Please cite as: Jones, T.J., McNamara, K., Evchenne, J., Rust, A.C., Cashman, K.V., Scheu, B. and Edwards, R., 2016. Primary and secondary fragmentation of crystal-bearing intermediate magma. Journal of Volcanology and Geothermal Research, 327, pp.70-83. Typeset manuscript available here: https://doi.org/10.1016/j.jvolgeores.2016.06.022 Primary and secondary fragmentation of crystal-bearing intermediate magma Thomas J. Jones^{1,2*}, Keri McNamara², Julia Eychenne², Alison C. Rust², Katharine V. Cashman², Bettina Scheu³ and Robyn Edwards² [1] Department of Earth Sciences, Durham University, Durham, DH1 3LE, UK [2] School of Earth Sciences, University of Bristol, Wills Memorial Building, Bristol, BS8 1RJ, UK [3] Department of Earth and Environmental Sciences, LMU Munich, Theresienstr. 41, 80333 Munich, Germany

31 Abstract

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Crystal-rich intermediate magmas are subjected to both primary and secondary fragmentation processes, each of which may produce texturally distinct tephra. Of particular interest for volcanic hazards is the extent to which each process contributes ash to volcanic plumes. One way to address this question is by fragmenting pyroclasts under controlled conditions. We fragmented pumice samples from Soufriere Hills Volcano (SHV), Montserrat, by three methods: rapid decompression in a shock tube-like apparatus, impact by a falling piston, and milling in a ball mill. Grain size distributions of the products reveal that all three mechanisms produce fractal breakage patterns, and that the fractal dimension increases from a minimum of ~2.1 for decompression fragmentation (primary fragmentation) to a maximum of ~ 2.7 by repeated impact (secondary fragmentation). To assess the details of the fragmentation process, we quantified the shape, texture and components of constituent ash particles. Ash shape analysis shows that the axial ratio increases during milling and that particle convexity increases with repeated impacts. We also quantify the extent to which the matrix is separated from the crystals, which shows that secondary processes efficiently remove adhering matrix from crystals, particularly during milling (abrasion). Furthermore, measurements of crystal size distributions before (using x-ray computed tomography) and after (by componentry of individual grain size classes) decompression-driven fragmentation show not only that crystals influence particular size fractions across the total grain size distribution, but also that free crystals are smaller in the fragmented material than in the original pumice clast. Taken together, our results confirm previous work showing both the control of initial texture on the primary fragmentation process and the contributions of secondary processes to ash formation. Critically, however, our extension of previous analyses to characterization of shape, texture and componentry provides new analytical tools that can be used to assess contributions of secondary processes to ash deposits of uncertain or mixed origin. We illustrate this application with examples from SHV deposits. Keywords: Volcanic ash; Fragmentation; Broken crystals; Milling; Fractals; X-ray computed tomography

1. Introduction

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Volcanic ash is an inevitable product of nearly all explosive eruptions. Formed by fragmenting magma and/or rock, it is a particularly important hazard in the modern day, presenting a risk to aviation as well as to human health. Additionally, the associated risk is not confined to areas proximal to the volcano, but, as demonstrated by recent eruptions in Iceland and Chile, can also have far reaching impacts (e.g., Alfano et al., 2011). Understanding the origin of volcanic ash particles is thus critical for predicting the nature and extent of ash hazards. Volcanic ash can form by both primary and secondary processes; the former from volatile-driven decompression during magmatic ascent and the latter from post-fragmentation processes such as collisions within the conduit and transport in pyroclastic density currents (PDCs). Distinguishing between the products of primary and secondary fragmentation is necessary for a comprehensive understanding of volcanic ash deposits. Primary magmatic fragmentation occurs either through ascent-driven vesiculation and expansion of a volatile phase or by rapid decompression, such as occurs because of edifice collapse. Typical fragmentation studies relate tephra characteristics (e.g., grain size distributions, GSDs) to the energy of the primary fragmentation (e.g. Walker, 1973). Secondary fragmentation further decreases the average grain size of a pyroclastic deposit. In PDCs, secondary ash-forming processes include both impact and abrasion (e.g. Freundt and Schmincke, 1992), which cause fining of vent-derived particles with increased transport distance (Dufek and Manga, 2008; Kueppers et al., 2012). The total grain size distribution (TGSD) produced by primary and secondary fragmentation processes, in turn, affects the efficiency of heat transfer (e.g., Zimanowski et al., 2003), PDC mobility (Félix and Thomas, 2004) and formation of co-PDC plumes (Eychenne et al., 2012). For this reason, identifying the contributions of co-PDC ash to the total deposit is particularly important for modelling ash plumes, especially in large eruptions where such deposits may make up a large proportion of distal ash (e.g. Dartevelle et al., 2002; Cashman and Rust, 2016; Engwell and Eychenne, 2016; Engwell et al., 2014; Eychenne et al., 2015; Eychenne et al., 2012; Rose and Durant, 2009). The goal of our work is to

develop analytical tools to distinguish primary from secondary ash deposits.

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Controlled laboratory experiments have provided some links between natural variables (e.g. degree of overpressure, transport distance) and the characteristics of the associated tephra deposit. Laboratory study of primary fragmentation is typically performed by shock-tube experimentation at magmatic temperatures. Results show that for a given sample porosity, greater degrees of overpressure increase the fragmentation efficiency and thus decrease the overall grain size (Kueppers et al., 2006b). Experimental GSDs typically follow power-law distributions with fractal dimensions (D) of ~2.5 (Kueppers et al., 2006a). Variations about this value are thought to reflect the energy of fragmentation, with higher fragmentation energies producing deposits with higher D values (Perugini and Kueppers, 2012). The fractal dimension increases by secondary fragmentation (Kaminski and Jaupart, 1998). The effects of secondary fragmentation have been quantified using experiments involving both collision and abrasion (Cagnoli and Manga, 2004; Mueller et al., 2015). Results confirm that the fine ash content increases with time, and that experiments involving abrasion produce finer ash than experiments involving impact (Mueller et al., 2015). Also important is the original clast density, with grain size reduction dominated by breakage of the more vesicular fragments (Kueppers et al., 2012). This observation has particular relevance to explosively generated ash, which often has TGSDs dictated by the vesicle size distribution (Genareau et al., 2012; Liu et al., 2015a; Rust and Cashman, 2011).

Less well studied is the role of crystals in fragmentation and ash formation. Crystals commonly form a major constituent of intermediate magmas; thus it seems likely that the abundance and size distributions of crystals would affect the size, density and shape of explosively generated fragments. In fact, crystal concentrations in ash deposits show that individual crystals typically have a limited size and density range (e.g., Martel et al., 2001; Cashman and Rust, 2016; Sparks and Walker, 1977). Crystals in ash fractions are often broken or rounded, possibly due to the fragmentation and/or ash generation mechanism (e.g. Bachmann et al., 2002; Best and Christiansen, 1997; Carter et al., 1986). The presence of broken crystals in explosive eruption deposits has previously been attributed to two main processes: melt inclusion (MI) decrepitation and shock fragmentation. Fragmentation by MI decrepitation occurs when overpressure in the MI exceeds the tensile strength of the crystal. This may

be achieved by cyclic periods of overheating linked to the latent heat of crystallisation (Bindeman, 2005; Zhang, 1998) or rapid decompression of crystal-bearing magma during ascent (Best and Christiansen, 1997; Bindeman, 2005; Miwa and Geshi, 2012; Tait, 1992; Williamson et al., 2010). Crystal fragmentation by shock fragmentation can occur by extensional fracture during decompression and subsequent melt vesiculation (Chouet et al., 1994; Kennedy et al., 2005; Miwa and Geshi, 2012; Pallister et al., 1996).

Here we present a multi-component analysis of experimentally fragmented pumice samples. Although the effects of both rapid decompression and abrasion have been previously investigated, we extend these experiments in several important ways. First, we focus on fragmentation of a vesicle- and crystal-rich intermediate magma by a range of methods. Second, we explore the effects of both primary and secondary fragmentation on the resulting shape, size and componentry of the particle population. Primary fragmentation is simulated by rapid decompression experiments in a shock tube while secondary processes (such as abrasion and impact in PDCs) are simulated in the laboratory by ball mill and falling piston experiments, respectively. Controlled experiments with the same starting material – pumice from recent eruptions of the Soufriere Hills Volcano (SHV) – allow us to compare the ash-generating efficiency of the different processes. Additionally, adding componentry analysis allows us to evaluate the extent to which (broken) crystals affect the grain size, shape and fractal dimension of the tephra deposit. Finally, we compare our experimental results with natural SHV ash samples from fallout deposits produced by a dome collapse and a Vulcanian eruption.

