#### **ORIGINAL PAPER**



# Microlite orientation in obsidian flow measured by synchrotron X-ray diffraction

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Received: 16 April 2018 / Accepted: 31 May 2018 © Springer-Verlag GmbH Germany, part of Springer Nature 2018

#### Abstract

Clinopyroxene and plagioclase (andesine) microlites in an obsidian flow from Glass Mountain (NE California, USA) display strong alignment. Synchrotron X-ray diffraction, coupled with Rietveld analysis, was used to quantify crystallographic-preferred orientation (CPO). Clinopyroxene, with a rod-shaped morphology, shows a strong alignment of [001] in the flow direction and (010) aligned parallel to the inferred flow plane. Andesine, with a platy morphology, displays an alignment of (010) platelets in the flow plane. Some pole densities exceed 90 multiples of random distribution. Applying a model of rigid ellipsoidal inclusions in a viscous matrix, the local pure shear strains are between 2 and 3.

Keywords Obsidian · Microlite alignment · Texture analysis · Crystal-preferred orientation, Strain estimates

#### Introduction

Obsidian in natural lava flows preserves a number of textural features that record its eruption and emplacement (e.g., Smith 2002). Structural features range from folds and layering at centimeter-to-decameter scales (Fink 1983; Manley and Fink 1987; Castro and Cashman 1999), to deformed bubbles at the micrometer-to-millimeter scale (e.g., Rust et al. 2003; Gardner et al. 2017) and aligned elongated micrometer-sized crystallites at the smallest scales (Manga 1998; Castro et al. 2002). Together, these textures provide information about flow, fragmentation, and solidification processes.

Microlites are oriented by flow, and their orientation distribution thus offers an opportunity to determine the type of flow and strain experienced by the melt prior to quenching (e.g., Castro et al. 2002; Befus et al. 2014, 2015). Quantitative information, including the relationship between size and orientation, can be obtained with optical microscopy

Communicated by Timothy L. Grove.

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(e.g., Manga 1998; Castro et al. 2004). Because microlites are small, however, it is not usually possible to determine crystallographic orientation, and microlites with diameters smaller than about 1 µm cannot be characterized. Here, we use synchrotron X-ray diffraction and scanning electron microscopy to investigate two rhyolitic obsidian samples from the Glass Mountain lava flow, Medicine Lake Volcano, Northern California.

Synchrotron hard X-ray diffraction proved to be a very useful method for texture analysis in a broad range of different materials from metals (e.g., Wu et al. 2010) to complex clays (e.g., Wenk et al. 2010), slates (e.g., Haerinck et al. 2015), and serpentinite (e.g., Soda and Wenk 2014). This is a first application of the method to volcanic rocks. Contrary to optical microscopy referred to above that describes the shape-preferred orientation (SPO) of particles, X-ray texture analysis provides a quantitative description of the crystallographic orientation distribution of clinopyroxene and plagioclase, averaging over crystallite volumes, not over crystallite numbers. The analysis is by no means trivial in the case of obsidian, since crystallites contribute less than 2% of the volume and diffraction from those crystals is weak compared to scattering from the amorphous glass. Nevertheless, results for SPO and CPO are comparable and can be used to estimate local strain patterns based on Jeffery's (1922) solution for the motion of rigid ellipsoidal inclusions in a viscous matrix.



Sample 2 was also analyzed with a JEOL SEM at the Engineering Department of the University of Trento, Italy,

# **Experimental**

# **Samples**

The two samples investigated are obsidian fragments from Glass Mountain, CA, USA (41°36′15.5″N/121°28′34.5″W). This ~ 1 km³ compositionally zoned flow erupted about 1070 AD and is the youngest eruption at the Medicine Lake shield volcano (Grove et al. 1997, Donnelly-Nolan et al. 2016). Optical microscopy reveals a layered structure with layers 2–5 mm in thickness (Fig. 1a). Some layers are more uniform than others. At higher magnification, a strong alignment of needle-shaped microlites is apparent (Fig. 1b). Locally, there is distorted flow around larger inclusions of magnetite (Fig. 1c) and fragments of plagioclase (Fig. 1d).

