The quintet completed: The partitioning of sulfur between nominally volatile-free minerals and silicate melts

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ABSTRA CT

Magmatic systems are dominated by five volatiles, namely H₂O, CO₂, F, Cl, and S (the igneous quintet). Multiple studies have measured partitioning of four out of these five volatiles (H₂O, CO₂, F, and Cl) between nominally volatile-free minerals and melts, whereas the partitioning of sulfur is poorly known. To better constrain the behavior of sulfur in igneous systems we measured the partitioning of sulfur between clinopyroxene and silicate melts over a range of pressure, temperature, and melt composition from 0.8 to 1.2 GPa, 1000 to 1240 °C, and 49 to 66 wt% SiO₂ (13 measurements). Additionally, we determined the crystal-melt partitioning of sulfur for plagioclase (6 measurements), orthopyroxene (2 measurements), amphibole (2 measurements), and olivine (1 measurement) in some of these same run products. Experiments were performed at high and low oxygen fugacities, where sulfur in the melt is expected to be dominantly present as an S⁶⁺ or an S²⁻ species, respectively. When the partition coefficient is calculated as the total sulfur in the crystal divided by the total sulfur in the melt, the partition coefficient varies from 0.017 to 0.075 for clinopyroxene, from 0.036 to 0.229 for plagioclase, and is a maximum of 0.001 for olivine and of 0.003 for orthopyroxene. The variation in the total sulfur partition coefficient positively correlates with cation-oxygen bond lengths in the crystals; the measured partition coefficients increase in the order: olivine < orthopyroxene < clinopyroxene \le amphibole and plagioclase. At high oxygen fugacities in hydrous experiments, the clinopyroxene/melt partition coefficients for total sulfur are only approximately one-third of those measured in low oxygen fugacity, anhydrous experiments. However when the partition coefficient is calculated as total sulfur in the crystal divided by S²⁻ in the melt, the clinopyroxene/melt partition coefficients for experiments with melts between ~51 and 66 wt% SiO₂ can be described by a single mean value of 0.063 ± 0.010 (1 σ standard deviation about the mean). These two observations support the hypothesis that sulfur, as S²⁻, replaces oxygen in the crystal structure. The results of hydrous experiments at low oxygen fugacity and anhydrous experiments at high oxygen fugacity suggest that oxygen fugacity has a greater effect on sulfur partitioning than water. Although the total sulfur clinopyroxene-melt partition coefficients are affected by the Mg/(Mg+Fe) ratio of the crystal, partition coefficients calculated using S²⁻ in the melt display no clear dependence upon the Mg# of the clinopyroxene. Both the bulk and the S²⁻ partition coefficients appear unaffected by ^{IV}Al in the clinopyroxene structure. No effect of anorthite content nor of iron concentration in the crystal was seen in the data for plagioclase-melt partitioning. The data obtained for orthopyroxene and olivine were too few to establish any trends. The partition coefficients of total sulfur and S^{2-} between the crystals studied and silicate melts are typically lower than those of fluorine, higher than those of carbon, and similar to those of chlorine and hydrogen. These sulfur partition coefficients can be combined with analyses of volatiles in nominally volatile-free minerals and previously published partition coefficients of H₂O, C, F, and Cl to constrain the concentration of the igneous quintet, the five major volatiles in magmatic systems.

Keywords: Clinopyroxene/melt sulfur partitioning, equilibrium melts, magmatic volatiles, synchrotron micro X-ray fluorescence

INTRODU CTION

Five volatiles (the igneous quintet), H₂O, CO₂, F, S, and Cl,
dominate magmatic systems (e.g., Johnson et al. 1994; Symonds
et al. 1994). They play multiple roles in the character of magma-

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tism, from influencing the explosivity of volcanic eruptions to driving local or global environmental upheavals through poisoning of the ecosystems by acid fallout and net cooling or heating of the troposphere (e.g., Robock 2013). Quantitative estimations of volatile concentrations in magmas come primarily from two types of samples: glassy rinds of submarine pillow lavas and melt inclusions trapped in phenocrysts. The former is quenched at a

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1 high enough hydrostatic pressure to prevent efficient degassing 2 of volatiles (except C and H) from the melt, and the latter can be 3 trapped early enough in the magmatic system to be representative 4 of the pristine magmatic volatile concentrations. For instance, fo-5 cusing on sulfur, dredged mid-ocean ridge basalt (MORB) glasses were shown to contain 800-1300 ppm S (LeVoyer et al. 2014), 6 7 and melt inclusions enclosed in phenocrysts from {{auth: spell 8 out/define on first use}} OIBs up to 2100 ppm S (Azores, Rose-9 Koga et al. 2017), from flood basalts up to 1300 ppm (Laki, Ice-10 land; Hartley et al. 2017), and from arc magmas up to 2900 ppm (DeHoog et al. 2001; Johnson et al. 2009; Rusciutto et al. 2010). 11 12 These are concentrations typical of non-degassed melts (>800 13 ppm; Wallace and Edmonds 2011 and references therein) and are 14 highly informative of the composition and oxidation state of the 15 mantle source of these magmas. However, data from such kinds 16 of samples pertain mostly to recent magmatic manifestations, 17 while working with subaerial and/or ancient eruptions, magmatic 18 volatiles quantification gets more challenging. Subaerial eruptions 19 tend to efficiently degas their volatile budget, e.g., a maximum of 20 ~150 ppm S was measured in subaerial matrix glasses (Wallace 21 and Edmonds 2011). Ancient volcanics tend to alter, making it 22 hard to find either fresh matrix glasses to analyze or preserved 23 melt inclusions. Valuable exceptions exist, with a precious few 24 melt inclusions successfully analyzed from flood basalts and 25 subvolcanic rocks from Large Igneous Provinces (e.g., Self et al. 26 2008, Deccan Traps; Sibik et al. 2015; Black et al. 2012, Siberian 27 Traps). In the absence of melt inclusions, volatile concentrations 28 in melts may be determined by the combination of the measure-29 ment of volatile concentrations in natural minerals-typically at 30 the parts-per-million level in nominally volatile-free minerals 31 (NVFMs) such as olivine, orthopyroxene, clinopyroxene, and 32 plagioclase, with experimentally determined partition coefficients 33 between these crystals and melts.

