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Trapped bubbles keep pumice afloat and gas diffusion makes pumice sink

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Trapped bubbles keep pumice afloat and gas diffusion makes pumice sink

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Highlights

<u>X-ray</u> microtomography shows that hot (500 °C) and <u>ambient-</u> <u>temperaturepumice</u> trap gas.

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The size distributions of trapped gas clusters match <u>percolation</u> theory.

Gas trapping can explain why pumice floats on water.

Pumice sinks because of outward diffusion of trapped gas.

Abstract

Pumice can float on water for months to years – long enough for pumice to travel across oceans and facilitate the spread of species. Long-lived pumice floatation is unexpected, however, because pumice pores are highly connected and water wets volcanic glass. As a result, observations of long floating times have not been reconciled with predictions of rapid sinking. We propose a mechanism to resolve this paradox – the trapping of gas bubbles by water within the pumice. Gas trapping refers to the isolation of gas by water within pore throats such that the gas becomes disconnected from the atmosphere and unable to escape. We use X-ray microtomography to image partially saturated pumice and demonstrate that non-condensable gas trapping occurs in both ambient temperature and hot (500 °C) pumice. Furthermore, we show that the size distribution of trapped gas clusters matches predictions of percolation theory. Finally, we propose that diffusion of trapped gas determines pumice floatation time. Experimental measurements of pumice floatation support a diffusion control on pumice buoyancy and we find that floatation time τ scales as $\tau \propto L2D\theta 2$ where L is the characteristic length of pumice, D is the gas–water diffusion coefficient, and θ is pumice water saturation. A mechanistic understanding of pumice floatation is a step towards understanding how

pumice is partitioned into floating and sinking components and provides an estimate for the lifetime of pumice rafts in the ocean.

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Keywords

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buoyancy

capillary processes

percolation theory

submarine volcanism

X-ray microtomography

Notation

v	velocity	р	volume–area coefficient
μ	dynamic viscosity	τ	pumice floatation timescale
Y	surface tension	$D_{\scriptscriptstyle eff}$	effective diffusion coefficient
ρ	density	θ	water saturation
g	gravity	D	liquid–gas diffusion coefficient
к	permeability	d	mean pore throat diameter
$oldsymbol{\phi}$	connected porosity	Р	pressure
h	height	R	pore throat radius
V_{w}	volume of water absorbed	Т	temperature
t	time	$L_{\scriptscriptstyle w}$	glass wall thickness
S_a	pumice surface area	$D_{\scriptscriptstyle w}$	water thermal diffusivity
n	number of occurrences	T_i	initial temperature
S	sites or pores	T_{f}	final temperature
β	power law coefficient	V_i	initial volume
а	spatial dimension	$V_{\rm f}$	final volume
f	fractal dimension a cluster	ξ	gas saturation
S _{max}	maximum size of a trapped gas cluster	ξ_i	initial gas saturation
L	pumice diameter	ξ*	neutral buoyancy gas saturation
A	surface area of trapped gas clusters	$ ho_r$	glass density
V	trapped gas volume	${oldsymbol{ ho}}_{l}$	liquid density

1. Introduction

<u>Pumice</u> is a highly vesicular <u>volcanic rock</u> with a <u>porosity</u> high enough that it can float. Rafts of volcanic pumice can transit lakes and oceans and circle the globe (e.g., Richards, 1958, Risso et al., 2002, Bryan et al., 2004, von Lichtan et al., 2016). For example, pumice from the 1952 eruption of Volcán Barcena on Isla San Benedicto, 600 km west of Mexico, floated for at least 560 days and drifted over 8700 km (Richards, 1958). The 2012 eruption of Havre submarine volcano created a 1.5 km³ pumice raft that spread over 550,000 km² within three months (Carey et al., <u>2014</u>, <u>Jutzeler et al.</u>, <u>2014</u>). Pumice rafts have been shown to facilitate the dispersal of species such as barnacles, corals, algae, and gastropods (Bryan et al., 2012) because marine organisms grow on, and ocean currents advect, pumice (Richards, 1958, Jokiel, <u>1984, Bryan et al., 2004</u>). While pumice rafts are relatively common and it is well known that <u>ambient temperature</u> pumice can float for long periods of time, the enduring buoyancy of pumice is surprising because pumice pores are almost entirely connected and water wets pumice (Whitham and Sparks, 1986, Vella and Huppert, 2007). Quantitative models for pumice saturation predict that ambient temperature pumice should sink orders of magnitude more rapidly than is observed (Vella and Huppert, 2007). The floatation time discrepancy between observations and the Vella and Huppert (2007) model suggests that simple gas displacement by an infiltrating water front is not sufficient to explain why ambient temperature pumice can float for years. By comparison to ambient temperature pumice, hot pumice (e.g., >300 °C) sinks almost immediately and the tendency for air-filled pumice to sink increases with pumice temperature (Whitham and Sparks, 1986, Dufek et al., 2007, Allen et al., 2008, Jutzeler et al., 2016). Rapid water ingestion by hot pumice has been attributed to coolinginduced gas contraction (Whitham and Sparks, 1986, Cashman and Fiske, 1991, Allen et al., 2008) and hydrodynamic instabilities due to steam generation (Dufek et al., <u>2007</u>). Air-filled hot pumice placed in water does not, however, completely saturate even at high (500 °C) temperatures (Allen et al., 2008). As a result, we wish to understand how gas remains within initially hot pumice and what differences and similarities exist between saturation of ambient temperature and hot <u>non-condensable gas</u> filled pumice. Pumice, with porosities of 50 to >90 percent, is a porous medium. Water saturation of pumice is an example of two-phase flow in porous media and requires the replacement of a defending fluid (air or magmatic gases) with an invading fluid (liquid water). Twophase flow in porous media has been widely studied in the context of the vadose zone, oil recovery, CO₂ sequestration, and gas sparging. In addition, water infiltration of pumice is a manifestation of a particular type of two-phase flow, imbibition, because water is the wetting phase. During imbibition the arrangement of fluid, or wetting pattern,