2. Methods

All experiments were conducted using a single crystalline andesite pumice block that survived decompression during an explosive (Vulcanian) eruption of SHV in February 2010. We fragmented the pumice sub-samples in three different ways: rapid decompression at both room temperature and 880°C, impact and milling at room temperature - all within the brittle deformation field; (Alidibirov and Dingwell, 2000). To characterize the resulting material, we determined the total grain size distributions produced by each fragmentation method, as well as ash components, shapes and textures.

For the decompression experiments, we also compare crystal sizes and shapes before and after fragmentation.

2.1 Pre-fragmentation characterisation

The original sample had a dense-rock equivalent density of 2.69×10^3 kg m⁻³ and an estimated glass transition temperature (T_g) of ~790°C (Jones et al., 2013). For the decompression experiments we drilled four pumice cylinders, each measuring 60 mm long and ~25 mm in diameter. Prior to fragmentation the physical and textural properties were characterised. Crystal populations and preexisting crystal fractures were quantified with X-Ray computed tomography (XRCT) and optical microscopy of thin sections; helium pycnometry was used to calculate connected and total porosities for the four samples (see Table 1).

2.1.1 X-Ray computed Tomography

Radiographs for XRCT were collected with a Nikon Meterology 225/320kV Custom Bay scanner located at the Henry Mosely X-Ray Imaging Facility, University of Manchester. Acquisition conditions were 41 kV accelerating voltage, 239 μ A current and a 0.5 mm Al filter. An exposure time of 1000 ms was used for all scans. To reconstruct the scans, we first calculated the centre of rotation of all the radiographs and then applied a beam hardening correction using the Xtek CT Pro software.

Mineral modes, porosity and bulk sample crystallinity were determined with a larger sub-volume of $6x10^8$ voxels, which required significantly more computational capacity than the crystal volume measurements, and therefore a different approach. Firstly a median filter was applied to the entire image stack with ImageJ software (http://rsbweb.nih.gov/ij/). Next pores and mafic and felsic mineral phases in each image were separated based on grey scale values with an ImageJ plugin, PhaseQuant (Elangovan et al., 2012). The small density variation between different mineral phases, and the corresponding similarity of grey scales, made segmentation of specific minerals difficult so crystals are classified as either mafic or felsic phases (e.g. Cnudde et al., 2006). Each segmented volume (felsic, mafic and pores) was then converted into a binary image stack and imported into Avizo. The felsic crystal data required a 3D erosion and subsequent dilation by two voxels to remove background noise. Finally, surface generation and volume analysis was performed.

2.1.2 Textural characterisation

A 3D rendered volume of an example sub-sampled volume (Figure 1a) shows the distribution of crystals prior to experimental fragmentation. Broken crystals are common, and often have a jigsaw-like fit. Where the separation between these crystal fragments was sufficient to resolve in the tomographic analysis, each crystal fragment was counted as a separate crystal. The average DRE crystallinity for crystals $>32\mu m$ in diameter (5 ϕ) is 27 vol. % with \sim 21% felsic and \sim 6% mafic crystals and oxides. Point counts of petrographic thin sections of the same samples included smaller microlites, and thus yield a higher crystallinity (38%); this shows that about 11% of the total crystal volume is < 32 μm .

A crystal volume distribution (CVD) extracted from the rendered tomographic volumes is shown in Figure 1b. The crystals range in size from 10⁻⁵ to 10¹ mm³, with a mode of 10⁻³ mm³. Measurement of broken crystals as individual fragments means that the rendered volume distribution accounts for pre-existing crystal breakage prior to experimental fragmentation, within the resolution of the imaging technique.

2.3 Porosities and fragmentation threshold

Porosity provides information about the potential energy available for fragmentation during decompression (Kueppers et al., 2006b) and is used to calculate the minimum initial overpressure required for complete fragmentation in the decompression experiments (Spieler et al., 2004). To measure porosity, we cleaned the sample cores and measured their porosity by helium pycnometry, using a Quantachrome Instruments Ultrapync 1200e pycnometer housed at LMU Munich, Germany. Total porosity \varnothing_T was calculated from the solid density ρ_s and the bulk sample density ρ as:

 $\emptyset_T = 1 - \frac{\rho}{\rho_s}, \tag{1}$

where ρ_s is the measured density of powdered sample milled to a grain size less than the minimum pore size. The connected or open porosity \varnothing_C is calculated as:

$$\varnothing_C = \left(\frac{V_C - V_M}{V_C}\right) \times 100 , \quad [2]$$

where V_C is the geometrical volume of the cylindrical samples and V_M is the volume measured by He pycnometry. The isolated (closed) porosity, \varnothing_I can then be defined as:

$$\emptyset_I = \emptyset_T - \emptyset_C.$$
 [3]

The average total porosity is 66%, with a range of 1.25% between all sample cores. Individual porosity values for the cores used in the four rapid decompression experiments are reported in Table 1.

2.2 Experimental fragmentation

Samples from the Soufriere Hills Volcano (SHV) were fragmented experimentally by three methods: rapid decompression, hammer impacts and ball milling (Figure 2). Broadly, these aimed to replicate three natural volcanic processes: magmatic overpressure and decompression, impact and abrasion/milling in turbulent flow.

Rapid decompression-driven fragmentation was performed using a shock tube-like apparatus (Figure 2a) at the LMU Munich, Germany (e.g. Scheu et al., 2008; Spieler et al., 2004). By using two

experimentally calibrated diaphragms we achieved a pressure differential of either 15 or 30 MPa; this exceeds the fragmentation threshold by 12 and 27 MPa (Spieler et al., 2004). At each differential pressure, experimental fragmentation was conducted at both 20°C and 880°C. The experimentally generated pyroclasts were left to settle in the low pressure tank for 1-2 hours and then sieved to 125 μ m using a pressure washer filled with distilled water. All material that passed through the 125 μ m sieve was left to settle for 2 days in a sediment collection tank.

Impact experiments were performed by placing a single rectangular block (~ 4 x 5 x 2cm) of SHV pumice into a detachable steel cup, as illustrated in Figure 2b. A 10 kg steel piston was then repeatedly dropped from a height of 50 cm. The fives experimental runs, each run with a different block, comprised 5, 10, 15, 20 and 25 piston drops; at the end of each run the entire sample was recovered from the basal cup by flushing with distilled water.

Ball mill grinding was performed using a Planetary Ball Mill PM 100 manufactured by RETSCH Ltd. The initial GSD for the ball mill experiments, represented by "0 min" in Figure 3c, was created by crushing ~ 10cm square blocks of SHV pumice. For each experiment an aliquot of this input sample was weighed and placed into the agate jar of the ball mill (Figure 2c) along with six 1.6 cm diameter agate balls. The container was sealed and rotated at 450 rpm for 0.5, 1, 2.5 or 5 minutes, and the sample removed for subsequent analysis.

2.3 Post-fragmentation characterisation

After fragmentation by the three methods (Figure 2) the experimental pyroclasts were characterised by grain size, componentry, shape and fracture surface morphology.

2.3.1 GSD measurements

All particles coarser than 125 μ m were separated into grain size fractions by manual sieving into size bins of -3, -2, -1, 0, 1, 2, 3 phi (ϕ). Particle size analysis for material \leq 250 μ m was performed at the Environmental Change Laboratory, University of Western England, Bristol, UK using a Mastersizer 2000 laser diffraction particle size analyser, manufactured by Malvern

Instruments Ltd. Sieve and Mastersizer data were combined using the overlapping 3 φ size fraction to produce a total grain size distribution for each fragmentation experiment.