34 Only a few, recent studies report measurements of the parti-35 tioning of H₂O (or H), C, F, and Cl between basaltic melts and 36 olivine, orthopyroxene, clinopyroxene, and plagioclase (cf. La 37 Tourrette 1995; Hauri et al. 2006; Guggino 2012; Hamada et al. 38 2013; Callegaro et al. 2014; Rosenthal et al. 2015; Lloyd et al. 39 2016; Dalou et al. 2012, 2014; Bénard et al. 2017; Urann et al. 40 2017; Beyer et al. 2012, 2016; cf. review by Webster et al. 2018). 41 Of these studies, only a small subset included S measurements 42 (Hauri et al. 2006; Callegaro et al. 2014; Rosenthal et al. 2015; 43 Lloyd et al. 2016), but none was focused specifically on sulfur 44 partitioning, except that of Callegaro et al. (2014), whose results 45 we include in the present contribution for discussion.

46 To better use NVFMs as probes of melt volatile concentrations, 47 additional measurements of partition coefficients are needed. 48 In addition, the influence of magmatic variables such as melt 49 composition (including water concentration), crystal structure 50 and chemistry, and oxygen fugacity on partitioning needs to be 51 examined, particularly for sulfur because of its change from an 52 S²⁻ to an S⁶⁺ species in the melt with increasing oxygen fugacity 53 (Wilke et al. 2011; Moretti and Baker 2011{{auth: not listed in 54 **Ref list please add or fix**}). Here we present the results obtained 55 from piston-cylinder experiments designed to investigate the par-56 titioning of sulfur between NVFMs and silicate melts as a function 57 of these variables and compare these results to the partitioning 58 behavior of the other volatiles constituting the igneous quintet. 59

EXPERIMENTAL TECHNIQUES

Experiments on basaltic compositions were performed with a powdered MORB and a powdered basalt (AN-31) of the Central Atlantic Magmatic Province (CAMP), collected from a lava flow in Morocco (Marzoli et al. 2019). Two intermediate-composition glasses were also used as starting materials; one andesitic glass (AT-29D) was made from a mixture of 95% Aleutian andesite and 5% diopside glass, added to facilitate clinopyroxene crystallization, and the other (AT-150) was a synthetic dacitic glass, whose composition was similar to a natural Aleutian rock. Starting samples AN-31, AT-29D, and AT-150 were enriched in sulfur through the addition of finely ground pyrrhotite and dry mixed in a horizontal rotary mill to homogenize them. Electron microprobe analyses of super-liquidus glasses of these starting materials are provided in Table 1. The MORB contained approximately 800 ppm sulfur, AN-31 approximately 900 ppm S, and AT-29D and AT-150 both approximately 300 ppm S (Table 1). Starting samples were ground by hand in a mortar to less than 50 µm in size and stored in a drying oven at ~120 °C before experiments.

Low oxygen fugacity experiments were performed in graphite-lined platinum capsules. These double capsules minimize iron loss and create oxygen fugacity conditions approximately 1.5 to 2 log units below the fayalite-quartz-magnetite buffer (FMQ), or FMQ-2 (e.g., Médard et al. 2008); at these conditions sulfur dissolved in the melt exists in a sulfide complex, S2- (Wilke et al. 2011). Capsules for anhydrous experiments were loaded with starting materials (~10 mg) and dried in the oven before welding. Hydrous conditions were achieved by first adding liquid water and then the other starting materials before welding with the capsule immersed in water to keep the metal cool and prevent volatile loss during welding. All hydrous capsules were heated at 110 °C for at least 2 h to test the weld, and any capsules whose weight changed either during welding or after heating were discarded.

High oxygen fugacity experiments were performed in Au₇₅Pd₂₅ capsules. These capsules mitigate iron loss and in our piston-cylinder assembly create oxygen fugacities approximately 1 to 2 log units above FMQ, i.e., FMQ+1 to FMQ+2 (Dalpé and Baker 2000; Liu et al. 2007); at these f_{02} values much of the sulfur in the melt is present as a sulfate complex, S^{6+} (Wilke et al. 2011). The loading, drying, and heating procedures for these capsules were identical to those used for the graphite-in-Pt capsules.

Experiments were performed in a piston-cylinder using NaCl-pyrex-crushable alumina assemblies following the techniques of Baker (2004). Hydrous experimental capsules were surrounded by pyrophyllite or Al(OH)3 powder to reduce water loss (Freda et al. 2001). Experiments were simultaneously heated and pressurized to conditions above the liquidus and held at those conditions for 1 to 2 h to homogenize the melt and destroy any crystals in the starting material before cooling to subliquidus conditions at a rate of 1 °C per minute. Upon reaching the desired, sub-liquidus temperature the experiments were held at that temperature for a duration of approximately 24 h, allowing crystal growth (Table 2). This duration has previously been shown sufficient for the andesite AT-29 and the MORB basalt to reach equilibrium conditions at anhydrous conditions and similar temperatures and pressures, even with residual melt compositions as rich in silica (67 wt% SiO₂) as those in this study (Baker and Eggler 1987; Baker 2008).

ANALYTICAL TECHNIQUES

Run-product phases were analyzed for major element concentrations on a JEOL 8900 electron microprobe (McGill University). We used an accelerating voltage of 15 kV, a beam current of 20 nA, and a beam diameter of 20 μm for the glasses and 1 µm for the crystals. We used 20 s counting time for the peaks of major elements and 200 s for S analyses of the quenched melts; background counting times

TABLE 1. Compositions of starting materials based upon microprobe analysis of super-liquidus glasses

	MORB basalt	AN-31 CAMP tholeiite	AT-29D andesite	AT-150 dacite							
SiO ₂ (wt%	6) 49.5 (0.70)	50.2 (0.18)	55.99 (0.58)	63.22 (0.26)							
TiO ₂	1.28 (0.06)	1.17 (0.04)	0.86 (0.04)	0.54 (0.03)							
AI_2O_3	15.4 (0.12)	11.1 (0.06)	16.33 (0.08)	17.82 (0.08)							
FeO*	9.37 (0.27)	11.4 (0.17)	7.86 (0.37)	4.81 (0.09)							
MnO	0.18 (0.04)	0.18 (0.05)	0.18 (0.02)	0.01 (0.01)							
MgO	8.89 (0.11)	12.8 (0.12)	4.00 (0.16)	1.73 (0.03)							
CaO	11.7 (0.16)	9.07 (0.11)	8.19 (0.25)	5.32 (0.08)							
Na₂O	2.4 (0.08)	1.63 (0.04)	3.48 (0.07)	4.30 (0.05)							
K ₂ O	0.1 (0.01)	0.66 (0.03)	1.94 (0.07)	1.76 (0.03)							
P_2O_5	0.11 (0.02)	0.12 (0.02)	0.23 (0.01)	0.01 (0.01)							
S (ppm)	842 (39)	911 (36)	366 (67)	263 (23)							
Total	98.93	98.33	99.07	99.46							
Note: Standard deviations in parentheses.											