can range from one where nearly all the pores are filled with the invading fluid to one where the defending fluid remains trapped in clusters (e.g., Lenormand and Zarcone, 1984). Trapped gas clusters (Fig. 1), pockets of non-wetting fluid that are surrounded by the wetting fluid, are not only characteristic of two-phase flow in porous media but are very difficult to mobilize because of surface tension. Indeed, gas trapping is a mechanism employed for long term CO_2 sequestration (e.g., Ide et al., 2007, Benson and Cole, 2008).



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Fig. 1. Mechanisms of gas trapping. Illustrations of gas trapping by (a) bypass and (b) snap-off trapping in <u>pumice</u> with connected <u>pores</u>. In both cases capillary forces draw water into pores such that water completely surrounds the <u>gas phase</u> and the gas is unable to escape.

We hypothesize that pumice floats for long periods of time because of the occurrence of gas trapping (either air or non-condensable magmatic gases) in isolated gas clusters during water infiltration. We use X-ray microtomography to test the hypotheses that gas trapping occurs in both hot and ambient temperature pumice, that gas trapping can result in a high enough residual gas saturations to keep pumice afloat, and that percolation theory can describe gas trapping in pumice. While trapped gas may buoy pumice, we hypothesize that the outward diffusion of gas trapped in bubbles eventually causes pumice to sink. We test this gas diffusion hypothesis by conducting experiments where we measure the floatation time of dry and ambient temperature

pumice on artificial seawater in a controlled laboratory setting. We then compare our results and pumice floatation times from four other studies with a prediction for pumice floatation time based on gas-diffusion out of a porous medium.

1.1. Gas trapping in porous media

Gas trapping has been observed in experiments, dictates wetting patterns, and controls residual non-wetting saturation of porous media (e.g., <u>Blunt and Scher, 1995</u>, <u>Iglauer et al., 2013</u>, <u>Geistlinger and Mohammadian, 2015</u>). A key element that promotes gas trapping is the slow advance of the invading fluid such that capillary forces dominate over viscous forces. In other words, the Capillary number

(1)Ca=vμγ,

where *v* is the characteristic velocity, μ is the wetting fluid viscosity, and *y* is surface tension on the interface between the two fluids, is very small, Ca \ll 1 (Lenormand and Zarcone, 1984, Wilkinson, 1984, Blunt and Scher, 1995).

We calculate a Capillary number for water infiltration into dry pumice using μ =10–3Pas for the viscosity of water at room temperature, γ =0.072Nm–1 for the surface tension at the <u>air–water interface</u>, and by estimating the velocity of water infiltrating pumice using <u>Darcy's law</u>. For pumice floating on water, the maximum head gradient is set by the <u>hydrostatic pressure</u> at the bottom edge of the pumice and the <u>capillary pressure</u>. By assuming a constant pore radius and a hemispherical gas– water meniscus we can write the liquid velocity as

(2)v=κμφ(ρgh+2γR),

where κ is permeability, ϕ is connected porosity, ρ is water density, g is gravity, h is the height of pumice in water, and R is pore throat radius. Pumice porosities can vary widely, but typical values are 50 to 90 percent (pumice with rock equivalent densities between 2.4 and 3.0 g cm⁻³ must have porosities of at least 58 to 67 percent, respectively, to initially float). Pumice permeabilities are more difficult to estimate, but measured values range from 10–14–10–10m2 (e.g., Klug and Cashman, 1996, Tait et al., 1998, Saar and Manga, 1999, Klug et al., 2002; Rust and Cashman, 2004, Rust and Cashman, 2011; Mueller et al., 2005; Wright et al., 2006, Wright et al., 2009; Degruyter et al., 2010). We note that permeability may also be a function of saturation, i.e., relative permeability is not unity. We consider pumice that is immersed 0.01–1 m in water and pores that have radii of 0.1–0.001 mm. From these input parameters, $3.8 \times 10-11 < Ca < 4.3 \times 10-4$, which shows that capillary forces dominate in pumice.