In this study, we focus on homogeneous layers in two samples. In sample 1, microlites are almost exclusively clinopyroxene (Fig. 1b). Sample 2 is a mixture of plagioclase and clinopyroxene microlites. Views of sample 2 with plane polarizers (Fig. 2a, c) and crossed polarizers (Fig. 2b, d), both parallel to the flow direction (Fig. 2a, b) and perpendicular to the flow direction (Fig. 2c, d), show the morphology and the strong alignment of microlites. Clinopyroxenes are needles aligned in the flow direction and plagioclase microlites are platelets aligned in the flow plane. Plagioclase platelets are typically larger in size than clinopyroxene needles which is most obvious with crossed polarizers.

to identify the phases and determine their morphology. Back-scattered images, again parallel and perpendicular to the flow direction, are shown in Fig. 3. Pyroxenes are easily recognized, with high contrast. They are prismatic needles with a pseudohexagonal cross section, roughly 2–10 µm long and 0.1–2 µm in diameter. Contrast for plagioclase is weak because of a similar chemical composition as the glass matrix. Some crystals are marked in Fig. 3. They are larger than pyroxenes and have a platelet morphology (see also Fig. 2). They range in size from 2 to 25 µm long to 2-5 µm wide. The chemical compositions of the phases were analyzed with an EDAX EDS system (Fig. 4; Table 1). The pyroxene is identified as ferrous intermediate augite/pigeonite (Ca<sub>0.18</sub>, Mg<sub>0.43</sub>, Fe<sub>0.36</sub>,  $Na_{0.03}$ )  $SiO_3$ , plagioclase as calcic andesine ( $Ca_{0.40}$   $Na_{0.52}$  $K_{0.08}$ )(Si<sub>0.72</sub>, Al<sub>0.28</sub>)O<sub>8</sub>, and glass as high in SiO<sub>2</sub> (> 70%) with significant aluminum, potassium, and sodium. Hard X-ray synchrotron diffraction

Samples for the X-ray diffraction experiment were  $\sim 0.7$  mm-thick slabs with the flow plane perpendicular to the slab surface. The diffraction measurements were carried out at beamline 11-ID-C at the Advanced Photon Source (APS) of Argonne National Laboratory. Experiments were performed in transmission geometry. A beam of X-rays ( $\lambda = 0.10738 \text{ Å}$ ), 0.5 mm in diameter, irradiates the sample, and diffraction patterns were recorded by a Mar345 image plate detector

Fig. 1 Optical micrographs of obsidian in sample O1. Plane polarized light. a Structure with compositional layering (Os2a), b detail of layering with microlites, c microlites around an inclusion of magnetite, and d inclusion of plagioclase. In c, the curves show streamlines for pure shear around a spherical and rigid inclusion

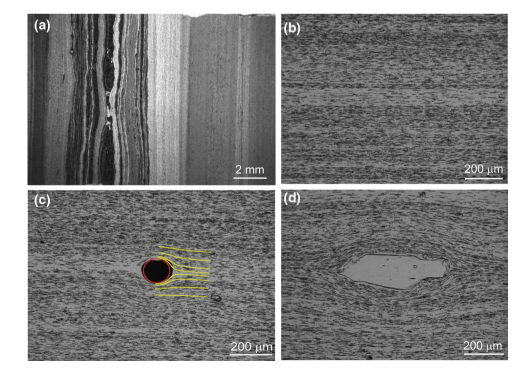
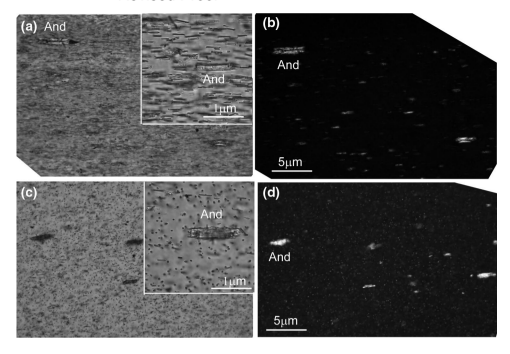




Fig. 2 Optical micrographs of obsidian in sample O2, a, b parallel to the flow direction and c, d perpendicular to the flow direction; a, c are plane polarized light and b, d crossed polarizers. Microlites are andesine (And) and clinopyroxene needles (lines in a and dots in c). In the upper right corner of a and c, an enlarged segment is displayed



 $(3450 \times 3450 \text{ pixels})$  placed about 2 m behind the sample. Exposure time was 900 s. High energy maximizes sample penetration. The beam size was chosen to give a sufficiently large sample volume (~0.2 mm<sup>3</sup>) to be representative of a single flow layer. The diffraction images display Debye rings that have intensity variations as a function of the azimuth because of preferred orientation of crystallites (Fig. 5). A dark broad ring is due to scattering from glass and sharp rings are scattering from crystallites.

The sample slab was mounted on a goniometer and was rotated around a horizontal axis perpendicular to the flow plane and the incident X-ray beam. The sample was tilted in 15° increments to seven positions  $(-45^{\circ}, -30^{\circ}, -15^{\circ}, 0^{\circ},$ 15°, 30°, and 45°) to obtain a good coverage of orientation space for texture analysis. This results in seven diffraction images for each data set. The flow layers are perpendicular to the tilt axis to avoid overlapping of layers at high tilt angles. For both samples, diffraction images were recorded for two layers (O11, O12, O22, and O25).

