produce glass that is devoid of vesicles. While cooling reduces 509 the overall porosity, it also allows distorted vesicles to be preserved for longer periods of time, 510 and to lower porosities. Finally, bimodal mixtures of coarse and fine ash result in significantly 511 more heterogeneous vesicular textures than unimodal samples, including the creation of large 512 pockets of large vesicles between relict coarse particles. In addition, the overall reduction of 513 porosity and relaxation of vesicle shapes also takes longer than in unimodal samples. 514 These results provide a framework in which to interpret observed vesicle textures in natural 515 obsidian pyroclasts. In general, we conclude that obsidian pyroclasts form by sintering of 516 mostly fine-grained ash on conduit walls. The sintering ash also cools sufficiently slowly that 517 the majority of obsidian pyroclasts have vesicle abundances well below those produced in 518 isothermal sintering experiments. The various scales of heterogeneous vesicle abundances and 519 volatile concentrations indicate that the sintering particle pack is not only unimodal fine ash

- 520 and that some pieces of obsidian incorporated by the eruption column are re-plastered onto
- 521 the walls to re-sinter with other fragments and more ash.
- 522

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530

531 Appendix A

532

533 Whether there is enough time for silicate melt to flow viscously to allow the bubbles to shrink depends on the timescale for H₂O to resorb from vesicles (λ_{γ}) and the characteristic time 534 required for the melt to flow (λ_{η}). Both timescales are functions of solubility (C_e), diffusivity 535 536 (D), and melt viscosity (η), all of which are functions of temperature (Figure A1), and so they 537 change through the cooling ramp. The curves for $C_e(T)$ and D(T) also require a dissolved water 538 content to be assumed. We choose $C_{e,0}$, which is the initial solubility of water in the melt, in 539 wt.%, at the start of the cooling ramp. This is the lowest dissolved water content anticipated 540 during the experimental run. This value yields the highest viscosity and slowest diffusivity, and 541 so constitutes the most conservative assumption. In our analysis we use the time-averaged values of each parameter over the linear cooling ramp: $\overline{C_e}$, \overline{D} , and $\overline{\eta}$ respectively. 542 In order to estimate λ_{γ} we first calculate the mass (M_0) of H₂O contained within a bubble of 543 544 initial radius (R_0) at the start of the cooling ramp, using the equation of state of Pitzer and

545 Sterner (1994). We then calculate the volume (*V*) of melt that would be required to resorb this 546 mass of water:

547

548
$$V = \frac{M_o(R_o, P, T_o)}{\left[\frac{(\overline{Ce} - C_{e,o})}{100}\rho_m\right]},$$
 (A1)

549

where $M_0(R_0, P, T_0)$ indicates that M_0 is a function of initial bubble radius, experimental pressure, and initial temperature (via the equation of state), and ρ_m is density of the melt, taken as 2300 kg m⁻³. We assume that, if resorption goes to completion, then *V* will be a sphere of hydrated melt with radius $R_h = \sqrt[3]{3V/4\pi}$. The characteristic diffusion length scale at the end of the cooling ramp is estimated as $l_D = \sqrt{Dt}$. By setting $l_D = R_h$, we can estimate the duration of the cooling ramp required to fully resorb the water in the bubble by diffusion:

557
$$\lambda_{\gamma} = \frac{1}{D} \left(\frac{3M_o(R_o, P, T_o)}{4\pi\rho_m \frac{(\overline{C_e} - C_{e,o})}{100}} \right)^{\frac{2}{3}}.$$
 (A2)

558

The characteristic time required for the melt to flow viscously to allow the bubble to collapse is estimated by assuming that it scales with the ratio of the viscosity of the melt to the stresses driving collapse, which are the surface tension (Laplace) stress σ and the confining pressure, given by

564
$$\lambda_{\eta} = \frac{\overline{\eta}}{\left(\frac{\sigma}{R_{o}} + P_{o}\right)}.$$
 (A3)

567	Figure A–1: Solid curves are solubility (C_e), diffusivity (D), and viscosity (η) as functions of
568	temperature, all at 22 MPa H_2O pressure. Dash lines are time-averaged values of each
569	during constant rate cooling from 750°C to 550°C. Diffusivity and viscosity are calculated
570	under the conservative assumption that dissolved H_2O content is given by solubility at
571	750°C.