We also use pumice saturation measurements from <u>Whitham and Sparks (1986)</u> to estimate water infiltration velocities according to,

(3)v=∆Vw∆tSa¢

where Vw is the volume of absorbed water, *t* is time, Sa is pumice surface area, and ϕ is connected porosity. We estimate Sa by assuming the pumice <u>clasts</u> are spherical. Fourteen pumice saturation measurements by <u>Whitham and Sparks (1986)</u> reveal average initial infiltration velocities over the first five minutes of water exposure of 2.74±0.97×10–2cmhr–1 and Capillary numbers of 7.62±2.69×10–8. These Ca estimates indicate that capillary forces dominate. Thus, the saturation of pumice should be considered at the <u>pore scale</u> and may lead to gas trapping. In the next sections we introduce two mechanisms that allow gas trapping to occur, the percolation models that simulate them, and the gas-trapping predictions percolation theory makes.

1.1.1. Bypass trapping

Bypass trapping (Fig. 1a) causes trapping through the sequential filling of pore throats such that the defending fluid becomes surrounded by the invading fluid before it can escape (e.g., Chatzis et al., 1983). Invasion percolation models simulate bypass trapping by (1) representing the porous medium as a network of spheres (pores) and cylinders (pore throats); (2) prescribing a capillary entry pressure for each throat and pore; and (3) filling throats and pores from highest to lowest pressure. As a result, a pore or series of pores with low capillary pressures may be completely surrounded before being filled and thus become trapped (but may contract or expand if compressible).

1.1.2. Snap-off trapping

Snap-off trapping occurs because for very <u>low flow</u> rates (or capillary numbers) the wetting fluid can flow along edges of the pore walls due to <u>surface</u> roughness (e.g., <u>Lenormand and Zarcone, 1984</u>). This wall-hugging thin film can swell as the wetting fluid invades. As the film swells in the smallest throats, it completely displaces the non-wetting phase and can disconnect the non-wetting phase from any neighbors (Fig. 1b). Snap-off trapping is simulated by Bond percolation. When snap-off and bypass trapping are both possible, bypass trapping is favored because it occurs for higher capillary pressures.

1.1.3. Percolation model predictions

Both invasion and bond percolation theory make predictions about the size distribution of residual trapped gas clusters. Because percolation systems are scale invariant at

the <u>critical point</u> (when fully percolated), the number of occurrences, n(s), of trapped gas clusters containing *s* sites or pores scales according to a power-law, $(4)n(s) \sim s - \beta$,

where β =a+ff and *a* is the spatial dimension and *f* is the <u>fractal</u> dimension of the cluster. In 3D: a=3, f=2.52, and β =2.19 (<u>Stauffer, 1979</u>, <u>Wilkinson and Willemsen, 1983</u>). The maximum size of the trapped gas cluster is limited by the samples size, *L*, smax~Lf(<u>Wilkinson, 1986</u>). Experimental studies of imbibition in porous media have shown that trapped gas size distributions match Equation (4) (e.g., <u>Geistlinger and</u> <u>Mohammadian, 2015</u>). Examining the size distribution of trapped gas bubbles in a natural system, such as pumice, can therefore help distinguish if percolation theory (either Bond or Invasion) applies.

2. X-ray microtomography

X-ray microtomography (μ XRT) allows us to see both the internal structure of pumice (e.g., Polacci et al., 2006, Wright et al., 2006, Degruyter et al., 2010, Giachetti et al., 2011, Voltolini et al., 2011, Carey et al., 2013) and, when multiple fluids are present within the pores, the distribution of those fluids (e.g., Wildenschild et al., 2002). We use μ XRT to (1) test the hypothesis that gas trapping occurs in pumice; (2) determine if percolation theory can describe the size distribution of trapped gas; (3) examine the size distribution of trapped gas for insights into the mechanisms (bypass or snap-off trapping) that occur in pumice.

2.1. Experimental set-up

We used μ XRT to image the internal distribution of liquid and air in six uncut pumice <u>clasts</u>. Before the μ XRT, we conducted saturation experiments on <u>ambient</u> temperature and hot (500 °C) pumice from Santa Maria, Guatemala and Medicine Lake, California (<u>Table 1</u>). We set ambient temperature pumice on a 13 wt% solution of <u>potassium iodide</u> (KI) for ~20 h such that the pumice could adsorb the liquid. KI, a common chemical dopant, increases contrast of the μ XRT images (greatly improves the segmentation of air and water in the images), while maintaining a <u>surface tension</u> within 1% of water (<u>Aveyard and Saleem, 1976; Wildenschild et al., 2002</u>).

Table 1. µXRT samples (uncut clasts) and measurements.