#### **Rietveld analysis**

The seven diffraction images for each sample location were then analyzed simultaneously with the Rietveld method as implemented in the software MAUD (Lutterotti et al. 2014). Intensities on each diffraction image (Fig. 5) were integrated over 5° azimuthal sectors, resulting in 72 spectra. In the Rietveld method, instrumental parameters, structural and microstructural characteristics, and preferred orientation patterns are refined until an optimal fit is obtained with  $7 \times 72 = 504$  diffraction spectra.

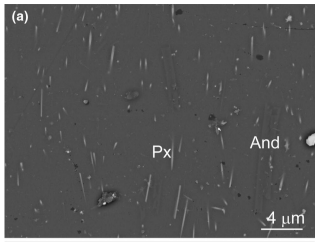
First, instrumental parameters such as sample detector distance, detector orientation, and Caglioti parameters defining the diffraction resolution were refined from diffraction data collected for a CeO<sub>2</sub> standard. These instrument parameters were then applied to each of the four obsidian sample data sets.

The Rietveld refinement requires as input the crystallographic characterization of the composing phases. For clinopyroxene, the C2/c structure was used (Clark et al. 1969, # 1000035). For the texture analysis of monoclinic crystals in MAUD, the first setting (z monoclinic axis) has to be applied (Matthies and Wenk 2009), but results are shown in the more conventional second setting (y monoclinic axis). For plagioclase andesine, I1 was used which had to be converted to P1 (Fitzgerald et al. 1986, #9001030). The glass phase, giving rise to the diffuse diffraction peaks at  $Q \sim 1.6$  and 4.3 Å<sup>-1</sup>  $(Q = 2\pi/d)$ , is more difficult to describe. For some refinements, a nanocrystalline form of tetragonal  $\alpha$  (low) cristobalite (Hosemann et al. 1991, spacegroup P4<sub>1</sub>2<sub>1</sub>2) was used, which was first applied to silica glass by Le Bail (1995) and later by Lutterotti et al. (1998).

From seven diffraction images at different tilts for each, polynomial background peaks, phase volume fractions, lattice parameters, grain sizes of phases, and textures (CPO) of clinopyroxene and andesine were refined. Grain sizes (average coherently diffracting domain size) were refined with the isotropic Popa model (Popa 1998), converging for glass at ~8 Å which is about the size of the unit cell, for clinopyroxene at ~300 Å and for plagioclase at ~800 Å.

In a first cycle three phases, glass (modeled as nanocrystalline cristobalite), clinopyroxene, and andesine were





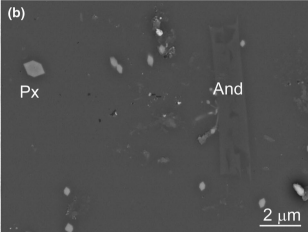
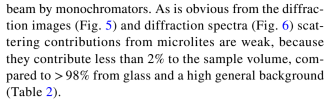


Fig. 3 SEM back-scattered electron images of sample O2, parallel to flow direction (a) and perpendicular to flow direction (b). The contrast of andesine is very close to that of glass and, therefore, difficult to recognize. The flow plane is vertical

included in the refinement to obtain phase proportions, though the assumptions of pure SiO<sub>2</sub> for cristobalite are obviously an approximation. Data and fit for one image of sample O22 are shown for an average spectrum (combining all 72 azimuthal spectra) (Fig. 6a) and for the stack of azimuthal spectra with measured data at the bottom and Rietveld fit on top (Fig. 6b). Note that  $Q = 2\pi/d$  is used as horizontal axis, rather than  $2\theta$  which depends on the rather arbitrary experiment wavelength, with  $2\theta = 2\arcsin(\lambda/2d)$ . The large and broad diffraction peak at  $Q \sim 1.6 \text{ Å}^{-1}$  (Fig. 6a) is from glass and corresponds to cristobalite (101). Two characteristic peaks of andesine and clinopyroxene are labeled. Volume fractions are summarized in Table 2.