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2.3.2 Componentry

Componentry analysis was conducted under a binocular microscope for grain sizes -3 to 3 φ. Grains ≥ 500 µm were picked manually; smaller grains were classified from analysis of digital microscope images. Where possible, at least 300 grains were analysed for each grain size fraction. Grains were separated into three component categories: (1) free mafic crystals, (2) free felsic crystals, and (3) clasts (i.e. anything that is not a free crystal), where a free crystal is defined as having matrix adhered to less than 20% of the surface of the crystal. Additional 2D componentry of the 2 φ ash fraction was performed using backscattered electron (BSE) mode of Scanning Electron Microscopy (SEM) on a Hitachi S-3500N SEM at the School of Earth Sciences, University of Bristol. This grain size fraction was chosen for detailed analysis because all experiments produce sufficient samples of this size fraction. Ash grains were set in resin and polished to expose grain interiors before being carbon coated. A mosaic of 25 images was taken of each sample using BSE mode from a working distance of 18 mm, a 15Kv accelerated voltage and a magnification of 60x. Each image had a 1024 x 768 pixel resolution. Componentry was performed manually by placing a grid on the image and counting grains. Grains were subdivided into three components: (1) crystals with no attached matrix, (2) crystals with matrix attached, and (3) vesicular matrix clasts (all matrix grains were vesicular). The 3D morphology of fine ash samples was examined using secondary electron mode on the

SEM. Ash grains within the 2 ϕ sieve fraction were mounted on carbon-based stubs and Au coated to

a thickness of approximately 5 nm. For grains measuring 125-250 μm or 3 φ (the finest fraction

studied), silver paint was applied to the SEM stub prior to Au coating, to reduce the effects of

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3.2.3 Shape analysis

charging.

The 2D SEM image mosaics of the 2 ϕ grain size fraction were analysed with ImageJ software. Images of individual grains were thresholded and converted to binary format before

quantifying grain shapes using axial ratio and convexity. Only grains from the 'vesicular matrix' componentry category were analysed for shape as they displayed the most variation between sample runs. This component comprised between 10 and 60 % of the total depending on the fragmentation mechanism.

Axial ratio is the ratio of the axes of the particle's best-fit ellipse:

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$$Axial\ Ratio\ (AR) = \frac{Minor\ Axis}{Maior\ Axis}. \quad [4]$$

Convexity is defined as the ratio of the perimeter of the grain and its convex hull (the smallest convex polygon that contains the 2D shape):

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$$Convexity = \frac{Perimeter\ of\ convex\ hull}{Perimeter}.$$
 [5]

- It is a measure of the surface roughness of the external shape boundary (Liu et al., 2015b), such that a high value of convexity indicates a smooth external surface.
- Where a crystal comprised at least 10% of the entire grain, areas of crystals and matrix were also 289 290 calculated separately by manually varying the greyscale threshold to generate separate binary images, then calculating the pixel area of each. Abrasion is expected to decrease the relative proportion of 291 glassy matrix adhering to crystals. For this reason, Meyer (1971) define the "abrasion index", which is 292 the ratio of the area of a crystal to the area of attached matrix. However, abrasion may not be the only 293 294 process to affect the crystal-matrix ratio. Furthermore, Meyer's "abrasion index" is unbounded (i.e. 295 becomes infinite if there is no matrix). For these reasons, we introduce a related parameter we call the 296 Adherence Factor:

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$$Adherence\ Factor\ (AF) = \frac{area\ of\ matrix}{area\ of\ matrix + area\ of\ crystal}\ [6]$$

The adherence factor is bounded between one (matrix only with no dominant crystal) and zero (for a crystal with no adhered matrix) and therefore we expect AF to decrease with increasing abrasion.

4. Results

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4.1 Grain size distributions

The grain size distributions (GSD) of all sample sets are limited to a maximum of -3φ because of initial sample size. Rapid decompression (Figure 3a) creates a sample suite with a median size of -2.30φ to 0.23φ. Most striking is the effect of changing fragmentation temperature, where elevated temperatures yield coarser GSDs. The grain size distributions produced from the falling piston experiments are notably more fine-grained than pyroclasts produced by rapid decompression. Median sizes range from 1.70φ to 5.03φ and generally decrease with increased number of hits (Figure 3b). The grain size resulting from 10 hits, however, is abnormally small relative to the other data sets; we attribute this to natural heterogeneity within the pumice block that was the source of the starting material of each experiment. Experimentally generated pyroclasts from ball milling show a systematic increase in fine particles with increased milling duration. All GSDs produced by milling show a pronounced fine tail and are skewed to smaller sizes than GSDs from the rapid decompression and falling piston experiments.

4.2 Ash shape

Vesicular grains (i.e. all those that were not dominated by a single large crystal of over 10% of the total grain) were analysed for two shape parameters: axial ratio and convexity. Axial ratio is a representation of particle elongation, while convexity quantifies the smoothness of the grain exterior (Liu et al., 2015b). Axial ratio values vary considerably depending on the experimental fragmentation method (Figure 4a-c). Ash produced by rapid decompression has a greater range of axial ratio values than those produced by milling or impact. Ash grains from the milling experiments showed the smallest range of axial ratios although the range increased with milling time: axial ratios of grains milled for 1 minute have a lower mode and extend to larger values than axial ratios of grains milled for 0.5 minutes (Figure 4c). In contrast, axial ratio values changed little with number of impacts (Figure 4b). However, convexity data show that increased impacts smoothed out the irregular grain surfaces in the comparative grain size fraction: the ash grains produced by 10 piston impacts have greater average convexity values, and thus less irregular exteriors, than grains produced by 5 impacts (Figure 4e).

4.3 Experimentally generated pyroclast componentry

Componentry analysis of the pyroclasts generated by rapid-decompression required the proportions of free mafic and felsic crystals (those with < 20% adhered matrix) to be discriminated from other matrix dominated clasts (Figure 5a). No free crystals were observed at grain sizes greater than -1 φ . At finer grain sizes the proportion of crystals varies considerably for different grain size fractions. Within these coarse-grained rapid decompression products, free crystals (mainly felsic) are most abundant in the 1φ size fraction, contributing ~40 weight % to this grain size bin. The maximum abundance of mafic crystals varies between 1 and 2 φ for different experimental runs. We observed no systematic componentry changes in the decompression experiments (Fig. 5a) as a function of either ΔP or fragmentation temperature; the small variations between experimental runs likely relate to slight sample heterogeneities prior to experimental fragmentation.

Within the 2φ fraction (Fig. 5b), there are only small differences in SEM componentry between the hot and cold rapid decompression products. Both have similar proportion of free crystals (with no attached matrix) although there is a higher proportion of crystals with attached matrix in the products of hot decompression (Figure 5b). More dramatic is the comparison with data from other fragmentation mechanisms, where products of the milling and impact experiments show a higher proportion of free crystals than the decompression products. Additionally, the sample milled for longer (1 min) has a higher proportion of free crystals than its counterpart that was milled for 30 seconds. A similar trend is observed in the impact products: the sample that underwent 10 hits has a greater proportion of free crystals than the sample that was hit five times. These observations show that repeated fragmentation removes crystal coatings and frees individual crystals from the matrix.

4.4 Crystal volume distributions

The crystal populations within the original SHV pumice cores and the pyroclasts produced by rapid decompression were quantified by analysis of X-Ray CT (Figure 1) and SEM analysis (Figure 5a), respectively. To effectively compare these pre-experiment and post-experiment crystal populations,

the crystal volume distributions generated by X-Ray-CT were converted to a length scale. We assume that all crystals within the volcanic rock samples form euhedral cuboid volumes,

$$V_{xal} = h \times w \times l$$
 [7]

where h and w represent the height and width of the smallest crystal face and l is length. Assuming that crystals have aspect ratios of 2:1 and that the minor and intermediate axes are the restrictive dimensions of the sieve size (d), an effective particle diameter is calculated as:

$$d = \left(\frac{V_{xal}}{2}\right)^{1/3} \quad [8]$$

Effective diameters are converted to the ϕ scale, the volumes falling in each ϕ bin summed and converted to weight % to form the "Rock" data set in Figure 6.

The crystal population generated by decompression is determined by averaging the mass of mafic/felsic crystals within each size fraction of all decompression experiments. We average because no significant difference in componentry was observed when changing fragmentation overpressure or temperature. These data are then converted to weight % of total crystals and form the "Ash" data set in Figure 6.