(pt.	Р	Т	Time	f_{O_2}	H ₂ O	Phase	n ^d	S	S ⁶⁺ /S _{tot} ^e	D	D
	(GPa)	(°C) ^a	h⁵	ΔFMQ ^c	(wt%)			(ppm)		(S tot) ^f	(S ^{2–}) ^g
				Starting m	aterial MORB -	basaltic glas	s				
B2012-36	1.0	1350/1240	2/20.1	-2	n.a. ^h	glass	12	1032 (84)	0		
						Срх	[8]	29 (7)		0.028	0.028
						PI	[1]	37 (1)		0.036	0.036
RB2012-38	1.2	1350/1240	2/20	-2	n.a.	glass	12	1090 (27)	0		
						Срх	[8]	25 (11)		0.023	0.023
52014 12	1.0	1250/1240	2/24	1.0			10	017 (20)	0.72		
S2014-13	1.0	1350/1240	2/24	1.8	n.a.	glass	10	917 (36)	0.73	0.017	0.065
						Срх	[4]	16 (13)		0.017	0.065
RB2015-1	1.0	1150/1060	2/24	1.5	7.6 (0.9)	glass	18 [3]	1156 (62)	0.63		
1020151	1.0	1150/1000	2/27	1.5	7.0 (0.2)	Cpx	[5]	21 (7)	0.05	0.018	0.049
			Startin	a material AN	-31 - basaltic gl					01010	010 15
RB2012-29	0.8	1350/1240	2/24	-2	n.a.	glass	18	933 (28)	0		
						Орх	[9]	3 (2)		0.003	0.003
						Ö	[2]	1 (0.2)		0.001	0.001
RB2012-35	1.0	1350/1240	2/20.1	-2	n.a.	glass	12	1116 (22)	0		
						Срх	[6]	29 (9)		0.026	0.026
RB2012-37	1.2	1350/1240	2/20	-2	n.a.	glass	12	1096 (19)	0		
						Срх	[8]	31 (9)		0.028	0.028
S2014-14	1.0	1350/1240	2/24	1.7	n.a.	glass	12	1037 (42)	0.73		
						Орх	[3]	2 (1)		0.002	0.007
				Starting ma	terial AT-29D -	andesitic gla	ISS				
S2014-9	0.8	1300/1160	2/24	-2	n.a.	glass	10	571 (60)	0		
						Срх	[4]	33 (7)		0.058	0.058
						PI	[1]	57 (8)		0.100	0.100
S2014-5	0.8	1300/1140	1/24	-2	n.a.	glass	11	717 (145)	0		
						Срх	[3]	54 (22)		0.075	0.075
						PI	[4]	47 (13)		0.065	0.065
CS2014-3	0.8	1300/1118	1/24	-2	n.a.	glass	22	551 (141)	0		
						Срх	[2]	38 (11)		0.069	0.069
						PI	[4]	44 (20)		0.080	0.080
S2014-30	0.8	1150/1000	2/24	-2	1.1 (0.3)	glass	15	689 (181)	0		
						PI	[2]	61 (36)		0.089	0.089
						Amp	[5]	87 (58)		0.127	0.127
DRB2015-2	1.0	1150/1060	2/24	1.5	6.3 (0.3)	glass	19	742 (55)	0.62		
						Срх	[6]	13 (7)		0.018	0.046
S2014-19	0.8	1150/1000	2/24	0.9	11.2 (0.5)	glass	14	580 (80)	0.33		
						Срх	[4]	25 (12)		0.043	0.064
					aterial AT-150						
S2014-31	0.8	1150/1000	2/24	-2	1.1 (0.3)	glass	10	292 (35)	0		
						Срх	[2]	21 (8)		0.072	0.072
62014 22	0.0	1150/1000	2/2 4			PI	[4]	67 (20)	0.11	0.229	0.229
S2014-20	0.8	1150/1000	2/24	1.1	5.7 (0.8)	glass	10	237 (44)	0.41	0.030	
						Срх	[1]	9(1)		0.038	0.064
						PI	[2]	93 (10)		0.393	0.666
						Amp	[2]	29 (14)		0.123	0.208

were all one-half of those on the peaks. The lower detection limit of S in glasses was approximately 100 ppm. Synthetic pyrrhotite was used as the sulfur standard for analyses of experiments at low oxygen fugacity, whereas barite was used for experiments at high oxygen fugacity. The standards used for glass analyses were a basaltic glass, VG-A99 (Jarosewich et al. 1980), for Na, Al, Fe, Si, Mg, Ca, and Ti; a rhyolitic glass for K; a spessartine for Mn; and a fluorapatite for P. Basaltic glass standards VG-2 (1410 ppm S) and VG-A99 (125 ppm S) were repeatedly analyzed to ensure the accuracy of our analyses (cf. Liu et al. 2007; Fortin et al. 2015). The standards used for mafic crystal analyzes were diopside for Ca and Mg and Olivine for Fe; feldspars were used as Na, K, and Al standards for all other elements were the same as those used for glass analyses.

by the sulfur peak shift method pioneered by Carroll and Rutherford (1988). The wavelength of the sulfur peak in the glass of each high oxygen fugacity experiment was found using the electron microprobe by scanning the peak of 20 different spots in the glass and then summing the scans to increase the peak-to-background ratio. A 10 µm diameter beam with a 15 kV potential and 20 nA current was used to minimize possible sulfur oxidation during analysis. This measured peak position was then compared to the sulfur peak positions measured in sphalerite, the sulfide standard, and barite, the sulfate standard, to determine the fraction of sulfur dissolved as sulfate in the melt (Carroll and Rutherford 1988). From this sulfate fraction, the log of the oxygen fugacity relative to the FMQ buffer was calculated following Wilke et al. (2011).