Sample name	SM01	SM02	SM04	SM05	ML01	ML02
Sample description	1902 Santa Maria	1902 Santa Maria	1902 Santa Maria	1902 Santa Maria	Medicine Lake	Medicine Lake
Pumice mass (g)	0.07	0.19	0.13	0.04	0.15	0.76

Sample name	SM01	SM02	SM04	SM05	ML01	ML02
Pumice dimensions mm ³	$9 \times 4.5 \times 4$	$11 \times 7 \times 4.5$	$11 \times 6 \times 5$	$6.5 \times 3 \times 3$	$9.5 \times 5 \times 5$	$5 \times 5 \times 4$
Temperature	ambient	ambient	500 °C	500 °C	ambient	ambient
Ending buoyancy	floating	neutrally buoyant	not floating	not floating	not floating	barely floating
Porosity	0.70	NA	0.73	NA	0.63	0.67
Connected porosity	0.70	NA	0.73	NA	0.55	0.65
Liquid volume/total volume	0.30	NA	0.56	NA	0.13	0.31
Gas volume/total volume	0.40	0.36	0.17	0.13	0.50	0.36
Liquid saturation (liquid volume/pore volume)	0.43	NA	0.77	NA	0.21	0.46
Calculated wet density/KI density	0.86	NA	1.07	NA	0.83	0.93
Trapped gas bubble number density (cm ⁻³)	9.6 × 10⁵	3.0×10^{6}	4.4×10^{6}	4.2×10^{6}	7.2×10^{6}	2.3 × 10 ⁶
Vesicle number density (cm ⁻³)	8.3 × 10 ⁶	NA	1.5 × 10 ⁷	NA	$9.6 imes 10^{6}$	5.3 × 10 ⁶
β : Power law exponent	2.02	1.90	1.95	2.10	1.83	1.51
Smallest bubble fit to power law (mm³)	5.9 × 10-5	1.3 × 10 ⁻⁵	2.3 × 10 ⁻⁵	3.1 × 10 ⁻⁶	1.3 × 10-4	1.0×10^{-6}
p: Trapped bubble volume to surface area exponent	0.82	0.82	0.77	0.76	0.75	0.83
Size of analyzed volume (mm³)	0.94	0.37	0.37	0.94	0.94	0.37

While μ XRT analysis is done on subvolumes of larger <u>clasts</u>, we expect the subvolumes to be representative of the larger <u>pumice</u> in part because the pumice are relatively small (i.e., <1 g). Furthermore, the selected pumice samples do not have bread crust textures such that we expect differences between rim an internal <u>porosities</u> to be minimal.

We heated two of the pumice samples (SM04 and SM05) to 500 °C, quenched these pumice clasts in the KI dopant, and allowed them to stay in the solution for ~10 min. All pumice pores were filled with air at the time of KI exposure. To preserve the internal fluid distributions for μ XRT imaging, we then rapidly removed and encased the pumice in wax.

We carried out the μ XRT imaging at the Lawrence Berkeley National Lab Advanced Light Source on beamline 8.3.2. We conducted the scans using 30 kev monochromatic X-rays and a 5X lens (resolution of 1.22 μ m/pixel). We used the TomoPy gridrec algorithm to reconstruct the 3D image stacks (<u>Gürsoy et al., 2014</u>). To minimize ring artifacts, we selected a subvolume from each data set of either 0.37 or 0.94 mm³ for data processing. We used Fiji's Trainable Weka Segmentation plugin – which employs multiple machine learning algorithms – to segment air, water, and rock within each image sequence (Hall et al., 2009). To train classifiers for each image sequence, we manually outlined vesicles and trapped air on fifty different images. After checking and retraining the classifiers as necessary, we applied the trained classifiers across the images sequences to segment air and rock (thereby creating two sets of binarized image sequences for each sample).

We also made the image sequences binary by applying a greyscale threshold in Fiji, but determined that the machine learning method reduced the effect of annular ring artifacts and better preserved thin glass walls compared to the traditional threshold based binarization method. Despite the advantages of the machine learning method, two image sets (SM02 and SM05) had glass walls that were too thin to resolve and we do not report porosities or vesicle size distributions for these samples.

After we segmented each data set, we loaded the binary images into Avizo where we identified and quantified the volume, surface area, and orientation of individual bubbles and vesicles. Here we refer to the pores of the pumice (that can be filled with either gas or liquid) as vesicles and areas where the <u>gas phase</u> is present as bubbles. Because most samples had highly interconnected porosities, we separated connected vesicles using a watershed algorithm before measuring vesicles sizes and orientations (Supplementary information). No separation was applied to the gas bubbles. As a result, any observed and reported gas bubbles were truly isolated. We note, however, that some gas bubbles may appear connected (by one or two voxels) when they are not if glass walls are thinner than one pixel. To correct for very thin glass walls we use a "neighborhood" value of six in the Avizo labeling module such that bubbles must share at least one voxel face to be considered connected.

Errors in generating the bubble and vesicle size distributions stem from two main sources: (1) ring artifacts in the original greyscale image and (2) vesicle walls that are thinner than the voxel resolution and that may also lead to overestimates of connected porosity (Fig. S1). Through examination of twenty 2D images from each dataset, we estimate that ring artifacts result in the mischaracterization of gas or water in <1 volume percent of each pumice. While the machine learning method for image segmentation reduced the effects of ring artifacts compared to threshold based segmentation, it introduced a number of very small (artifact) bubbles and vesicles that do not contribute to the total volume fraction of pores or trapped gas. As a result, we filtered the data to

exclude any bubbles or vesicles with less than a 4.16 pixel radius or smaller than 10–6mm3.