The size distribution of crystals within the ash relative to those in the SHV pumice (Figure 6) shows that rapid decompression caused the crystal size to decrease. Grain size reduction is concentrated over the central portion of the size range investigated. In coarser grain size fractions $(2 - 0 \, \phi)$ the crystal abundance in the pumice is greater than the abundance in the ash and is dominated by felsic crystals. Conversely, in the finer fractions $(1 - 3 \, \phi)$ crystals are more abundant in the ash than in the original pumice. Moreover, there are no crystals within the -2 ϕ ash fraction, although they were present in the pumice prior to decompression and fragmentation. This suggests that crystals were not simply freed from the matrix during fragmentation but were also reduced in size by crystal breakage.

4.5 SEM imagery of fracture surfaces

Photomicrographs of the 2φ ash fraction produced by rapid decompression reveal characteristics of broken crystal surfaces (Figure 7) that can be classified as: (1) smooth, clean surfaces with negligible topographic relief (Figure 7a) and (2) rougher, often highly irregular surfaces displaying intense riverline fracturing and hackles (Figures 7c and 7d). On some crystals, a vesicular glass coating is observed (Figure 7b); it is slightly more common in the hot fragmentation experiments relative to the room temperature runs. No other differences were observed in the fracture surfaces or styles between the products of hot and cold experiments. About 90% of broken crystals have clean broken surfaces with negligible topographic relief. We interpret these as breakage along a cleavage plane or a preexisting internal fracture (Figure 7a). Where river-line fractures are observed on these surfaces they are widely spaced and have low relief. Only about 10% of broken crystal surfaces are much rougher and complex due to fractures cross-cutting cleavage planes (e.g. Figure 7d).

Finally, sub-circular cavities, including some with protruding glass strands, are a rare but ubiquitous feature of crystal surfaces in the experimental products. Similar features were observed by Williamson et al. (2010) in natural SHV pumice and interpreted as melt inclusions that burst within a plastic groundmass due to a major decompression event (e.g. dome collapse). Melt extension is a non-brittle process that could not have been generated through experimental fragmentation at room temperature. The fact that we observe these features in the experiment products indicates that they must already have existed in the dome/pumice samples prior to decompression in the laboratory.

5. Discussion

We have explored several techniques to characterise both experimentally generated ash and its parent material (Table 1). We now use these results - from grain size measurements, fractal analysis, componentry and shape descriptors - to examine the effects of different fragmentation mechanisms on pyroclast characteristics. For samples that were rapidly decompressed, we couple this analysis with XRCT and SE SEM imagery to examine how rapid decompression-driven fragmentation alters the crystal population. Lastly, to illustrate our techniques we compare experimentally generated SHV ash to two natural SHV ash examples.

5.1 Fractal behaviour of products

Since the idea of fractal behaviour was introduced to Earth Sciences (e.g., Korvin, 1992; Turcotte, 1986), power law exponents (fractal dimension D values) have been used to quantify the size distributions of volcanic pyroclastic products (e.g., Kaminski and Jaupart, 1998; Perugini et al., 2011; Taddeucci et al., 2004). Higher power law exponents represent tephra deposits that are dominated by finer ash fractions. Therefore fractal dimensions are commonly used to infer the fragmentation efficiency and have been linked to the energy available for fragmentation (e.g., Kueppers et al., 2006a).

It has long been noted that tephra produced from volcanic eruptions commonly has a much higher fractal dimension (larger fine ash component) than expected from simple crushing or rock disaggregation (Hartmann, 1969). The source of this fine ash is an open question, but is clearly dictated ultimately by the vesicle size distribution (Rust and Cashman, 2011). Kaminski and Jaupart (1998) proposed a model of secondary fragmentation based on experimental fragmentation, where they assumed that piston impact causes primary fragmentation while grinding in a ball-mill caused secondary fragmentation. In this scenario, primary fragmentation initially creates $D=2.5\pm0.1$, then ongoing selective re-fragmentation through particle-particle collisions increases the values of $D\geq3$ preserved in fall deposits. However, many experimental fragmentation experiments fail to replicate the high power law exponents observed in natural pyroclastic deposits (Table 2). This leads to a question about the mechanism(s) capable of producing the finer ash, particularly the role of the original bubble population in controlling the final grain size distribution (Rust and Cashman, 2011).

We analysed the fractal dimension of the total GSD for all experimental fragmentation methods by converting raw data (mass proportion of particles in different size bins; Figure 3) to number-based data to aid comparison to other published data sets (e.g. Kueppers et al., 2006a). Over the size range analysed here, all experimentally produced ash samples follow a power law distribution, that is, they plot on a straight line in log(N)–log(L) space (Figure 8), where N represents the number of grains

larger than corresponding fragment size (L) and the slope defines the fractal dimension (D). The power-law distributions were fitted to the total GSD recovered from the experiments. In some cases one or two data points are excluded at the coarse or fine tail (open circles in Figure S1).

The log(N) vs. log(L) data for the rapid decompression experiments (Figure 8a) form linear trends with fractal dimensions between 2.03 and 2.24, in good agreement with other studies that have used the same fragmentation method (Kueppers et al., 2006a; Kueppers et al., 2006b; Perugini and Kueppers, 2012). Pyroclasts from the impact (falling piston) experiments also show a fractal distribution (Figure 8b) with a mean D = 2.60. This value is comparable to values from simple crushing and disaggregation of rocks (Table 2; Hartman, 1969). The falling piston sample set shows the most variation. Here D values do not vary systematically with number of impacts, although the power law exponents do broadly increase with increased number of impacts. In these analyses, we excluded the tails of the fragment size distribution when calculating D (represented as open circles in Figure S1), which may account for minor disparities. The milling experiments follow a power law distribution, which shows a systematic increase in D from 2.33 at 0.5 min to 2.45 at 5 mins (Figure 8c). This progressive rise in D suggests that milling not only creates fine ash but also causes the fine ash component to become increasingly dominant in the TGSD.

Our experimental data show that secondary fragmentation can progressively increase D from an initial value resembling rapid decompression-driven fragmentation to higher values. Yet, even after considerable milling durations (5mins) or a large number (25) of successive impacts, the experimental products do not exceed D values of 3. Furthermore, analysis of the 2ϕ fraction shows that the crystals were not being broken through milling, rather they were being stripped of adhering groundmass. This suggests that during secondary fragmentation of crystal rich tephra, the crystal population may help to sustain a relatively coarse control on the GSD. This interpretation is supported by data from eruptions of Heimaey, Fuego and Oshima, where the basaltic pyroclasts are rich in microphenocrysts and the TGSDs have fractal dimensions D \sim 1.9-2.3. Silicic deposits (Mount St. Helens, El Chichon and Quizapu), in contrast, have both low (or no) groundmass crystallinity and D values >3 (Table 2).

Assuming all of the eruptions listed in Table 2 involved secondary, as well as primary, fragmentation (through both milling and impacts, either with the conduit wall or within the particle-rich plume at depth), the differences in observed D seem most easily explained by variations in groundmass crystallinity. Where large proportions of crystals are present in the starting magma they provide a coarse control on the grain size distribution and prevent production of high proportions of fine ash; this keeps D values low (< 3). The 'final' D value is therefore a function of both magma and fragmentation characteristics, including magma porosity and permeability, magma crystallinity, fragmentation overpressure and the degree of mechanical processing.

Our analysis confirms results from previous experimental studies, including the effect of mechanical

fragmentation in shifting the GSD to smaller sizes (Cagnoli and Manga, 2004; Kueppers et al., 2012;

5.2 Fragmentation control on pyroclast characteristics

Mueller et al., 2015) and the high efficiency of abrasion relative to impact experiments in producing fine ash (Mueller et al., 2015; Figure 3). We also found that the proportion of vesicular fragments decreased with increased milling time and number of impacts (Figure 5b), consistent with Kueppers et al. (2012).

Our experiments also show that the mechanism of fragmentation has a significant effect on the characteristics of individual pyroclasts. Of particular note is the variation in the axial ratios of matrix fragments between the three fragmentation mechanisms studied (Figure 4). The axial ratios of impact and milling products are large compared to the decompression products, which suggests that increased mechanical abrasion prevents, or rapidly reduces, the likelihood of producing elongated fragments. Additional milling further increases the mean and range of axial ratio values of ash particles. This is interesting because the small ash particles mimic shape variations previously observed in larger pumice clasts, both in rock tumbling experiments and natural PDC deposits (e.g., Manga et al., 2011). Studies of pumice clast rounding further suggest that both rounding and ash production via comminution are most efficient proximal to volcanic vents, where PDCs are most energetic.