The oxygen fugacity in all high oxygen fugacity experiments was determined

Water was measured in the run-product glasses of all hydrous experiments by 57 Raman spectroscopy (see Supplementary¹ Fig. S1), following Fortin et al. (2015) 58

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TABLE 3. Electron microprobe analyses of the glass and crystal phases obtained from the experiments

	Expt.	Phase	nª	SIO ₂	TiO ₂	Al ₂ O ₃	FeO*	MnO	MgO	CaO
Starting	DRB2012-36	glass	12	48.98 (0.3)	1.48 (0.11)	16.48 (0.12)	10.71 (0.23)	0.19 (0.02)	7.18 (0.09)	10.38 (0.07
material MORB		Срх	9	52.07 (1.41)	0.42 (0.17)	5.55 (2.06)	7.25 (1.)	0.19 (0.04)	19.78 (2.46)	13.68 (2.56
		PI	1	53.44 (0.53) ^b	n.a.c	28.81 (0.29)	0.33 (0.03)	n.a.	0.16 (0.02)	12.14 (0.12
	DRB2012-38	glass	12	49.33 (0.14)	1.53 (0.05)	17.21 (0.11)	10.54 (0.12)	0.17 (0.03)	6.42 (0.06)	9.80 (0.05
		Срх	10	50.18 (0.37)	0.61 (0.11)	8.36 (0.53)	6.95 (0.59)	0.18 (0.03)	17.10 (0.57)	15.27 (0.93
	CS2014-13	glass	10	50.70 (0.22)	1.39 (0.04)	16.57 (0.24)	8.54 (0.16)	0.18 (0.02)	8.46 (0.2)	10.93 (0.08
		Срх	28	49.27 (0.72)	0.63 (0.18)	8.78 (1.39)	6.90 (0.42)	0.18 (0.03)	15.93 (1.26)	17.01 (1.04
	DRB2015-1	glass	18	51.88 (0.25)	0.72 (0.03)	18.23 (0.12)	4.03 (0.12)	0.16 (0.03)	5.71 (0.15)	8.35 (0.17
		Срх	52	48.04 (2.09)	0.68 (0.29)	6.58 (1.77)	7.46 (1.09)	n.a.	15.76 (1.77)	20.17 (0.86
Starting	DRB2012-29	glass	18	50.83 (0.48)	1.29 (0.07)	13.28 (0.16)	10.40 (0.21)	0.18 (0.05)	8.80 (0.13)	10.50 (0.18
material AN-31		Орх	42	55.52 (0.55)	0.18 (0.06)	2.04 (0.59)	8.65 (0.48)	0.17 (0.03)	30.89 (1.34)	2.07 (0.53
		OI	19	39.66 (0.23)	0.01 (0.02)	0.03 (0.01)	14.53 (0.44)	0.19 (0.02)	45.35 (0.48)	0.24 (0.02
	DRB2012-35	glass	12	48.85 (0.24)	1.52 (0.03)	14.21 (0.1)	10.84 (0.12)	0.18 (0.03)	8.04 (0.07)	10.35 (0.05
		Срх	10	55.31 (0.41)	0.18 (0.05)	2.22 (0.28)	9.82 (0.39)	0.18 (0.03)	28.91 (1.5)	3.33 (1.54
	DRB2012-37	glass	12	49.10 (0.23)	1.40 (0.04)	14.14 (0.1)	11.11 (0.12)	0.17 (0.04)	7.92 (0.08)	10.09 (0.06
		Срх	10	54.91 (0.29)	0.16 (0.03)	2.60 (0.31)	10.10 (0.52)	0.20 (0.02)	28.72 (1.2)	3.20 (1.26
	CS2014-14	glass	12	50.22 (0.24)	1.25 (0.03)	12.76 (0.15)	10.62 (0.18)	0.18 (0.03)	10.00 (0.25)	10.02 (0.19
		Орх	31	54.32 (1.13)	0.14 (0.05)	3.47 (0.56)	7.45 (0.54)	0.19 (0.03)	31.01 (0.73)	1.96 (0.25
Starting	CS2014-9	glass	10	57.67 (0.36)	1.23 (0.06)	15.27 (0.13)	9.78 (0.25)	0.20 (0.03)	2.77 (0.09)	5.95 (0.12
material AT-29D		Срх	16	51.23 (0.9)	0.50 (0.15)	3.68 (0.96)	12.55 (1.53)	0.40 (0.05)	16.76 (1.53)	13.54 (2.74
		PI	20	58.06 (0.47)	n.a.	25.82 (0.26)	0.44 (0.07)	n.a.	0.10 (0.02)	8.57 (0.33
	CS2014-5	glass	11	56.67 (0.29)	1.41 (0.04)	14.55 (0.14)	10.55 (0.33)	0.20 (0.02)	2.69 (0.07)	5.90 (0.12
		Срх	18	51.53 (0.5)	0.55 (0.13)	3.19 (0.76)	12.69 (1.51)	0.40 (0.05)	16.91 (1.27)	13.57 (2.39
		PI	22	57.90 (0.32)	n.a.	26.09 (0.25)	0.47 (0.11)	n.a.	0.12 (0.04)	8.86 (0.21
	CS2014-3	glass	22	56.28 (0.3)	1.07 (0.14)	15.35 (0.47)	9.34 (0.45)	0.20 (0.03)	3.53 (0.26)	6.94 (0.34
		Срх	19	51.67 (0.42)	0.62 (0.12)	3.89 (0.67)	10.06 (1.28)	0.33 (0.05)	16.65 (1.19)	16.07 (2.09
		PI	13	57.34 (0.67)	n.a.	26.20 (0.35)	0.41 (0.14)	n.a.	0.13 (0.04)	9.19 (0.29
	CS2014-30	glass	15	59.49 (1.07)	2.04 (0.18)	13.59 (0.35)	10.32 (1.02)	0.18 (0.04)	1.05 (0.08)	3.93 (0.32
		PI	17	58.55 (0.83)	n.a.	25.59 (0.68)	0.46 (0.07)	n.a.	0.05 (0.01)	8.13 (0.7)
		Amp	5	55.67 (0.91)	1.22 (0.51)	10.50 (3.71)	11.70 (3.71)	0.32 (0.08)	6.35 (1.87)	8.36 (2.42
	DRB2015-2	glass	19	55.76 (0.43)	0.54 (0.04)	16.39 (0.15)	3.44 (0.09)	0.16 (0.02)	2.92 (0.06)	6.57 (0.09
		Срх	29	44.38 (0.44)	0.85 (0.14)	8.62 (0.36)	10.81 (0.47)	n.a.	11.58 (0.36)	21.78 (0.15
	CS2014-19	glass	14	57.84 (0.33)	0.57 (0.05)	17.13 (0.14)	3.37 (0.12)	0.14 (0.02)	2.44 (0.1)	5.67 (0.12
		Срх	17	44.94 (1.08)	1.13 (0.21)	8.88 (0.52)	10.03 (1.17)	0.25 (0.03)	11.50 (0.71)	21.36 (0.26
Starting	CS2014-31	glass	10	66.11 (0.8)	0.90 (0.07)	13.74 (0.58)	6.66 (0.43)	n.a.	1.00 (0.19)	3.43 (0.23
material AT-150		Срх	10	52.64 (0.39)	0.32 (0.05)	2.27 (0.4)	20.16 (1.32)	n.a.	20.08 (1.09)	4.15 (0.42
		PI	25	58.99 (1.14)	n.a.	25.74 (0.72)	0.41 (0.14)	n.a.	0.08 (0.02)	8.48 (0.57
	CS2014-20	glass	10	61.34 (1.24)	0.46 (0.07)	16.78 (0.7)	3.27 (0.27)	n.a.	1.44 (0.14)	4.47 (0.34
		Срх	4	45.34 (1.39)	0.66 (0.14)	10.03 (1.51)	9.69 (0.67)	n.a.	10.61 (0.71)	21.84 (0.08
		Ρl	4	57.61 (0.39)	n.a.	25.87 (0.11)	0.90 (0.09)	n.a.	0.07 (0.01)	9.05 (0.09
		Amp	3	55.23 (2.87)	0.51 (0.18)	14.14 (1.3)	5.79 (1.61)	n.a.	5.39 (1.71)	13.30 (2.59
		Opq	1	0.00	12.14 (0.12)	2.14 (0.02)	73.97 (0.74)	n.a.	1.42 (0.01)	0.05 (0.01