On the "mapplot" (Fig. 6b), a weak azimuthal intensity variation of the strong glass peak at  $Q \sim 1.6 \text{ Å}^{-1}$  is present. It was established that this is not due to alignment of disordered silica structure components but an artifact due to the horizontal polarization of the X-ray



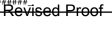
In a second cycle of refinements, glass was omitted as a phase, and the contribution of clinopyroxene and andesine microlites was extracted above a "background" modeled with a Gaussian peak to account for the contribution from glass, in addition to a fifth-order polynomial and using a more limited *Q* range (1.8–4.2  $\text{Å}^{-1}$  for sample 1 and 1.4–4.0  $\text{Å}^{-1}$  for sample 2). For the texture refinement, the EWIMV method was applied with a resolution of 5° for clinopyroxene and 7.5° for andesine. Rietveld fits for one image of each scan are shown in stacked spectra (Fig. 7). Looking at diffraction peaks in Fig. 6a (bottom), there is a lot of overlap of peaks for these low symmetry minerals and caution is required to have high resolution and avoid artifacts from peak overlaps. The reliability of the fit is expressed in map plots with the good agreement of the fit (Fig. 7 top) with the experimental data (Fig. 7 bottom). These refinements provided the orientation distributions that were exported from MAUD and then used in the software BEARTEX (Wenk et al. 1998) for pole figure calculations. The intensity variations (Figs. 5, 6, 7) are very strong, especially for sample 1 with only clinopyroxene microlites. Average refined lattice parameters are listed in Table 3 and a summary of texture strength is given in Table 4.

#### Results

Optical microscopy provides an overview of the microstructure. The two samples of obsidian are composed of layers, 0.1-2 mm thickness defined by differences in microlite volume fraction, and some layers are more heterogeneous than others (Fig. 1a). Four fairly homogeneous layers were selected for this study (e.g., Figs. 1b, 2). Clinopyroxene microlites are 5–15 μm in length and 0.2–0.5 μm wide (inset in Fig. 2a, c). Many needles have (010) parallel to the inferred flow plane. The needle axis [001] [~ $(10\overline{2})$  normal] is strongly aligned in the flow direction. Figure 2c, with a view in the flow direction, displays point-like crystallites. SEM images confirm a prismatic morphology with a pseudohexagonal cross section. The prism (010) planes are preferentially aligned in the flow plane (Figs. 3b, 8a).

Andesine has a platelet morphology with (010) parallel to the platelet (Figs. 2, 8b). Grains are overall larger than clinopyroxene, 10-100 µm in length and 1-10 µm in width. These platelets are aligned, which is most obvious when viewed with crossed polars where they all shine up simultaneously





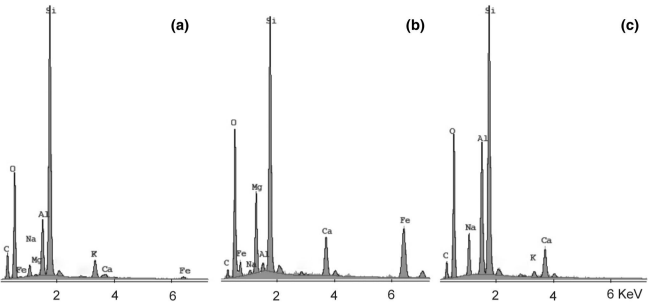


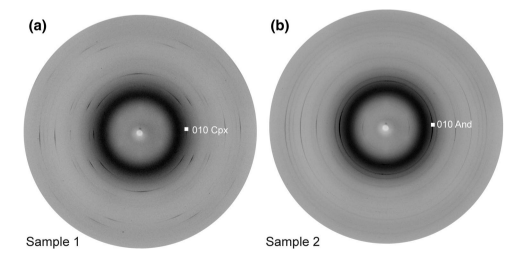
Fig. 4 EDS spectra for a glass, b clinopyroxene and c andesine (compared with Table 2)

Table 1 Chemical composition of glass and microlites in sample 2, measured at two locations with SEM EDS analyses

Glass			Andesine		Pigeonite		
Na	4.3	4.8	10.0	8.9	1.7	1.2	
Mg	0.4	_	_	_	21.0	19.0	
Al	14.0	14.5	23.4	23.1	2.1	4.4	
Si	72.5	71.5	58.4	58.7	49.3	49.9	
K	6.3	6.2	1.4	1.4	0.1	_	
Ca	1.1	1.3	6.9	7.9	7.5	10.2	
Fe	1.3	1.7	_	_	18.4	15.3	

Atomic% (compare with Fig. 4)

Fig. 5 Selected diffraction images for sample 1 (O11) and sample 2 (O22). Some diffraction peaks are indexed. The broad ring at low angles is due to scattering from silica glass



at a certain rotation (Fig. 2b, d). SEM images display a skeletal morphology of andesine, with some inclusions of clinopyroxene.