A sizeable portion of the experimental products includes crystals with adhered matrix (Fig. 5b). Past studies have used these fragments as a marker for the amount of abrasion (Freundt and Schmincke, 1992; Meyer, 1971). We quantified this effect using an 'Adherence Factor' (AF: Equation 6; Figure 9). Our data show that the decompression experiments have a substantially higher AF (more adhered matrix) than the products of impact and milling. Hot decompression causes particles to retain slightly more matrix material than cold decompression. Interestingly, increasing the number of impacts or time of milling does not appear to dramatically alter the AF, but the products of milling have a lower AF, overall, than the products of impact.

To interpret the mechanisms responsible for adhered matrix produced by different fragmentation methods, it is also important to consider the effect of crystal breakage on AF. We identified broken crystals by recording whether the crystal displayed at least one intact ring of zoning or whether the zoning was interrupted. In the products of rapid decompression, the crystals with a lower AF value were typically broken. However, there is no obvious correlation between crystal breakage and AF in the products of milling and impacts (Figure S2); this lack of correlation may be due to these two mechanisms not dramatically increasing the number of broken crystals. As a consequence, the matrix is stripped from the outside without breaking the crystal, resulting in many whole crystals with a low AF value. In contrast, the low AF of broken crystals within products of rapid decompression suggests that, as crystals break, internal crystal surfaces are exposed and the area of adhered matrix decreases. Therefore we suggest crystal breakage is the main mechanism for reducing AF during rapid decompression.

5.3 The influence of crystals on fragmentation during rapid decompression

Crystals commonly form a major constituent of erupting magmas and there is the potential for the abundance and size distributions of crystals to affect the size, density and shape of fragments generated in explosive eruptions. The componentry analysis presented here demonstrates that free crystals contribute a significant mass (up to 40%) to specific grain size fractions produced by rapid decompression (Figure 5a). The size range in which crystals dominate appears independent of

fragmentation pressure and temperature and is directly related to the crystal size distribution in the starting material. Over the grain size range studied (-3 to 3 φ), plagioclase dominates the total crystal population within each size fraction of the experimentally generated fragments. This agrees with modal analysis of the starting material that showed it to be the most abundant mineral. As illustrated in Figure 5a, free crystals are non-existent or of trivial abundance in the grain size fractions coarser than 0 φ , consistent with the initial crystal size population in the starting material (Rock curve in Fig.6). The mafic crystals show a broader peak at smaller grain sizes, which is likely related to the initial size populations. Therefore the textural characteristics of the starting volcanic material (magma analogue) directly control the experimentally produced tephra (PDC analogue).

High proportions of crystals within a grain size fraction have the potential to alter the bulk ash density, especially at coarser grain sizes. This, in turn, will cause differential settling of crystals relative to glass fragments of similar size, particularly if the glass is vesicular (e.g., Sparks and Walker, 1977). This effect has been documented in natural eruptive products of Mount St Helens and Quizapu (Cashman and Rust, 2016), where the proportion of free crystals decreases faster with distance from the vent than pumice or glass shards from the same eruption. Currently, ash dispersion models (e.g. Tephra 2; Bonadonna et al., 2010) commonly use a single vesicular glass ash density to represent all grain size fractions from a volcanic eruption. Improvements could therefore be made to more accurately represent crystal-rich pyroclasts, particularly in eruptions characterised by PDCs.

Crystal size distributions may also help to explain the production of crystal pyroclasts devoid of glass coatings. If a whole (micro)phenocryst were to be extracted from the fragmenting magma, the surface would likely be at least partially coated in a glass. However, if internal broken fragments of a once larger phenocryst are extracted, then the relative chance of a glass-coated surface is low. Knowing the surface properties of ash has implications for remote sensing applications and surface leachate studies. Our componentry study suggests that free crystals in the ash generated by rapid decompression represent broken fragments from once larger whole crystals; this is likely because during rapid

decompression fragmentation, the crystals cannot accommodate deformation and hence fracture (Cordonnier et al., 2009).

Crystal fracture topographies indicate that breaking of crystals in our decompression experiments was dominantly along cleavage surfaces (type 1 fractures) with only about 10% of fractures cross-cutting cleavage planes (Figure 7). Crystals tend to break or 'cleave' along a particular crystallographic orientation because cleavage planes have a relatively low surface energy (Hull, 1999; Kelly and Macmillan, 1986). Experimental studies and natural observations of plagioclase identify intracrystalline fractures controlled by crystallography and preferential fracture along cleavage planes (e.g. Borg and Heard, 1970; Brown and Macaudiére, 1984). Further, Kennedy et al. (2005) use fractures in SHV hornblende crystals to infer the orientation of tensile unloading and therefore the shape of the fragmentation front. Under mode I tensile failure, cleavage surfaces have a direct and primary effect on fragmentation behaviour. When cleavage planes are orientated perpendicular to the tensile force (parallel to the unloading wave) then simple cleavage fracture is expected to occur (e.g., Figure 7a). However, when cleavage planes are orientated obliquely to tensile forces, then mode II failure can aid fracture by the formation of shear couples (Figure 7c). Therefore, the orientation angle, defined as the angle between the cleavage plane and the tensile force vector, will influence the relative proportions of mode I and II failure.

5.4 Comparison with natural samples

To compare with our experimental results, we also studied two ash samples from fallout deposits produced by a dome collapse event (on 31 March 1997) and a Vulcanian eruption (on 26 September 1997) of Soufriere Hills Volcano. The dome collapse event generated PDCs but no plume at the vent, making the fallout deposit entirely co-PDC in origin (Bonadonna et al., 2002; Engwell and Eychenne, 2016). The Vulcanian eruption produced both an eruptive column at the vent and PDCs on the volcano's flank; the fallout deposit consequently includes ash from both vent-derived and co-PDC plumes (Bonadonna et al., 2002). These two natural events, with distinct eruption characteristics, allow us to demonstrate how our analysis techniques can be used on natural material to identify

secondary fragmentation. To compare with experimental samples, we analysed the componentry, AF (Figure 9) and proportion of broken crystals in the 2 φ grain size fraction (Figure S2) of each sample. The 2 φ fraction of both natural samples comprises approximately 30% dense material (crystals, phenocrysts and vesicle-free glass). The Vulcanian sample however, contains 15% crystals with adhered glass whereas the dome collapse sample contains approximately 20% (Figure 10). Hence the Vulcanian sample comprises approximately 55% vesicular fragments, while the ash produced by dome collapse contains less than 50%. Of this vesicular portion, the Vulcanian sample is comprised mostly of microlite-free vesicular glass fragments. In comparison, the dome collapse sample is comprised entirely of the microlite-rich vesicular glass. The higher proportion of vesicular microlitefree glass in the Vulcanian sample indicates that in addition to microlite-bearing magma stored at shallow levels (top of the conduit or dome), deep magma was erupted in the Vulcanian explosion. In contrast, the dome collapse event involved only material previously extruded in the dome, comprising microlite-rich glass of variable vesicularity. The two natural samples also differ in patterns of matrix adherence. The AF distribution of the Vulcanian sample is trimodal, with a major mode at AF ~0.2 (low adhering matrix) and two minor modes at AF ~0.6 and 0.8 (Figs. 9 and S2). In comparison, the dome collapse sample shows a uniform AF distribution (Figure 9) and a higher proportion of broken crystals at low AF values (Figure S2). Although this natural case study is illustrative only, and uses a single grain size fraction at a single location for each eruption, both the components and the AF distributions are distinctive. The differences in componentry (vesicle and microlite content) of the natural samples can be explained by the fact that they experienced different ascent and fragmentation histories and different transportation modes (PDCs vs. plumes); both of which should affect the mechanical properties. The Vulcanian sample includes both particles transported directly in a plume and particles that were first transported in PDCs before being entrained into co-PDC plumes. These two events can be observed, respectively, in Figure 9 as a group of poorly abraded particles (high AF values) and a set of highly abraded particles (low AF values). The dome collapse sample is dominated by microlite-rich particles that were entirely transported in PDCs. The uniform AF distribution suggests that abrasion during

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transport did not affect all the phenocrysts uniformly. Differences might reflect: (1) the robustness of particles due to their high microlite content and moderate vesicularity compared to the highly vesicular, microlite-free particles produced by the Vulcanian eruption; and/or (2) the coarser initial grain size distribution in the dome collapse block-and-ash flows compared to the explosively produced Vulcanian pumice flows, whereby larger initial particle sizes would require more abrasion to release phenocrysts from the matrix. This points to a limitation of our analysis, which is that we have not attempted to assess variations as a function of time/distance. At Tungurahua volcano, it has been suggested that dense crystal-rich PDCs become increasingly crystal rich with time/distance as the vesicular material is removed through abrasion (Douillet et al., 2013). This hypothesis could be tested by applying our techniques to an appropriate selection of samples from a range of locations.