Notes: Totals for the glass phases do not include SO₃ wt%, nor H₂O and Cl wt% (where analyzed).

³⁴ ^a Number of analyses by electon microprobe for major and minor elements.

35 ^bWhere only a single analysis is available, the uncertaintly is the analytical uncertainty from counting statistics.

36 ^c n.a. = not analyzed; Ol = olivine; Cpx = clinopyroxene; Opx = orthopyroxene; Pl = plagioclase; Amp = amphibole; Opq = opaque phase.

(Extended on next page)

using as standards a set of andesitic and basaltic glasses previously analyzed by ion
 microprobe (Fortin et al. 2015).

Sulfur in the crystals was measured by synchrotron X-ray microfluorescence 40 (SXRF) on beamline I18 at the Diamond Light Source synchrotron, U.K. (Mos-41 selmans et al. 2009). Synchrotron X-ray fluorescence analyses of silicon and sulfur 42 concentrations in the crystals were performed in a helium atmosphere using a 3 keV beam focused to $6 \times 6 \,\mu\text{m}$ by a pair of Kirkpatrick-Baez mirrors, and the fluorescence 43 spectra of the samples were measured with a Vortex silicon drift detector. Sulfur 44 concentrations were determined from the spectra by PyMca (Solé et al. 2007) using 45 the silicon concentration of the minerals as the internal reference for quantification. 46 The SXRF analytical technique for sulfur was tested by analysis of two in-house clinopyroxene crystal standards. The crystals were gem-quality DeKalb diopside 47 (USNM #R18685) and F-14 clinopyroxene from Frosty Peak, Alaska, U.S.A. 48 (collected by D.R.B.). Bulk analyses of these crystals for sulfur were made at the 49 Saskatchewan Research Council Geoanalytical Laboratories using a LECO induc-50 tion furnace carbon and sulfur analyzer (https://www.src.sk.ca/labs/geoanalytical-51 laboratories). Three different aliquots of DeKalb diopside yielded a mean sulfur concentration of 32 ± 15 ppm (1 St.dev.), and one aliquot of the F-14 clinopyroxene 52 contained 32 ppm sulfur (Supplementary¹ Table S1). Aliquots of international 53 standards BHVO-2, JP-1, and JB-2 were analyzed in the same analytical batch, 54 and the results reproduced the published recommended values for these standards 55 (Erdman et al. 2014). Eight SXRF analyses of DeKalb diopside crystals produced a mean sulfur concentration of 32 ± 18 ppm and six analyses of F-14 clinopyroxene 56 crystals yielded a concentration of 22 ± 9 ppm sulfur. The agreement of bulk sulfur 57 analyses and the average of the SXRF analyses is within 10 uncertainty for DeKalb 58 and just outside the 10 uncertainty for F-14, although both crystals display heterogeneity in sulfur concentrations. Analyses of the same crystals were attempted by ion microprobe (CAMECA ims 1280 at Nordsim Laboratory, Stockholm Natural History Museum, Sweden) using glass standards for calibration and yielded sulfur concentrations in DeKalb and F-14 less than 1 ppm. Because of the significant difference between the bulk sulfur analyses and those obtained by ion microprobe, we did not use the latter in this study. We suggest that the difference between ion microprobe and SXRF analyses may be due to inadequate standards for ion probe sulfur analysis in mafic crystals.

We calculated a detection limit of approximately 1 ppm for our SXRF analyses by two different methods (p. 446, Goldstein et al. 2003; Rousseau 2001). Based upon a relative uncertainty of 10% in our electron microprobe analyses as well as 10% uncertainty seen in our peak fitting areas, we calculate through error propagation (Rousseau 2001) an analytical uncertainty of 14% relative for samples with 6 ppm and greater. At 2 ppm sulfur, the uncertainty in the peak fitting areas reaches 37%, and the analytical uncertainty becomes 38% relative.