2.2. X-ray microtomography results

We find trapped gas clusters in all pumice we imaged (Fig. 2, Fig. 3). Trapped gas cluster size varies and gas clusters fill part of, single, and multiple vesicles (Fig. 2, Fig. 3). Because connected porosities for most samples are close to or equal to the total porosity, isolated gas pockets are due to trapping by liquid instead of isolation by rock (Table 1). Furthermore, we found that ambient temperature pumice contain larger volume percentages of trapped gas (36–50%) than the hot pumice (13 and 17%) (Table 1).



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Fig. 2. X-ray microtomography images. 2D slices through μ XRT images of <u>pumice</u> containing water and trapped gas. While hot pumice (SM04 and SM05) contain trapped gas, they hold smaller volume percentages compared to <u>ambient</u> <u>temperature</u> pumice. <u>Contacts</u> between rock, liquid, and gas demonstrate that the liquid is the wetting fluid (see red box). While glass walls are too thin to resolve in places (particularly in SM02 and SM05), we do not see any evidence in the μ XRT that the thin glass walls in pumice are broken or damaged due to cleaning in an <u>ultrasonic</u> bath. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



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Fig. 3. Shapes of trapped gas bubbles. 3D μ XRT images of trapped gas bubbles within <u>pumice</u>. Colors in this figure are chosen at random to identify separate gas bubbles. The top rows show the largest gas bubbles and the bottom row shows all gas bubbles within a single pumice. A single interconnected gas bubbles extends throughout many <u>pores</u> in pumice SM01. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

We use µXRT measurements of phase (liquid and water) saturations to estimate pumice buoyancy, assuming glass densities of 2.4 g cm⁻³, and find that the observed trapped gas saturations are high enough to allow pumice to float. These density calculations match our observation that many of the pumice clasts were floating or neutrally buoyant when we encased them in wax. Our density calculation of ML01, however, does not match our observation that ML01 was not floating (Table 1). Discrepancies between observations and density estimages suggest that gas and liquid saturation within pumice is likely heterogeneous. Indeed, we visually observed areas of more and less gas saturation within µXRT images.

The size distributions of both trapped gas (black) and segmented pores (grey) are shown in Fig. 4. For the majority of samples (ML02, SM01, SM02, and SM05) at least a third of the trapped gas is contained within one large cluster. These clusters, in cases where the vesicle size distributions are known (SM01 and ML02), greatly exceed the maximum vesicle size and Fig. 3 shows how these largest trapped bubbles can extend throughout multiple vesicles. The trapped bubble size distributions, however, show that there are multiple modes of trapped bubbles sizes. In sample ML01, the trapped bubble size distribution mirrors the vesicle size distribution.



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Fig. 4. Bubble and vesicle size distributions. Trapped gas bubbles, locations where only the gas is present, are shown in black, and vesicles, <u>pumice pores</u> that can be filled with liquid or gas, are in grey. At least a third of the trapped gas is contained with a single large bubble for most of samples. The existence of large trapped bubbles, in combination with the vesicle size distribution, demonstrates that at least the largest gas clusters often extend through multiple pores. Bubbles and vesicles are binned into fifty logarithmically spaced bins and each bin is divided by the total volume of air or vesicles. <u>Volume fraction</u> is not weighted by bin width such that the sum of all bins equals one.

2.3. Comparison to percolation theories

Here we test the hypothesis that percolation theory can describe gas trapping in pumice by fitting Equation (4) to the observed distributions of trapped gas clusters using the maximum likelihood method (Clauset et al., 2009, Iglauer and Wülling, 2016). We find that Equation (4) fits the observed distributions well and that fitted power-law coefficients range from 1.51–2.10, which (other than the coefficient for ML01) are close to the value predicted by percolation theory (β =2.19) (Fig. 5; Table 1). These power-law fits lend support to a percolation theory treatment of pumice saturation.



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Fig. 5. Power-law fits of bubble-size distributions. Measured distributions of trapped bubble volume (grey circles) and the fitted <u>power-law distributions</u> (dashed lines). The *y*-axis shows the probability that a bubble is larger than a given volume, *V*. With the exception of ML02, the fitted power-law coefficients, β , are consistent with the value predicted from <u>percolation</u> theory, $\beta = 2.19$. Because our <u>machine learning</u>segmentation method generated very small (artifact) bubbles, we removed bubbles smaller than 10⁻⁶ mm³ before fitting a power-law coefficient.

Percolation theory also predicts:

A∝Vp,

where *A* is the surface area of the trapped clusters, and *V* is the volume of the trapped clusters (<u>Stauffer, 1979</u>). From our data sets, we empirically find p=0.75-0.83 and the R2values for these fits are 0.95-0.99 (<u>Table 1</u> and Fig. S2). Values of *p* greater than 2/3 demonstrates that the gas bubbles are non-spherical.

3. Mechanisms that cause pumice to sink

The previous sections demonstrated that gas trapping occurs in <u>pumice</u>, that <u>ambient</u> temperature pumice traps more gas than hot (500 °C) pumice, that gas trapping can lead to high enough <u>residual gas</u> saturations to allow pumice to float, and that <u>percolation</u> theory can describe the distribution of trapped gas clusters. The occurrence of gas trapping does not, however, explain why pumice, after floating for days or months, eventually sinks.