6. Conclusions

Three fragmentation methods (rapid decompression, impact and milling) have been explored to constrain fragmentation associated with three different eruptive processes (Figure 11). Analysed GSDs are fractal, which means that they can be characterised by the fractal dimension D. Our data show that an initial GSD with D \approx 2.1 produced by rapid decompression can be altered by secondary fragmentation processes that generate fine particles. Importantly, however, D values produced by secondary processes in our experiments reach only \sim 2.7 (Fig. 8), which is far from D >3 observed in many silicic eruptions (e.g., Kaminski and Jaupart, 1998; Rust and Cashman, 2011). We suggest that the relatively low D values reflect the role of the groundmass crystal population, which prevents extensive crushing and grain size reduction.

We support this hypothesis by analysis of particle shape (Fig. 4), componentry (Fig. 5) and adherence factor (AF; Fig. 9). These data show that secondary fragmentation by either impact or milling dramatically reduces the matrix component (Fig. 5b); ash generated by milling becomes increasingly rounded (less elongate), and ash exteriors generated by impacts becomes increasingly smooth (Fig. 4). Both milling and impact also remove adhering matrix from crystals (decrease AF), with milling the

most efficient at this process. Importantly, products of primary fragmentation retain much more adhering matrix than products of secondary fragmentation by either milling or impact (Fig. 9). Together these data provide guidelines for assessing primary vs. secondary contributions to the total grain size population within an eruptive deposit.

To further explore the role of crystals, we analysed the crystal content of volcanic ash derived from rapid decompression. Most crystals fracture along cleavage planes, leaving a smooth and clean breakage surface. During the evacuation of a crystal-rich magma body driven by rapid decompression, crystal fragmentation is inevitable and indeed characteristic of the associated tephra fallout. The proportion of free crystals depends on the grain size considered. X-Ray CT measurements combined with componentry allowed us to describe the shift in crystal size during fragmentation by rapid decompression. These data are important because the size fractions dominated by juvenile crystals show enhanced sedimentation because crystal densities exceed those of vesicular clasts. Thus we suggest that consideration of crystal sizes and proportions could improve settling calculations that inform hazard maps and dispersion models.

Finally we illustrate the application of our analysis techniques to two samples from Soufriere Hills Volcano, Montserrat: one dome collapse sample with co-PDC ash only, and one Vulcanian sample with contributions from both primary and co-PDC fragmentation. These samples are distinct in both their components and their grain characteristics, particularly AF. The Vulcanian sample is dominated by deep-derived microlite-poor and highly vesicular glass. Ash particles have polymodal AF distributions, with a dominant mode at low AF (little adhering glass) and two other modes at higher AF. We suggest that the high AF modes reflect primary fragmentation, while the low AF mode is the signature of co-PDC ash. The dome collapse sample, in contrast, is dominated by microlite-rich glass of variable vesicularity. In this sample, the AF distribution is approximately uniform, and most likely reflects the wide range of groundmass textures (both vesicularity and crystallinity). This latter observation brings up one further point, which is that both primary and secondary fragmentation processes are strongly dependent on the original magma components. As a result, a full understanding

of ash attributes produced by different mechanisms requires a systematic study using starting materials with different bubble and crystal attributes.

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822	Figure Captions:
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824	Figure 1: 3D surface reconstructions of the crystal phases. Processed using Avizo software from X-
825	Ray computed tomography generated image stacks. (a) SHV subvolume. (b) Cumulative frequency
826	diagram showing the relative crystal volume distributions within SHV pumice extracted from 3D
827	reconstructions.
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829	Figure 2: Diagrams of the various fragmentation apparatus used in this study. (a) shock tube
830	(modified after Alatorre-Ibargüengoitia et al. (2010)), (b) falling piston and (c) ball mill.
831	
832	Figure 3: Grain size distributions for (a) rapid decompression shock tube, (b) falling piston and (c)
833	ball mill experiments.
834	
835	Figure 4: Shape analysis from BSE SEM images of 2φ grain size fraction. Histograms show shape
836	data gathered from 2D SEM image mosaics. Where an axial ratio of one implies a perfectly round
837	grain and a convexity of one implies a smooth grain with the same perimeter as its convex hull. (a)
838	The difference in axial ratio of the grains produced by hot and cold decompression. (b) The change in
839	the axial ratio value for grains that were hit 5 and 10 times. (c) The change in the axial ratio value for
840	grains that were milled for 30 seconds and one minute. (d) The difference in convexity values of the
841	grains produced by hot and cold decompression. (e) The change in value of convexity in grains hit 5
842	and 10 times. (f) The change in the value of convexity for grains that were milled for 30 seconds and
843	one minute.
844	
845	Figure 5: Componentry results for experimentally generated ash. (a) SHV average componentry
846	results from rapid decompression, normalised per grain size fraction. (b) Individual ash grain
847	componentry from the 2ϕ grain size fraction using BSE SEM images of products of varying

fragmentation techniques.

850 Figure 6: Normalised crystal distributions for SHV represented as a cumulative distribution. Solid lines represent the crystal population present in the volcanic core prior to experimental fragmentation. 851 Dashed lines represent the experimentally fragmented crystal population. 852 853 854 Figure 7: SEM micrographs of crystals from the SHV 2φ ash fraction. (a) Hornblende crystal showing mainly a clean flat surface however extremely irregular at the edges. (b) Broken crystal with 855 856 a thin vesicular glass coating. (c) Hornblende crystal with a stepped fracture surface. (d) A plagioclase 857 crystal showing multiple river line fractures on broken crystal surface. 858 859 Figure 8: Power law plots for (a) the rapid decompression experiments, (b) the falling piston experiments and (c) the ball mill experiments. Data markers represent those included in the linear 860 861 regression. 862 Figure 9: Adherence Factor plot of the three fragmentation techniques: Milling (0.5 minutes and 1 863 minute), Impact (5 hits and 10 hits) rapid decompression (cold and hot) and natural samples 864 (Vulcanian and dome collapse). Adherence Factor (AF) = area of matrix / (area of matrix + area of 865 866 crystal) against the total cumulative per cent. A high value for AF represents a higher proportion of adhered matrix and thus the upper curve shows a greater amount of 'matrix stripping' than the lower 867 curve. Normalised Grain No. is the value of AF in descending order, normalised to one. 868 869 Figure 10: Componentry of two natural SHV samples. The samples are from fallout deposits 870 produced by a dome collapse event (on 31 March 1997) and a Vulcanian eruption (on 26 September 871 1997) of Soufriere Hills Volcano. 872 873 Figure 11: A summary cartoon relating the three experimental fragmentation mechanism to the 874

natural volcanic scenario. Primary fragmentation through rapid decompression produces pyroclasts

with D ~2.2. Then through abrasion and milling, selective secondary fragmentation increases D and adds to the fine ash component to the GSD. Table 1: Summary of all experimental runs and characterisation. Grey fields indicate that the characterisation has been performed. **Table 2:** A review of relevant fragmented material and the associated fractal dimensions obtained. Figure S1: Individual power-law plots for each experimental fragmentation experiment: (a) rapid decompression (b) impact and (c) milling. Open circles represent the data points not included in the linear fitting. Figure S2: Histogram of adherence factor values for decompression (hot and cold), impact (5 hits and 10 hits), milling (30 seconds and 1 minute) and natural samples (Vulcanian and dome collapse). Shown on bars is the proportion of broken crystals amongst the crystal population where a broken crystal is defined as a crystal that clearly does not display an intact ring of zoning.