RESULTS

The experimental conditions and the analyses of the run product phases are presented in Tables 2 and 3. Experimental conditions were chosen for the crystallization of clinopyroxene, but plagioclase, low-calcium pyroxene, olivine, amphibole, and opaque phases were also present in selected experiments. Iron sulfide phases, either quenched from an immiscible liquid or present as a stable sulfide crystal, were observed in three experiments

TABLE 3.—EXTENDED

	Expt.	Phase	nª	Na₂O	K ₂ O	P_2O_5	Total
Starting	DRB2012-36	glass	12	2.92 (0.06)	0.13 (0.01)	0.12 (0.07)	98.57
material		Срх	9	0.41 (0.1)	0.02 (0.01)	0.04 (0.03)	99.40
MORB		PI	1	4.29 (0.43)	0.05 (0.01)	n.a.	99.35
	DRB2012-38	glass	12	3.13 (0.02)	0.13 (0.01)	0.16 (0.05)	98.42
		Срх	10	0.62 (0.07)	0.02 (0.01)	0.06 (0.08)	99.34
	CS2014-13	glass	10	2.65 (0.08)	0.11 (0.01)	0.13 (0.01)	99.65
		Срх	28	0.63 (0.15)	0.02 (0.02)	n.a.	99.35
	DRB2015-1	glass	18	3.06 (0.09)	0.13 (0.01)	0.15 (0.02)	92.40
		Срх	52	0.50 (0.06)	0.02 (0.01)	n.a.	99.21
Starting	DRB2012-29	glass	18	1.98 (0.05)	0.81 (0.03)	0.15 (0.02)	98.22
material		Орх	42	0.05 (0.06)	0.02 (0.03)	n.a.	99.60
AN-31		OI	19	n.a.	0.02 (0.01)	0.04 (0.02)	100.08
	DRB2012-35	glass	12	2.25 (0.04)	0.91 (0.02)	0.15 (0.07)	97.31
		Срх	10	0.09 (0.04)	0.02 (0.01)	0.05 (0.06)	100.11
	DRB2012-37	glass	12	2.28 (0.03)	0.94 (0.02)	0.16 (0.08)	97.31
		Срх	10	0.09 (0.04)	0.02 (0.01)	0.06 (0.04)	100.07
	CS2014-14	glass	12	1.94 (0.03)	0.78 (0.02)	0.15 (0.01)	97.91
		Орх	31	0.06 (0.02)	0.02 (0.01)	n.a.	98.62
Starting	CS2014-9	glass	10	3.42 (0.04)	2.80 (0.06)	0.36 (0.02)	99.45
material AT-29D		Срх	16	0.38 (0.1)	0.03 (0.01)	n.a.	99.07
		ΡI	20	5.74 (0.12)	1.16 (0.13)	n.a.	99.90
	CS2014-5	glass	11	3.39 (0.06)	2.92 (0.04)	0.41 (0.03)	98.73
		Срх	18	0.32 (0.08)	0.03 (0.01)	n.a.	99.19
		ΡI	22	5.73 (0.08)	1.01 (0.09)	n.a.	100.18
	CS2014-3	glass	22	3.42 (0.07)	2.43 (0.14)	0.29 (0.05)	98.85
		Срх	19	0.38 (0.06)	0.03 (0.03)	n.a.	99.70
		ΡI	13	5.56 (0.14)	0.88 (0.1)	n.a.	99.71
	CS2014-30	glass	15	3.29 (0.1)	4.24 (0.21)	0.67 (0.06)	98.79
		PI	17	5.99 (0.23)	1.19 (0.33)	n.a.	99.95
		Amp	5	1.96 (0.79)	2.04 (0.77)	n.a.	98.11
	DRB2015-2	glass	19	3.54 (0.06)	2.13 (0.04)	0.25 (0.02)	91.72
		Срх	29	0.79 (0.04)	0.02 (0.01)	n.a.	98.83
	CS2014-19	glass	14	3.20 (0.1)	2.05 (0.06)	0.25 (0.01)	92.68
		Срх	17	0.59 (0.04)	0.03 (0.03)	n.a.	98.70
Starting	CS2014-31	glass	10	3.65 (0.1)	2.69 (0.1)	n.a.	98.19
material		Срх	10	0.10 (0.05)	0.04 (0.04)	n.a.	99.78
AT-150		Ρİ	25	6.15 (0.25)	0.49 (0.11)	n.a.	100.33
	CS2014-20	glass	10	4.62 (0.08)	1.64 (0.03)	n.a.	94.04
		Срх	4	0.82 (0.04)	0.02 (0.01)	n.a.	99.02
		PI	4	6.09 (0.12)	0.29 (0.01)	n.a.	99.87
		Amp	3	1.90 (0.22)	0.97 (0.37)	n.a.	97.23
		Opq	1	0.00	0.031 (0.01)	n.a.	89.74

(CS2014-3, -5, and -30), but in each experiment the modal proportion of sulfides was less than 5%. The typical morphologies of the crystals in low oxygen-fugacity experiments with basaltic compositions were euhedral to subhedral (Fig. 1a), whereas crystals in experiments with basaltic compositions at high oxygen fugacity and experiments with andesitic and dacitic bulk compositions were typically subhedral to anhedral (Fig. 1b). Most crystals were approximately 100 µm in their minimum dimension; however, some were as low as 25 µm and some as high as 400 µm across. All the crystals were significantly larger than the beam sizes of all applied analytical techniques ($6-2 \mu m$). The smallest crystals allowed for only one analysis each, while several analyses (for instance at the core and at the rim) were performed on the largest crystals. In all run products, the rims (within 10 µm of the melt) of the most euhedral crystals were used for the measurement of major element and sulfur concentrations, because some crystals displayed major element compositional zoning. Although we tried to analyze at least five crystals in each experiment, in some cases, this was impossible. Additionally, some of our analyses contained anomalously high sulfur concentrations indicative of an analysis of a mixture of crystal+glass, marked also by a higher Cl concentration. Such obviously incorrect analyses were removed from those used to calculate the mean sulfur concentrations of the crystals. In one case (CS2014-20) only one clinopyroxene

analysis was deemed acceptable, but for most experiments, 1 three or more analyses were used for the calculation of the mean 2 and standard deviation (see Table 3). The compositions of the 3 melts and crystals obtained from all successful experiments are 4 reported in Table 3. The many anhydrous experiments at high 5 oxygen fugacity that produced crystals too small for microbeam 6 analysis were not considered. Notably, the melt compositions, 7 sulfur concentrations in clinopyroxenes, and the resulting parti-8 tion coefficients for experiments DRB2012-29, -35, -36, -37, and 9 -38 were previously presented in Callegaro et al. (2014) and are 10 included in this study for comparison.