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- 670

671 **Figure Captions**:

672

673	Figure 1: Sample porosity (ϕ , in vol.%) as a function of experimental sintering time. High and
674	low porosities of heterogenous samples are linked by vertical tie lines. All porosities are
675	precise to $\pm 10\%$. Experiments of coarse ash only ran for 30 and 60 minutes, but did not
676	sinter. The equilibrium porosity of ϕ = 3 vol.% is shown as a dashed line (Wadsworth et al.,
677	2016; Gardner et al., 2018).

678

679 Figure 2: Photomicrographs of sintered unimodal ash. (a) Binary threshold image of G-1766 (fine 680 ash, t = 5 minutes) with incipiently sintered particles (white) dispersed in a continuous porous 681 network (black). (b) Reflected light image of G-1728 (fine ash, t = 20 minutes) with dense glass 682 and dispersed, contorted vesicles (in black). (c) Transmitted light image of the same sample 683 in (b) with isolated spherical (arrows) and highly contorted vesicles. Dashed line traces one 684 chain of connected vesicles. (d) Transmitted light image of G-1759 (fine ash, t = 47 minutes) 685 with isolated, contorted vesicles (arrows). (e) Transmitted light image of G-1758 (fine ash, t 686 = 60 minutes) with only spherical vesicles. (f) Reflected light image of G-1772 (wet ash, t = 30687 minutes) with incipiently sintered particles dispersed in a continuous porous network, now 688 mainly filled with epoxy.

689

Figure 3: Sintering textures of fine ash samples (open circles) as a function of equilibrium melt viscosity (η_e , in log Pa s) and experimental time. Solid lines demark approximate phases of constant texture. In Phase 1, individual particles remain loose and lack cohesion. In Phase 2,

particles are sintered only at their contacts and porosity is fully open. In Phase 3, particles
are merged together and vesicles are nearly sealed, but many vesicles are still multi-cuspate
shaped. In Phase 4, samples are dense glass with isolated vesicles. Dashed curves (with small
numbers) are for coarser ash sintered in Gardner et al. (2018).

697

698 Figure 4: Photomicrographs of sintered bimodal samples (all scale bars are 200 µm long). (a) 699 Reflected light image of G-1724 (50:50 fine:coarse ash, t = 20 minutes;) with incipiently 700 sintered particles in a continuous porous network, now mainly filled with epoxy. (b) Reflected 701 light image of G–1760 (50:50 fine:coarse ash, t = 45 minutes) with highly porous matrix 702 between closely spaced large particles; dashed line traces a large continuous pocket of large 703 vesicles >1 mm long. (c) Reflected light image of same sample as (b) but with low porosity 704 matrix where large particles are far apart. (d) Transmitted light image of G-1762 (50:50 705 fine:coarse ash, t = 60 minutes) with homogeneous matrix with spherical vesicles (arrows) 706 between large relict particles. (e) Reflected light image of G-1763 (75:25 fine:coarse ash t =707 45 minutes) with homogeneous porous matrix amongst large particles (outlined in white). 708 Large vesicle in the middle is part of a longer pocket \sim 300 μ m long. (f) Reflected light image 709 of G–1767 (50:50 fine:wet ash, t = 20 minutes) with highly porous matrix between closely 710 spaced large particles; dashed lines trace continuous pockets of large vesicles.