3.1. Pumice floatation experiments

To examine why pumice sinks, we conducted pumice floatation experiments using pumice from the 1902 plinian eruption of Santa Maria Volcano, Guatemala. We placed the pumice in artificial seawater and measured the time it took for the pumice to sink. Before the experiments, we cleaned the pumice in an ultrasonicator for four hours and dried the pumice in an oven at 65 °C for 12 h. We specifically chose a temperature lower than 100 °C to ensure that we did not break pumice walls during the drying process and used μ XRT to confirm that the pumice was dry prior to experiments. Table 2 shows pumice weights, sizes, and characteristics. While we did not measure the volume of each pumice clast, we estimated volume by assuming porosities of 80% and glass densities of 2.4 g m⁻³.

Sample name	Dry weight (g)	Estimated volume (cm ³)	Floatation time (days)
SM_F02	0.30	0.63	17.3
SM_F03	0.25	0.53	22.7
SM_F05	0.46	0.95	20.2
SM_F06	0.68	1.42	17.0
SM_F09	0.61	1.28	195.5
SM_F10	0.06	0.13	2.6
SM_F11	0.55	1.15	28.2
SM_F21	0.54	1.12	2.5
SM F22	0.12	0.24	20.1

Table 2. Floatation time measurements for Santa Maria <u>p</u>	<u>umice</u> .
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To initiate the floatation experiments, we dropped the pumice from a height of 4 cm into individual containers of artificial seawater. These containers were covered to prevent evaporation and contamination. We monitored the pumice with a time lapse camera that could determine when each clast sank to the nearest minute.

3.2. Pumice floatation timescale

Here we propose that the diffusion of trapped gas bubbles out of the pumice (and eventually to the atmosphere) causes pumice containing trapped gas clusters to sink. We envisage that the diffusion process is analogous to <u>Ostwald ripening</u> where small (and thus higher pressure) trapped gas clusters diffuse into larger clusters and eventually to the atmosphere. If <u>gas diffusion</u> does control pumice buoyancy through time, then pumice floatation time should scale like a diffusive process where $(5)\tau$ ~L2Deff,

and where *L* is the mean diameter of the pumice, and Deff is the effective <u>diffusion</u> <u>coefficient</u>. The effective diffusion coefficient for trapped gas in pumice, as in other <u>porous media</u>, is not just the gas–liquid diffusion coefficient, but should be weighted by the connectivity of the porous media (i.e., porosity and partial saturation). From <u>Hunt et al. (2014)</u> we write effective <u>diffusivity</u> as Deff=D02, where θ is water saturation (fraction of <u>pore space</u> filled by water) and *D* is the liquid–gas diffusion coefficient, such that

(6)τ∝L2Dθ2.

We test the hypothesis that diffusion of trapped gas out of pumice allows pumice to sink by comparing experimental measurements of pumice floatation time and volume (Whitham and Sparks, 1986, Manville et al., 1998, Risso et al., 2002, White et al., 2001) with Equation (6) and where L~V2/3 (Fig. 6). Because we do not know θ for any individual pumice, and because, if our hypothesis is correct, θ changes through time, we consider Equation (6) with a range of saturation values (8–80%).



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Fig. 6. Floatation time as a function of <u>pumice</u> volume. Measurements of pumice floatation from this study, <u>Whitham and Sparks (1986)</u>, <u>Manville et al. (1998)</u>, <u>Risso et al. (2002)</u>, and <u>White et al. (2001)</u>. Pumice floatation time predictions from this study (Equation (6)) and <u>Vella and Huppert (2007)</u> are also shown. Measured floatation times fit the trend predicted by Equation (6) (grey bar). Filled symbols represent measured sinking times while open symbols represent projected sinking times.

3.3. Pumice floatation results and model comparison

Fig. 6 shows measurements of floatation time and volume (this study and four others) against predictions from Equation (6) and where $D=1.9\times10-5cm2s-1$ is the air–water diffusion coefficient at room temperature. Experimental measurements of pumice floatation times generally match predictions from Equation (6) (Fig. 6). We also list pumice floatation times from our experiments in Table 2.

It is worth noting, however, that if pumice are highly non-spherical, then we overestimate the effective length scale (shortest pumice axis). Furthermore, the

timescale for pumice floatation may depend strongly on θ , not only because θ affects the diffusivity, but also because more diffusion must occur to sink a pumice with an initially high trapped gas content.

While we are not the first to recognize that pumice floatation time scales like a diffusive process (<u>Manville et al., 1998</u>), diffusion of trapped gas out of the pumice has not previously been identified to be the controlling process. <u>Fig. 6</u>, in combination with observations of gas trapping, suggests that it is the diffusion of trapped gas out of pumice that causes raft and other floating pumice to eventually sink.

4. Discussion

4.1. Gas trapping timescale

We find that water saturation of <u>pumice</u> is a two-step process. First, capillary and <u>hydrostatic pressures</u> drive water into <u>pores</u>. Water invasion often leads to gas trapping (e.g. section <u>1.1</u>) and ends when there are no longer pores to invade. Second, gas slowly diffuses out of trapped <u>gas pockets</u> thereby creating more space for the liquid (section <u>3</u>). Here we estimate the timescale for the first of these processes – the time for pumice to reach its residual saturation state.