Crystal-melt equilibrium was assessed by comparing 12 measured Fe-Mg partitioning between crystals and melts with 13 previous studies. With one exception, the clinopyroxene-melt 14 Fe-Mg partitioning of the low oxygen fugacity experiments was 15 within 25 relative percent of values calculated following Putirka 16 (1999). Experiment CS2014-31, performed at 1000 °C, is an 17 exception; we do not attribute its difference to disequilibrium, 18

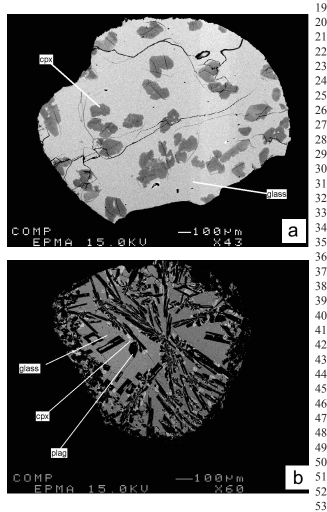


FIGURE 1. Backscattered electron images of two representative run products. (**a**) Basaltic run product of experiment DRB2012-38 (1.2 GPa, 1240 °C, anhydrous, low oxygen fugacity) containing clinopyroxene (Cpx) and glass. (**b**) Andesitic run product of experiment CS2014-5 (0.8 GPa, 1140 °C, anhydrous, low oxygen fugacity) containing clinopyroxene, plagioclase (Plag) and glass; the scale bar in this image is 200 µm. 59

1 but to the fact that the experimental temperature is below the 2 calibration range of Putirka's study, and the melt composition is 3 richer in silica (66 wt%) than the melts used in Putirka (1999) to 4 calibrate Fe-Mg partitioning equations. The Fe-Mg partitioning 5 for both orthopyroxene and olivine in the low oxygen fugacity 6 experiments of this study were similar to those previously mea-7 sured for similar compositions at similar temperatures (Baker 8 and Eggler 1987). On the other hand, the high oxygen fugacity 9 experiments displayed partition coefficients significantly higher 10 than expected from Putirka's (1999) calibration. We attribute this 11 difference to the effect of oxygen fugacity on the ferric/ferrous 12 ratio in both the melt and the crystal, a variable not included 13 in Putirka's (1999) equations describing Fe-Mg partitioning 14 between clinopyroxene and melt.

15 No intracrystalline heterogeneity was observed for sulfur, but 16 many crystals were so small that only one analysis was made for 17 each. Boyd's homogeneity index was used to assess the homogene-18 ity of sulfur in the analyzed clinopyroxenes. According to Boyd 19 et al. (1967), if the ratio of the relative standard deviation (based 20 upon multiple analyses) to the relative uncertainty inherent in 21 the analyses (e.g., the counting statistics) is above 3, it should be 22 taken as highly suggestive of the presence of inhomogeneity in 23 the material (see also Harries 2014). Potts et al. (1983) suggested 24 that a higher value, above 4, should be considered as the threshold 25 between homogeneity and heterogeneity. We calculated Boyd's 26 homogeneity index using the relative standard deviation about 27 the mean of multiple S analyses on different crystals (Table 2) 28 and a relative analytical uncertainty for concentrations greater 29 than 6 ppm of 14%, as calculated in the Analytical Techniques. 30 Of the 13 clinopyroxene samples {{auth: okay?}} analyzed, we 31 find that 4 display a Boyd homogeneity index greater than 3, but 32 only 1 is greater than 4 (CS2014-13, with a value of 5.8). Thus, 33 despite the large standard deviations seen for some analyses, the 34 intercrystalline sulfur concentrations from each experiment do not 35 display significant evidence of heterogeneity. Although CS2014-36 13 displays evidence of heterogeneity, we retain it because we 37 believe that the mean sulfur concentration in the clinopyroxenes 38 is a reliable estimate.

39 The fraction of sulfur as sulfate (as measured at the electron 40 microprobe by the sulfur peak shift) in the high oxygen fugacity experiments varied from 0.33 to 0.73, and their calculated oxygen 41 42 fugacities range from FMQ+0.8 to FMQ+1.8 (Table 2). These high 43 oxygen fugacities are consistent with the presence of Fe-Ti oxide 44 minerals (e.g., CS2014-20 in Table 2) only in the high oxygen fu-45 gacity experiments. Additionally, only these experiments produced 46 clinopyroxenes whose mineral formula calculations indicated the 47 presence of ferric iron. Three low-oxygen-fugacity experiments 48 (CS2014-9, -30, and -31) were also measured and found to have all 49 sulfur dissolved as sulfide, consistent with previous measurements 50 of sulfide speciation in anhydrous and hydrous melts in graphite-51 lined Pt capsules (Fortin et al. 2015). Although the oxygen fugacity 52 cannot be calculated for these low oxygen fugacity experiments, 53 the lack of measurable sulfate indicates oxygen fugacities at, or 54 below, the FMQ buffer, which following Fortin et al. (2015) we 55 estimate as FMQ-2 in Table 2. The oxygen fugacity may possibly 56 be lower, but its minimum value is constrained by the lack of 57 metallic iron in the run products, i.e., the oxygen fugacity is above 58 FMQ-4 at the conditions studied. 59

Sulfur partitioning between clinopyroxene and melt

The mean concentration of sulfur in the clinopyroxenes varied from a minimum of 9 to a maximum of 54 ppm (Table 2), and the corresponding partition coefficients varied from 0.017 to 0.0750. Total sulfur partition coefficients are calculated by dividing the sulfur concentration in the crystal by the total sulfur concentration in the coexisting melt and are plotted as a function of the SiO₂ concentration in the melt in Figure 2a. This figure demonstrates that with the exception of one hydrous experiment with a high-SiO₂ melt (CS2014-31), the partition coefficients can be separated into low- f_{0_2} and high- f_{0_2} trends. Based upon the major element composition of the clinopyroxenes, approximately half were augitic and the other half pigeonitic. There appear to be no significant differences in the crystal-melt sulfur partition coefficients of augitic and pigeonitic clinopyroxenes; however, as discussed below, the Fe/Mg ratio of the clinopyroxene appears to affect the total sulfur partition coefficient.