711

Figure 5. Temperature versus total run duration for fine ash samples that were either sintered at
 750°C (open circles) or sintered at 750°C and then cooled (solid circles). Final porosity (vol.%)

714	values are listed next to each sample. Note that samples plot along nearly linear cooling lines
715	(solid curves), showing that cooling occurred at a relatively steady rate.

716

717 Figure 6: (a) Reflected and (b) transmitted light images of G-1729 that sintered for 30 minutes at 718 750°C and then guenched. (c) Reflected and (d) transmitted light images of G-1789 that 719 sintered for 10 minutes at 750°C and then cooled to 550°C, over 25 minutes. Both have ~3 720 vol.% vesicles, but all are distorted in G-1789 (some marked by arrows), whereas half are 721 spherical in G–1729. Images (a) and (c) are 400 μ m x 400 μ m; scale bars in (b) and (d) are 50 722 μm long. 723 724 **Figure 7:** Porosity (ϕ , vol.%) as a function of sintering time for samples that started with dry 725 ash; symbols are the same as in Figure 1; porosities are precise to ±10%. Curves for 726 solutions of equation 3 for low-Pc sintering are shown for unimodal fine ash (solid lines) and 727 unimodal coarse ash (dashed lines), assuming ϕ_i = 45 or 60 vol.%. All mixed fine-coarse 728 bimodal samples fall between the end-member cases. The equilibrium porosity of $\phi =$ 729 3 vol.% is given as a limiting value (Wadsworth et al., 2016; Gardner et al., 2018). 730 731 Figure 8: Estimated duration of cooling step (over interval 750-550°C) required to resorb bubbles 732 of different initial size. Calculations (Appendix A) are based on simple scaling arguments and 733 are indicative only. Solid black line is the cooling time required to diffusively resorb all H_2O 734 within a bubble; dashed black line is the characteristic timescale of viscous shrinkage of the 735 bubble. Solid red line is the cooling time to 550°C for experimental run G-1783; solid blue line

is the cooling time to 550°C for experimental run G-1729 (during rapid quench); grey box
indicates range of initial bubble sizes expected in that sample. See main text for details.

738

739 Figure 9: Representative photomicrographs of North Mono obsidian pyroclasts (scale bars 740 shown). (a) Transmitted light image of sample P4B–I with coexisting spherical (S) and 741 distorted (D) vesicles. (b) Transmitted light image of sample P2–F with uniform distribution 742 of spherical vesicles. (c) Transmitted light image of sample P10–E with bands of highly 743 distorted vesicles. (d) Binary image of slice from High Resolution X-Ray Computed 744 Tomography (HRXCT) scan of P10–E (same as in c) showing that vesicles (some marked by 745 arrows) make up 0.04±0.004 vol.% of the obsidian. Widest length of sample is ~3 mm; see 746 supplemental materials from Gardner et al. (2017) for scanning methods. (e) Transmitted 747 light image of sample P4B–C with large pockets of vesicles dispersed in relatively dense glass 748 with few vesicles. (f) Transmitted light image of sample P10–I that consists of bands of glass 749 with numerous distorted vesicles (within the dashed lines) and regions of dense glass with 750 very few vesicles. Watkins et al. (2017) found that vesicle-poor regions in P10-I tend to be 751 rich in H_2O , whereas the bands with abundant vesicles are rich in CO_2 .





Run duration (minutes)















Run duration (minutes)