If pumice can be modeled as a bundle of horizontal parallel cylindrical tubes and <u>capillary pressures</u> drive fluid into the tubes, then the Washburn equation can describe the timescale for water saturation over horizontal distance L,

(7)t=4L2 μ yd,

where *d* is mean pore throat diameter. Equation (7) suggests that pumice with 0.05 mm diameter pores should reach its residual saturation state very rapidly: 2.7 s for 5 cm pumice and 18 min for 1 m pumice. In other words, if a dry pumice is set on water, then capillary forces draw in liquid quickly and set the wetting pattern and trapped gas geometry in time *t*. We emphasize that pumice can remain buoyant following water invasion due to gas trapping.

4.2. Water saturation of hot versus cold pumice

We find that <u>residual gas</u> saturation is lower for hot pumice (500 °C) compared to <u>ambient temperature</u> pumice, and that these lower gas saturations can account for the buoyancy differences of hot versus cold pumice (<u>Table 1</u>). Because hot pumice may rapidly saturate due to <u>non-condensable gas</u> contraction (<u>Whitham and Sparks,</u> <u>1986</u>, <u>Cashman and Fiske</u>, <u>1991</u>, <u>Allen et al.</u>, <u>2008</u>), we compare the pressure differences generated by three processes that can drive fluid into pumice: water column weight, capillary action, and <u>gas cooling</u> and contraction. Gravity induces a pressure difference when pumice is submerged in water proportional to

(8) $\Delta P=(\rho I-\rho g)gL\sim 103Pa$,

choosing L=10cm. The pressure difference induced by capillary forces is $(9)\Delta P=2yR\sim 1.5\times 104Pa$,

where $R=10\mu m$ is pore throat radius. Lastly, the pressure difference created by gas contraction is proportional to the change in temperature of the gas by the ideal gas law such that

(10)∆P=Pi∆TTi.

Assuming that the initial pressure, Pi, of the gas is atmospheric (~105Pa), the initial temperature of the gas, Ti, is 800 K, and the change in <u>gas temperature</u>, ΔT , is 500 K, then $\Delta P \sim 3.7 \times 104Pa$.

Gas contraction can thereby produce pressure differences as large as those produced by capillary forces which suggests that, because <u>pressure gradients</u> drive <u>fluid flow</u>, gas contraction can be a relevant process for liquid ingestion. Furthermore, we expect that heat transfer from pumice to liquid is rapid (<1 s) because pumice walls are very thin, (11)t~Lw2/Dw,

where Lw~10-5–10–6m is the glass wall thickness (Fig. 2) and Dw~10–7m2s–1 is water thermal diffusivity. Rapid heat transfer suggests that gas contraction occurs simultaneously with capillary induced pore filling. We note, however, that gas contraction may not proceed prior to pore filling because hot gas must contact cool liquid for heat transfer (e.g., Stroberg et al., 2010).

Following the derivation of the Washburn equation (Equation (7)), we derive a new timescale for the initial stage of water ingestion into pumice when both gas contraction and capillary forces drive fluid flow:

(12)t=4μL2R2(2γR+PiΔTTi).

Gas contraction in hot pumice may help to explain the differences in residual gas saturation of hot and cold pumice. When pressure doesn't change, according to the ideal gas law

(13)ViVf=TiTf,

where Vi and Ti are initial volume and temperature of gas within a pumice and Vf and Tfare final volume and temperature, respectively. We therefore expect that when air is cooled from 800 to 300 K, Vf=0.37Vi. Our experiments showed that hot pumice contained gas in ~23% of <u>pore space</u> while ambient temperature pumice retained gas in 54–79% of its pore space. This <u>reduction</u> in trapped gas volume is ~0.23/0.7=0.32 and is approximately the value (0.37) predicted by gas contraction alone.

Trapped bubble size distributions matched <u>percolation</u> theory for both hot and ambient temperature pumice (Fig. 5). Invasion percolation theory works by prescribing a capillary entry pressure for each pore and by filling pores from highest to lowest entry pressure. Percolation theory may apply to hot and non-condensable gas-filled pumice because gas contraction introduces a new, yet additive, pressure difference across each pore and thus does not fundamentally change the mechanism for pore filling.

4.3. Bypass versus snap-off trapping

While the fit of Equation (4) to the measured trapped gas bubble size distributions supports a percolation control on pumice saturation, we have yet to determine if gas is trapped by bypass or snap-off mechanisms. The trapped bubble area-to-volume ratios show that the trapped bubbles are non-spherical (Table 1). Furthermore, we observe trapped gas within many interconnected pores (Fig. 3). Because snap-off trapping leads to gas bubbles trapped in single pores (e.g., Fig. 1), these observations support a bypass trapping mechanism. Furthermore, bypass trapping is topologically favored over snap-off trapping when both are possible.