The alignment of microlites is consistent with flow of rigid inclusions in a viscous matrix, particularly the disruption of flow around larger inclusions (Fig. 1c, d). Since





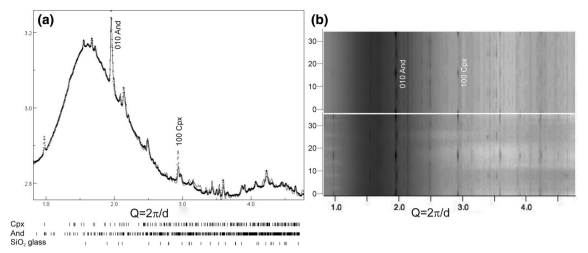


Fig. 6 a Azimuthal average of an image for sample O22 with Rietveld fit for clinopyroxene, andesine, and the broad peak for silica glass, approximated by nanocrystalline cristobalite. Dots are observed values and line is the Rietveld fit. Below the spectrum diffraction peaks of the three phases are indicated and some peaks are indexed.

**b** Stack of 72 diffraction spectra at 5° azimuthal intervals. Bottom is experimental data and top is Rietveld fit. A logarithmic intensity scale is used to emphasize weak diffractions. Horizontal axis is  $Q = 2\pi/d$ 

**Table 2** Volume fractions of glass and microlites based on Rietveld refinement

	O11	O12	O22		O25	
Glass	0.993	0.989	0.971		0.986	
Pigeonite	0.007	0.011	0.006	0.235(1)	0.008	0.679(3)
Andesine			0.024	0.765 (1)	0.006	0.321(3)

Standard deviations in parentheses; for sample 2, also volume fractions of crystallites

microlites have this euhedral morphology (Fig. 8), there is a perfect coincidence between shape-preferred orientation (SPO) and crystal-preferred orientation (CPO). The latter is explored with the diffraction experiments.

For the X-ray analysis, we selected layers with intermediate microlite fractions, large enough to provide sufficient diffracting volume and small enough to expect homogeneous flow. Glass volume fractions for O11 and O12 are > 99%. For O22 and O25, there is a slightly larger amount of microlites ( $\sim$  0.97 of glass). In O11 and O12, no andesine could be documented. In O22, there is  $\sim$  24% clinopyroxene versus  $\sim$  76% andesine, and in O25, there is  $\sim$  68% clinopyroxene versus  $\sim$  32% andesine (Table 2).

Figures 9, 10 show pole figures for selected crystal directions, after rotating all samples into the same orientation with the flow (layer) plane as projection plane and the flow direction horizontal. The pole figures confirm the qualitative optical results. There is, indeed, very strong preferred orientation in all scans with very high maxima for planes aligned in the flow plane and prism axes aligned in the flow direction as expressed in multiples of a random distribution in Table 3. For pigeonite, the poles to (010) and {110} are aligned preferentially perpendicular to the flow plane, and

[001] is parallel to the flow direction (Fig. 9). For andesine, (010) is the platelet plane and aligned in the flow plane (Fig. 10).

It should be emphasized that the quantitative data analysis is by no means trivial. One issue is the peak overlaps of the low symmetry microlite phases (Fig. 6a). A second problem is the very weak scattering from crystalline microlite phases compared to the broad background of dominant scattering from glass. Most diffraction peaks are less than 10% above background. To minimize problems from overlaps, textures of the two microlite phases were refined separately: first for andesine and then for clinopyroxene. Major overlaps were expected from diffraction on lattice planes aligned parallel to the flow plane and giving a signal at 0 azimuth (bottom of the stack of spectra). Therefore, these spectra were disabled from the refinement, but it turned out that it did not make any difference.

The preferred orientation does not only display very strong peaks (measured in multiples of random distribution, m.r.d.) but also very sharp peaks, especially for clinopyroxene with width of CPO peaks at half maximum ranging from 15° to 25°. All pole figures (Table 4) also show a significant background ranging from 0.1 to 0.5 m.r.d. Part



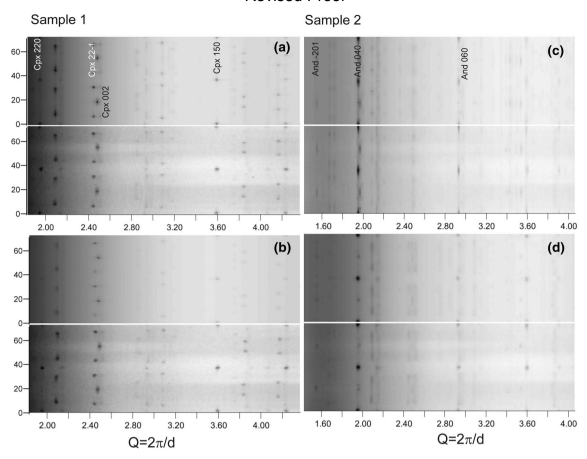


Fig. 7 Refinements of two scans on sample 1 and sample 2. Bottom is stack of 72 experimental spectra for different azimuths (c.f. Fig. 5). Top is corresponding Rietveld fit. a O11, b O12, c O22, and d O25. Horizontal scale is  $Q = 2\pi/d$  (Å<sup>-1</sup>). Logarithmic intensity scale