Figure A–1

Run ^a	P^b	T^{b}	t ^b	Fine ^c	Coarse ^c	Wet ^c	ϕ^d	$\log N_v^e$	Size ^e
	(MPa)	(°C)	(min)	(mg)	(mg)	(mg)	(vol.%)	(cm⁻³)	(µm)
G–1766	22	750	5	31.3	0	0	45.7	n.d.	-
G–1765	22	750	10	30.1	0	0	22.3	n.d.	-
G–1728	22	750	20	31.4	0	0	4.9	7.80	2–4
G–1729	22	750	30	31.9	0	0	3.2	7.66	1–15
G–1759	22	750	47	32.2	0	0	4.7	7.53	1–38
G–1758	22	750	60	32.8	0	0	5.4	7.50	2–48
G–1769	22	750	30	0	42.1	0	n.d.	n.d.	-
G–1768	22	750	60	0	40.2	0	n.d.	n.d.	-
G–1772	22	750	30	0	0	39.7	34.6	n.d.	_
G–1773	22	750	60	0	0	41.2	31.9	n.d.	-
G–1724	22	750	20	20.2	20.5	0	32.6	n.d.	_
G–1760	22	750	45	20.4	22.0	0	25.5/3.5	7.20	n.d.
G–1762	22	750	60	19.7	20.9	0	7.6	7.34	1–20
G–1761	22	750	20	30.0	10.0	0	16.2/6.7	7.54	1–10
G–1763	22	750	45	30.6	9.3	0	7.4/5.4	7.63	1–16
G–1764	22	750	60	30.0	10.2	0	3.1	7.59	1–84
G–1767	22	750	20	15.1	0	15.5	27.9/9.0	7.75	1–20

 Table 1: Experimental Run Conditions and Results

(a) Composition of obsidian used in all experiments (all oxides in wt.%, with all Fe reported as FeO: $SiO_2 = 76.53$;

TiO₂ = 0.06; Al₂O₃ = 13.01; FeO* = 0.79; MnO = 0.08; MgO = 0.02; CaO = 0.74; Na₂O = 3.87; K₂O = 4.91.

(b) Pressure (in MPa), temperature (in °C), and run duration (in minutes) of the experiment.

(c) Mass of "fine" (dry, \leq 45 µm), "coarse" (dry, 250–500 µm) and wet (hydrated, 125–180 µm) powders in sample.

(d) Porosity (in vol.%) of the experiment, including all vesicle types. If two values are listed, the first is porosity of matrix between closely spaced large particles, and the second is the porosity of the matrix far away from large particles.

(e) Number density (in numbers per cubic centimeter) of all vesicles present and range of sizes of spherical vesicles only; n.d. = not determined; – means no spherical vesicles present.

Table 2: 0	Cooling E	xperin	nents, Co	onditio	ns and Resu	ults				
Run	\mathbf{P}^{a}	T_s^a	t_s^a	T_f^b	t_f^b	Fine ^c	Coarse ^c	ϕ^d	$\log N_v^e$	Size ^e
	(MPa)	(°C)	(min)	(°C)	(min)	(mg)	(mg)	(vol.%)	(cm⁻³)	(µm)
G–1788	22	750	5	550	22:44.6	30.7	0	7.4	n.d.	n.d.
G–1787	22	750	10	650	12:06.6	32.3	0	10.4	n.d.	n.d.
G–1789	22	750	10	550	24:53.4	34.3	0	3.2	n.d.	n.d.
G–1786	22	750	30	700	6:10.2	29.4	0	0.9	6.83	n.d.
G–1784	22	750	30	650	11:17.3	30.3	0	0.3	5.41	15–35

0

n.d.

0*

0

15.9/1.8 6.25

... -

30

30

(a) Pressure (in MPa) and temperature (in °C) of sintering for duration t (in minutes).

550

650

(b) Final temperature (in °C) of cooling and duration of cooling (in minutes) to final temperature.

(c) Mass of "fine" (dry, \leq 45 µm) and "coarse" (dry, 250–500 µm) powders in sample.

(d) Porosity (in vol.%) of the experiment, including all vesicle types. If two values are listed, the first is the porosity of matrix between closely spaced large particles, and the second is porosity of matrix far away from coarse particles.

22:56.7

11:01.2

30.3

20.2

0

20.8

(e) Number density (in numbers per cubic centimeter) and range of sizes of all vesicles present; n.d. = not determined. N_{ν} in G–1783 is 0 per cm³.

G-1783

G-1785

22

22

750