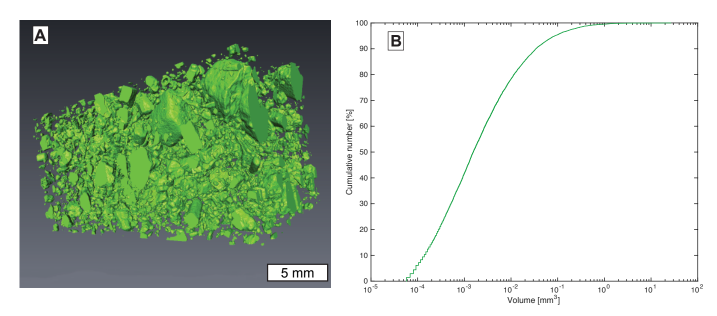


Figure 1: Jones et al (2016, JVGR) Size: 1.5 column

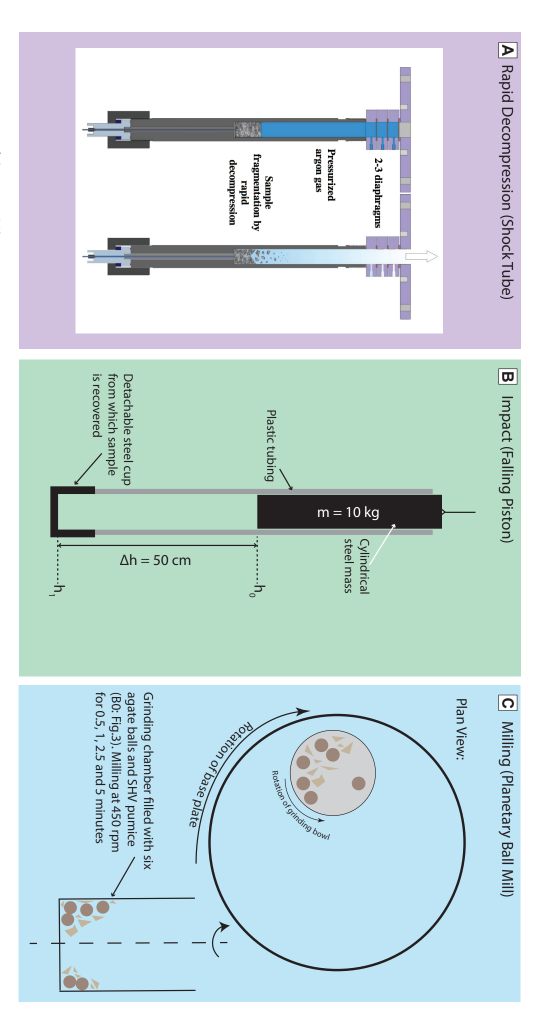


Figure 2: Jones et al. (2016, JVGR) Size: Double column

Figure 3: Jones et al. (2016, JVGR) Size: Double column

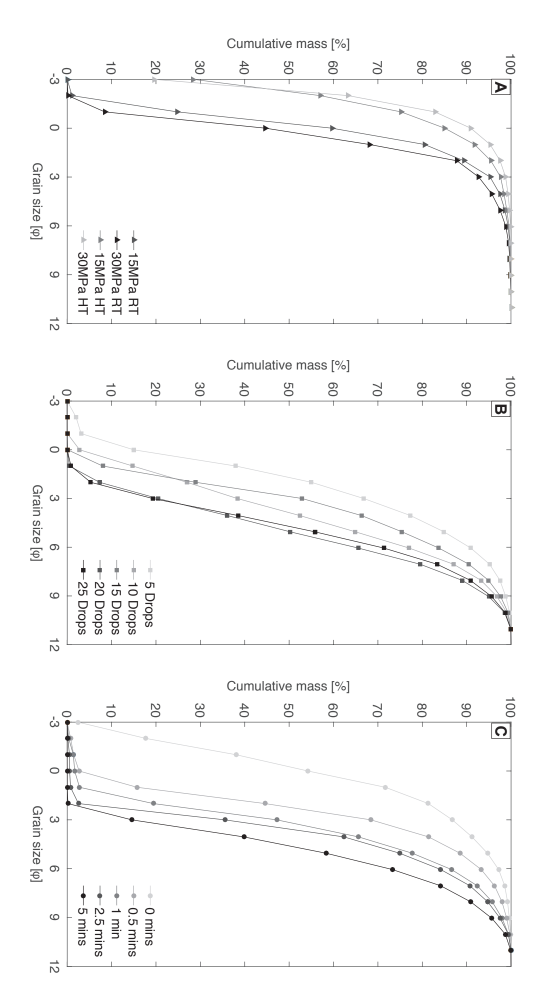
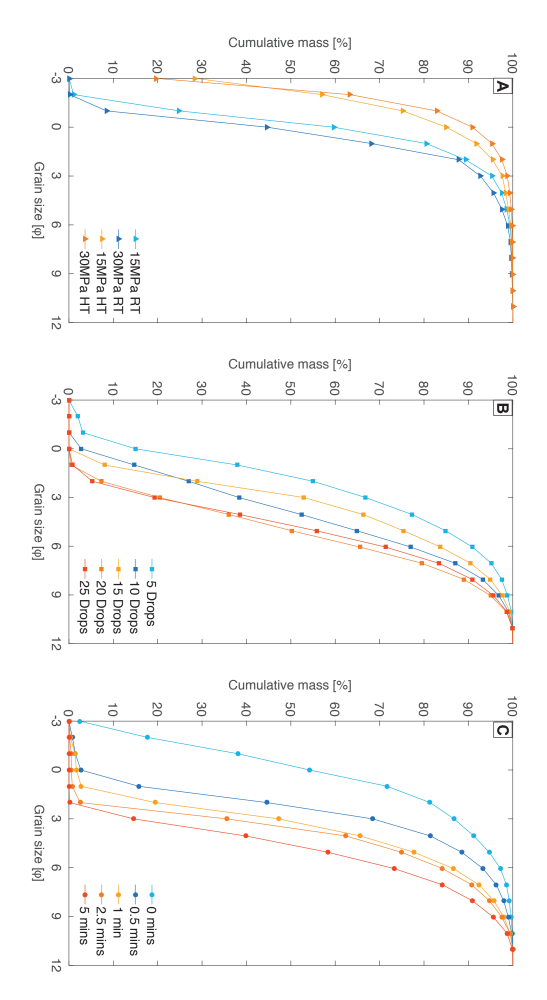


Figure 3: Jones et al. (2016, JVGR) Size: Double column



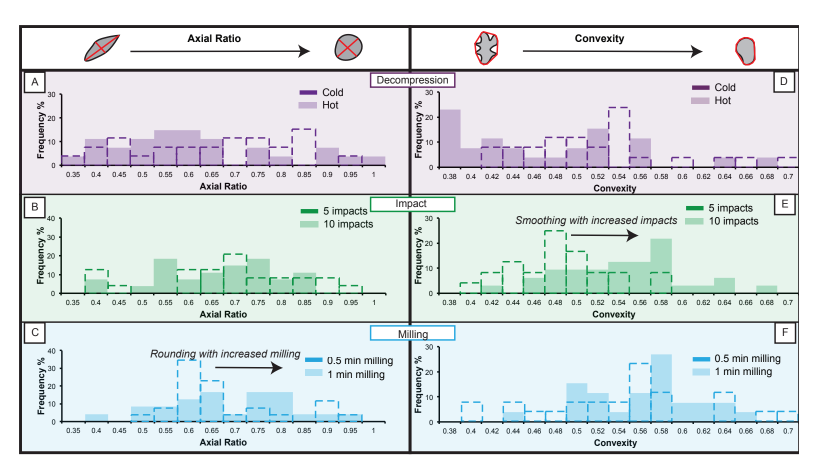


Figure 4: Jones et al. (2016, JGVR)