Table 3 Average lattice parameters for clinopyroxene and andesine based on Rietveld refinements

	a (Å)	b (Å)	c (Å)	α (°)	β (°)	γ (°)	
Clinopy	roxene C2/c (sec	cond setting)			'		
O12	9.7573 (8)	8.8956 (2)	5.2872 (2)	5.2872 (2) 106.810 (6			
O11	9.7482 (6)	8.8977 (3)	5.2846 (3)	5.2846 (3) 106.782 (5)			
O22	9.7123 (9)	8.9112 (5)	5.3022 (4)				
O25	9.7260 (6)	8.8942 (5)	5.2827 (3) 106.894 (6)				
Andesin	e I-1						
O22	8.1692 (3)	12.8796 (4)	7.1203 (2)	93.280 (4)	116.360(3)	90.029 (3)	
O25	8.1740 (8)	12.9005 (3)	7.1126 (3)	93.149 (11)	115.950 (6)	89.793 (10)	

Standard deviations in parentheses

of the background is due to randomly oriented crystallites such as clinopyroxene clusters or dispersion around inclusions (Fig. 1c, d) and nanocrystalline microlites but part may also be an artifact due to the high background and limited counting statistics. Yet, note that in measured diffraction images (Fig. 5) and measured diffraction patterns (bottom of Fig. 7), there is residual intensity for all azimuthal angles, not just for the intensity maxima. For scan O22, we have explored the effect of background and apparent high random contribution by first calculating pole figures, then subtracting from pole figures the random contribution, and then using these new pole figures to calculate new orientation distributions with WIMV in BEARTEX. It turns out that the new orientation maxima have increased by a factor 2–3 (bold numbers in Table 4). These are representative of the aligned crystallites during viscous flow.

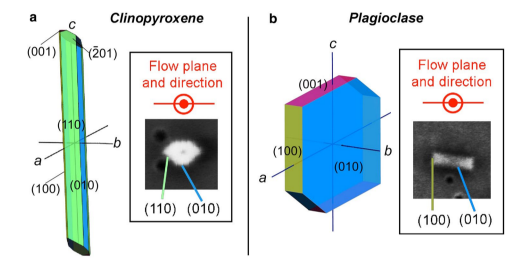


**Table 4** Pole figure maxima and minima in multiples of random distribution

	O11		O12		O22		O25		O25-sub
Pigeonite	(monoclini	c, second s	setting)						
(100)	13.0	0.2	10.4	0.3	5.8	0.2	7.6	0.5	13
(010)	92.6	0.1	58.3	0.3	23.7	0.2	33.3	0.4	43
[001]	35.3	0.1	27.5	0.3	12.5	0.3	19.1	0.4	30
Andesine									
(100)					4.0	0.4	4.0	0.4	8
(010)					19.5	0.4	29.4	0.3	49
(001)					3.0	0.4	4.8	0.4	10

For sample O25, we also show maxima after subtracting a background of 0.5 ("phon") from pole figures and recalculating ODF (bold)

Fig. 8 Schematic sketch of the morphology of a clinopyroxene and b andesine crystals. On the right side is an SEM image of a crystallite viewed in the flow direction



## Discussion

The alignment of microlites parallel to bands is the result of flow and confirms that the visibly identified layers in obsidian are in fact flow bands. Although the melt may have fractured and rewelded multiple times to create the heterogeneities that define the various layers (e.g., Tuffen et al. 2003; Gonnermann and Manga 2003, 2005; Gardner et al. 2017), the alignment of microlites with the macroscopic layers requires that much of the strain we see accumulated after the last welding event and that microlites were reoriented after the last welding event.

The measurements show that not only are microlites aligned according to their morphology (SPO) in the flow field, but also that their crystallographic axes are aligned (CPO). It is unclear whether the flow-induced orientation has consequences for the crystal habit and their growth during emplacement.