4.4. Gas diffusion model

In section 3 we demonstrated that the slow diffusion of trapped gas bubbles causes pumice to eventually sink. Here we plot solutions to the diffusion equation to examine how pumice floatation time varies as a function of pumice size, porosity, and initial trapped gas saturation. To write a solution to the diffusion equation we make several assumptions: (1) pumice is spherical; (2) the binary water + <u>gas mixture</u> in pumice can be described as a continuum, (3) initial trapped gas saturation is uniform within the pumice, (4) the <u>diffusion coefficient</u> is constant, (5) pumice pores are entirely connected, and (6) the pumice is entirely submersed in water. With these assumptions in place, the average saturation ξ (gas volume/pore volume) in a spherical pumice of diameter *L*, with initial interior gas saturation ξ i, and with gas saturation equal to zero at the boundaries ±L/2 is

$(14)\xi(t)=6\xi i\pi 2\sum n=1\infty 1n2e-Deffn2\pi 2t/(L/2)2$

(<u>Carslaw and Jaeger, 1959</u>, <u>Manville et al., 1998</u>). Here we let Deff=D θ i2=D(1- ξ i)2(e.g., Equation (<u>6</u>)). Pumice sinks when its average density is equal to the density of the surrounding liquid such that

 $(15)*(1-\phi)\rho r+\phi(1-\xi*)\rho l=\rho l,$

where ρr is the density of the solid phase, $*\xi*$ is the gas saturation when a pumice is neutrally buoyant and ρl is the liquid density.

By combining Equations (14) and (15) we can calculate pumice floatation time, the time it takes a pumice to reach gas saturation $*\xi*$, as a function of pumice size *L*, porosity ϕ , and initial gas saturation ξ i. Fig. 7 shows solutions to Equations (14) and (15) using contours of constant floatation time. Fig. 7 explores how pumice size, porosity, and initial gas saturation affect pumice floatation. We find that high porosity pumice need higher initial gas saturations to float for the same period of time as lower porosity pumice. Pumice size and porosity vary inversely along contours of constant floatation time and, for constant ϕ and ξ i, larger pumice float longer.



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Fig. 7. Floatation time as a function of <u>clast</u> size, <u>porosity</u> and initial saturation. We plot solutions to Equations (<u>14</u>) and (<u>15</u>) to demonstrate how floatation time varies with <u>pumice</u> size, porosity, and initial gas saturation, ξ_i . We assume $\rho_i = 2.4$ g cm⁻³, $\rho_i = 1.0$ g cm⁻³, and sum the first ten terms of Equation (<u>14</u>).

5. Conclusions

In this paper, we explore the processes that allow <u>pumice</u> to float and sink. To do this, we studied pumice saturation at the scale of <u>pore</u> level processes where capillary forces may be relevant. From estimates of the capillary number – which indicate that <u>surface</u>

tensiondominates over viscous forces in pumice – we hypothesized that pumice can trap isolated gas bubbles as water infiltrates its pores. We used X-ray microtomography to determine if gas trapping occurs in pumice and found that both <u>ambient</u> temperature and hot (500 °C) pumice trap gas. We observed, however, that hot pumice traps far less gas than ambient temperature pumice, consistent with the observation that hot pumice (>300 °C) often sinks after <u>contact</u> with water (Whitham and Sparks, <u>1986, Allen et al., 2008, Jutzeler et al., 2016</u>). That is, cold pumice can trap enough gas to keep the pumice afloat while gas contraction in hot pumice leads to lower <u>residual</u> <u>gas</u> saturations.

We examined the size distributions of trapped gas in both hot and ambient temperature pumice and found that the distributions fit a power-law prediction

from <u>percolation</u> theory. The power-law fits support a percolation theory treatment of water infiltration into pumice.

While trapped gas can buoy pumice, we hypothesized that outward diffusion of the trapped gas clusters causes pumice to eventually sink. We quantified this hypothesis with a model for pumice floatation time in terms of trapped gas diffusion (Equation (6)). To test the gas diffusion hypothesis, we conducted pumice floatation experiments and compared Equation (6) to our results as well as data from four other studies. We found that the observed pumice floatation times match a trapped gas diffusion prediction (Fig. 6). Furthermore, we plot solutions to the diffusion equation on a sphere to explore how pumice porosity, size, and initial gas saturation affect floatation time (Fig. 7). A mechanistic explanation for pumice floatation is an important step towards understanding when and how pumice is partitioned into rafts versus submarine deposits (e.g., Cashman and Fiske, 1991, Allen and McPhie, 2009; Cas and Giordano, 2014). Indeed, submarine volcanic deposits may not record eruption dynamics because noncondensable (i.e., CO₂ or air) gas trapping can make pumice buoyant. Furthermore, our results suggest that the lifetime of buoyant pumice rafts is not just limited by pumice abrasion (e.g., Carey et al., 2001, White et al., 2001), but also by gas diffusion. Noncondensable gas trapping and diffusion, by controlling pumice buoyancy, are therefore important controls on the transport and fate of pumice in the marine environment and location in the rock record.

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Appendix A. Supplementary material

The following is the Supplementary material related to this article.

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Supplementary information. S.1 Avizo marker-based watershed vesicle separation methods.

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