Size: 2 column

≻ wt.% 25 20 10 15 ယ် Clasts Mafic crystals Felsic crystals $\dot{\diamond}$ 6 0 N ω В Frequency (%) 100 20 30 90 10 40 50 60 70 8 5 hits Impact 10 hits Fragmentation method
Milling F 0.5 mins 1 min Rapid Decompression cold hot Crystal
Crystals with attached matrix
Vesicular clasts

Figure 5: Jones et al (2016, JVGR) Size: Double Collumn

Figure 6: Jones et al. (2016, JVGR) Size: Single Column Cumulative crystal weight % **N** - Ash - Rock

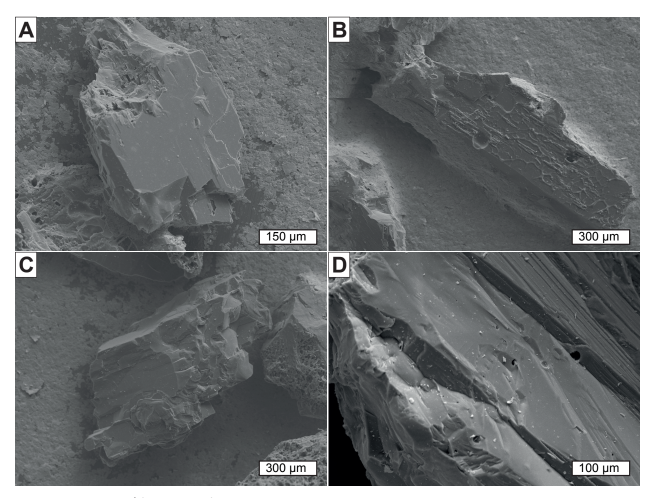


Figure 7: Jones et al (2016, JVGR) Size: Single column

Log(N>L) -½ ω Γ 10 N တ 0 ∞ — D = 2.07 (15 MPa 25°C) — D = 2.15 (30 MPa 25°C) — D = 2.24 (15 MPa 880°C) — D = 2.03 (30 MPa 880°C) 7 Log(L) [mm] \triangleright Log(N>L) . 3 ⊏ 10 တ 0 N ∞ D = 2.51 (5 Hits)
D = 2.73 (10 Hits)
D = 2.52 (15 Hits)
D = 2.69 (20 Hits)
D = 2.58 (25 Hits) Log(L) [mm] 0 W Log(N>L) 10 -\frac{1}{2} <u></u> ∞ 0 \sim -D = 2.33 (0.5 min) -D = 2.37 (1 min) -D = 2.43 (2.5 mins) -D = 2.45 (5 mins) Log(L) [mm] 0 C

Figure 8: Jones et al. (2016, JVGR) Size: Double column

Log(N>L) -½ ω Γ 10 တ 0 N ∞ — D = 2.07 (15 MPa 25°C) — D = 2.15 (30 MPa 25°C) — D = 2.24 (15 MPa 880°C) — D = 2.03 (30 MPa 880°C) 7 Log(L) [mm] \triangleright Log(N>L) . 3 ⊏ 10 တ 0 N ∞ -D = 2.51 (5 Hits) -D = 2.73 (10 Hits) -D = 2.52 (15 Hits) -D = 2.69 (20 Hits) -D = 2.58 (25 Hits) Log(L) [mm] 0 W Log(N>L) -2 -3 10 2 <u></u> ∞ 0 -D = 2.33 (0.5 min) -D = 2.37 (1 min) -D = 2.43 (2.5 mins) -D = 2.45 (5 mins) Log(L) [mm] 0 C

Figure 8: Jones et al. (2016, JVGR) Size: Double column

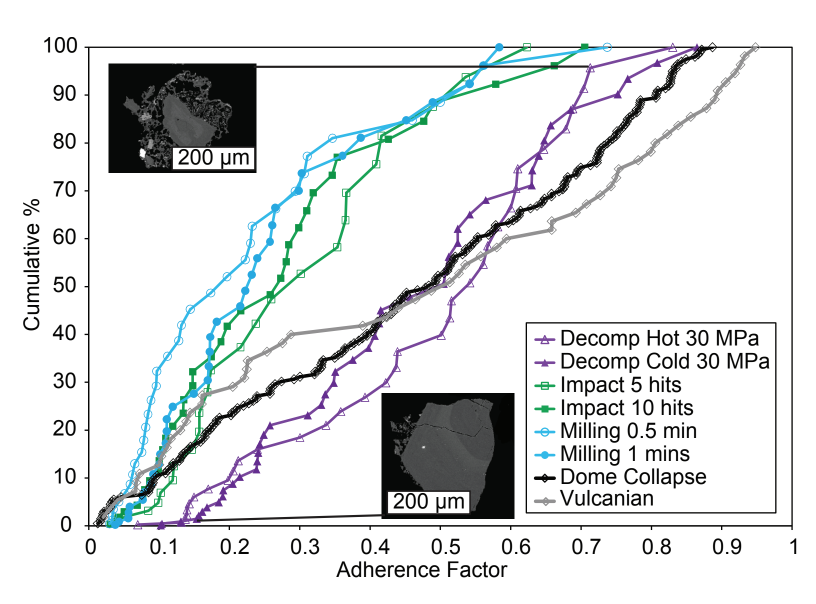


Figure 9: Jones et al. (2016, JGVR) Size: Single column

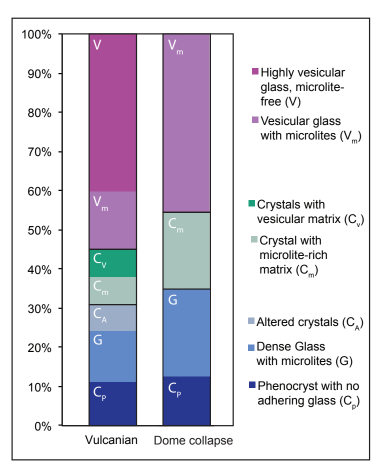


Figure 10: Jones et al. (2016, JVGR) Size: Single Column

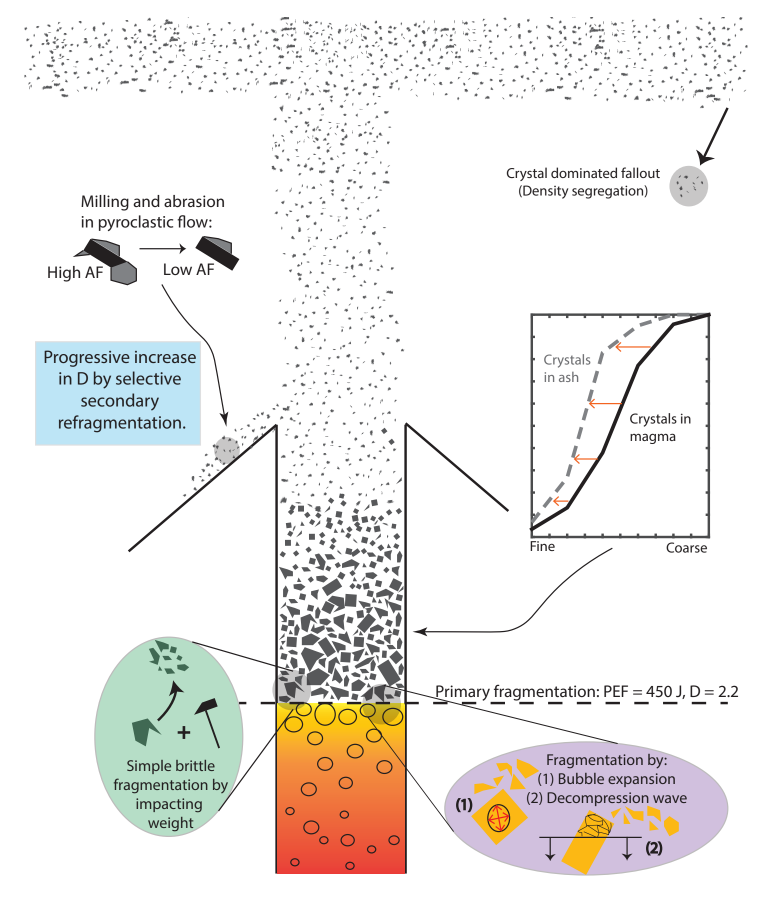


Figure 11: Jones et al (2016, JVGR) Size: Single collumn