The preferred orientation of microlites in obsidian is an example of alignment of rigid particles in a viscous medium. Flow is governed by Stokes equations for viscous flow, since the Reynolds number is  $\ll 1$  owing the high viscosity, 10<sup>8.6</sup> Pa s using the composition in Giachetti et al. (2015), the mean water content of 0.22 weight% reported by Tatlock et al. (1976), a temperature of 850 °C (Waters et al. 2015), and the viscosity model of Giordano et al. (2008). Jeffery (1922) solved the equations for the motion of an ellipsoid at low Reynolds number (Stokes flow). This solution was later advanced by March (1932) to establish the relationship between the orientation of ellipsoidal particles and the overall strain. Sander (1950) applied the March concept to explain preferred orientation of phyllosilicates in slates and there are several applications reviewed by Oertel (1983). If we assume a unit sphere with a random needle distribution and deform the sphere into an ellipsoid with axes  $E_1 > E_2$ ,  $E_3$ , the particles become aligned, mainly parallel to the principal stretch  $E_1$ . According to Oertel (1985):

$$E_1 = \rho_n^{1/3},$$

where  $\rho_n$  is the pole density of the needles. In addition, from the stretch, we obtain the strain

$$\varepsilon_1 = \rho_n^{1/3} - 1.$$



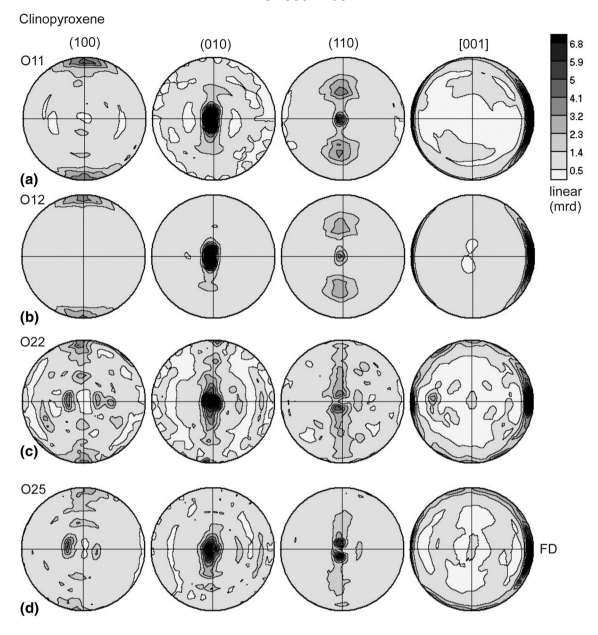


Fig. 9 Pole figures for clinopyroxene for samples 1 and 2 projected on the flow plane. FD is flow direction. Equal area projection, linear pole density scale

Platelet-shaped particles align with platelet normal preferably parallel to the smallest stretch  $E_3$ :

$$E_3 = \rho_p^{-1/3}$$

and strain

$$\varepsilon_3 = \rho_p^{-1/3} - 1.$$

Microlites in obsidian is an ideal case to apply the March model. In the case of O25, we can use the concentration of clinopyroxene [001] in the flow direction to estimate the extensional stretch and strain, and andesine (010) to determine the compressional stretch and strain. With  $\rho_n \sim 19$  m.r.d. for clinopyroxene, we obtain  $E_1 = 2.7$ , i.e., the long ellipsoid axis is 2.7 times longer than the original sphere diameter. This corresponds to a strain  $\varepsilon_1 = 1.7$ . For andesine platelets with  $\rho_p \sim 29$  m.r.d., we obtain a stretch  $E_3 = 0.32$  and a strain  $\varepsilon_3 = -0.68$ .

These values are for the case with a randomly oriented component that is not related to the alignment by flow. For the probably more realistic situation where the random component is subtracted (Table 4, right column, bold), pole density maxima increase greatly. With  $\rho_n \sim 30$  m.r.d. for



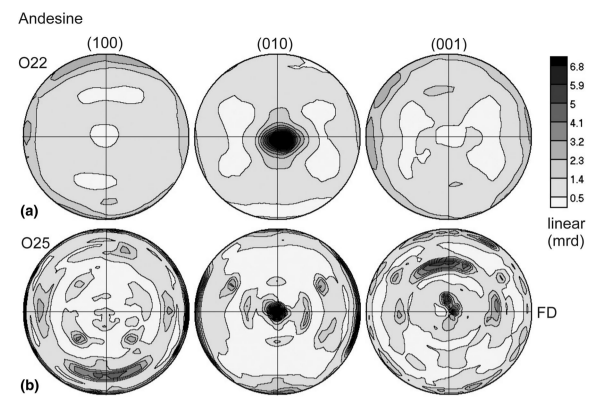


Fig. 10 Pole figures for andesine for sample 2 projected on the flow plane. FD is flow direction. Equal area projection, linear pole density scale

clinopyroxene, we obtain  $E_1$  = 3.1, and a strain  $\varepsilon_1$  = 2.1, and with  $\rho_p$  ~ 49 m.r.d. for andesine, we obtain  $E_3$  = 0.27, and a strain  $\varepsilon_3$  = -0.73.

There is a little evidence for deformation in the intermediate direction, suggesting overall plane strain and pure shear. Further support for the dominance of pure shear is provided by the orientation of the microlites around large inclusions. In Fig. 1c, we also show streamlines (lines parallel to the velocity) for pure shear around a spherical inclusion. As the flow is fully three-dimensional, we show the flow in the midplane of the particle and a cross section parallel to the flow direction. Equations for this problem are presented in Manga (2005). The orientations of microlites are very close to the orientation of streamlines for pure shear flow around a rigid inclusion. In contrast, Rust et al. (2003) measured the orientation distribution and aspect ratio of deformed bubbles in a sample from the same flow and inferred that deformation was dominated by simple shear. These two inferences, that there was pure and simple shear, need not be incompatible, since different parts of the flow will experience different strains: pure shear near the top, simple shear near the base (e.g., Buisson and Merle 2002; Ventura 2004). The sample analyzed here was collected from the flow front and about half-way between the surface and the bottom of the flow.

The magnitude of strain is consistent with the previous investigations of microlite orientations and bubble deformation that suggested strains of the order of 2–3 for pure shear (Manga 1998; Castro et al. 2002; Befus et al. 2014, 2015). These estimates all build on the solution of Jeffrey (1922) which is for motion of a single isolated particle. Even though the volume fraction of crystals is small (less than 3%), because the microlites are elongate, there are hydrodynamic interactions between the particles (Rahnama et al. 1995) and these do have a small effect on the orientation distribution for the observed aspect ratios and concentrations in obsidian (Manga 1998).

A number of studies have used optical microscopy to determine the orientation distribution of microlites and then to quantify kinematics: the total strain and flow type that accompanied rhyolite flow emplacement (Castro et al. 2002; Befus et al. 2014, 2015). Such measurements of microlite morphology are very time-consuming and rely on data from individual crystallites. Generally, larger crystallites are selected, because they are optically visible.

The diffraction measurements reported here are based on overall volume fractions. Quantitative texture analysis using hard synchrotron X-rays has advanced through a combination of experiments and data analysis (e.g., Lutterotti et al. 2014; Wenk et al. 2014). A single measurement, e.g., at beamline APS 11-ID-C, takes less than half an hour. Yet, it requires access to a synchrotron which must be obtained through proposals and is usually limited to one or two



sessions a year. Data analysis can be very involved depending on sample complexities and it needs some experience.

The CPO patterns are statistical averages over selected sample volumes, and for interpretation of pole figure data, additional information about crystal morphology is required (e.g. Fig. 8). However, the method provides data about multiple minerals and smaller crystals than those resolved with optical methods. It might also allow us to document when different crystal populations nucleated and become oriented, but this would require examining a suite of samples collected at different distances from the vent and at different depths within the flow. The measurements presented here do not provide direct constraints on dynamics (beyond kinematics) and hence emplacement time scales: additional constraints are needed about crystal nucleation and growth rates, or other complementary measurements that constrain time scales.

Diffraction methods provide complementary results compared to optical ones and a combination of the two can be useful. Other methods that could be combined are X-ray or electron nanotomography (e.g., Miao et al. 2016; Midgley et al. 2007; Stampanoni et al. 2010) that have provided information about microstructures in multiphase shales (e.g., Kanitpanyacharoen et al. 2013). Voltolini et al. (2011) used X-ray microtomography to determine shape-preferred orientation of both crystals and vesicles in volcanic rocks and Arzilli et al. (2015) combined tomography with SEM-EBSD to characterize the growth of spherulites from melts. Such examples highlight how diffraction- and imaging-based approaches provide complementary results and might be combined successfully in the future to systematically explore what can be learned from microlites in lava flows.

# **Conclusions**

This study introduces a new method to determine the orientation distribution of microlites in an obsidian flow. With synchrotron X-ray diffraction, we can statistically quantify the orientation of clinopyroxene needles in the flow direction and andesine platelets in the flow plane. Microlites are aligned parallel to layering, confirming that layers are, in fact, flow bands. Based on model of rigid particles in a viscous fluid, we obtain strains 2–3.

Acknowledgements HRW acknowledges support from NSF (EAR-1343908) and DOE (DE-FG02-05ER15637) and access to beamline 11-ID-C at the Advanced Photon Source (APS) of Argonne National Laboratory. MM is supported by NSF 1724469. We appreciate help from Yang Ren at beamline ID-11 ID-C of APS and Tim Teague for sample preparation. Part of this research was done during a research leave at the University of Trento/Mesiano and HRW is grateful to Luca Lutterotti for the hospitality and guidance in data analysis, and also for access to the SEM and help from Lorena Maines. The authors thank the

editor, J. Donnelly-Nolan, and an anonymous reviewer for constructive comments and suggestions. Authors are listed in alphabetical order.

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