Five hydrous, clinopyroxene-bearing experiments were performed. The water concentrations in the melts varied from 1.1 to 11.2 wt% H₂O (Table 2). Comparison of the one hydrous experiment producing a basaltic melt with 7.6 wt% H₂O (DRB2015-1) with the anhydrous experiment at the same pressure and oxygen fugacity (CS2014-13) produced similar total sulfur partition coefficients (Fig. 2a, Table 2). The clinopyroxene/melt partition coefficient for a hydrous dacitic melt (CS2014-31) at low oxygen fugacity is within the uncertainty of the extrapolation of the high oxygen fugacity, hydrous partition coefficients (Fig. 2a). Two hydrous experiments at high f_{02} produced melts with andesitic compositions coexisting with clinopyroxene (CS2014-19, DRB2015-2). On the other hand, no anhydrous experiments at high f_{02} were successfully performed for andesitic compositions. Therefore, we cannot make any direct comparison between results from hydrous and anhydrous high f_{02} and esitic experiments (Fig. 2a). Overall, these results lead to a conundrum: the total sulfur partition coefficients for basaltic melts indicate that the difference in oxygen fugacity is responsible for the two trends in Figure 2a; instead the clinopyroxene-melt partition coefficients for the dacitic melt suggest that the presence of water may be responsible for the different trends.

The oxygen fugacity and compositional dependence of the sulfur partition coefficient can be removed if the partition coefficient is calculated by dividing the sulfur concentration of the crystal by the sulfue concentration in the melt, calculated from the shift of the sulfur peak and the bulk sulfur concentration (Table 2). When this is done, all of the S^{2–} partition coefficients show a weak correlation (Fig. 2b). However, the partition coefficients for the melts with silica concentrations below ~51 wt% from Callegaro et al. (2014) all cluster slightly below the line suggesting the existence of a small compositional effect for clinopyroxenes crystallizing from low-silica melts. If the influence of these points is removed, the S^{2–} partition coefficient between clinopyroxene and melts with SiO₂ higher than ca. 51 wt% can be calculated from the mean of the points in Figure 2b: 0.063 ± 0.010 (1 σ St.dev, about the mean).

Partitioning of sulfur between melt and plagioclase

Plagioclase crystallized in 7 experiments (Tables 2 and 3). However, as discussed below, one of the experiments appears to have crystallized plagioclase during quench, hence only six parti-

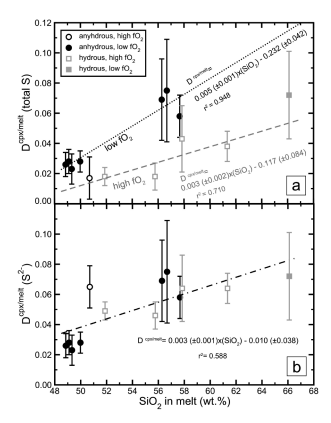


FIGURE 2. Correlation of sulfur clinopyroxene-melt partition coefficient, D, with melt composition at low and high oxygen fugacity as well as with and without added water. (a) Total sulfur partition coefficients for clinopyroxene-melt vs. silica concentration in the melt. Note that, with one exception, experiments at low f_{0_2} display higher partition coefficients (by a factor of approximately 3) than hydrous experiments at high f_{02} , as discussed in the text. The stippled line labeled low f_{02} is fit through the anhydrous experiments performed in graphite capsules and the dashed line labeled high f_{02} is fit to hydrous experiments performed in gold-palladium capsules (see Table 2). (b) S²⁻ partition coefficient between clinopyroxene and melt as a function of the SiO₂ concentration in the melt. Note that the two trends presented in a collapse into a single trend almost independent of the silica concentration in the melt, and that the S2- partition coefficient appears constant for melts above approximately 51 wt% SiO2. The uncertainties in the measured partition coefficients shown in this and subsequent figures are 10 uncertainties calculated from either the standard deviation about the mean (where multiple analyses were performed; see Table 2), or uncertainties calculated from counting statistics (where only single measurements were available). See text for further discussion.

tioning measurements are deemed reliable. The concentrations of sulfur in the crystals varied from 37 to 93 ppm and the calculated total sulfur partition coefficients from 0.036 to 0.393, but most of them are near the lower value (Fig. 3). In general, the partition coefficients for plagioclase were similar to, or slightly higher, than those for clinopyroxene. Most plagioclase crystals were in the compositional range of An_{40-45} , although one An_{61} crystal formed in DRB2012-36. With the exception of a hydrous experiment at high f_{02} (CS2014-20), no influence of the anorthite content of the crystals or their iron concentration on the total sulfur partition coefficients was detected, nor was any influence of oxygen fugacity

observed (Fig. 3). The few plagioclase crystals in the anomalous 1 experiment CS2014-20 with the high partition coefficient were 2 anhedral with morphologies suggestive of rapid growth during 3 quench, therefore we surmise that plagioclase-melt partition coefficients for this experiment are not valid. The lack of observable 5 influence of crystal composition and oxygen fugacity is also true 6 when the S^{2–} partition coefficient is calculated (not shown). 7

Partitioning of sulfur between melt and olivine or orthopyroxene

Olivine and orthopyroxene crystallized in one and two anhy- 11 drous experiments, respectively (Tables 2 and 3). The concen- 12 trations of sulfur in the orthopyroxenes and olivines were near, 13 or at, the detection limits of the SXRF analysis. Consequently, 14 the corresponding total sulfur partition coefficients (which are 15 the same as the S²⁻ partition coefficients for the low oxygen 16 fugacity experiments because all sulfur is found in the S²⁻ state) 17 were significantly lower than those of clinopyroxene (Table 2). 18 The olivine in DRB2012-29 was at the detection limit of our 19 analysis with a S concentration of 1 ± 0.2 ppm, yielding a maxi- 20 mum sulfur partition coefficient of 0.001. The orthopyroxenes 21 that co-existed with a basaltic melt in the low oxygen fugacity 22 experiment DRB2012-29 (at 0.8 GPa, 1240 °C) contained 3 ± 23 2 ppm S, and produced a total sulfur orthopyroxene/melt D of 24 0.003. The orthopyroxene crystallized from a basaltic melt in 25 the high oxygen fugacity experiment (CS2014-14 at 1.0 GPa, 26 1240 °C) contained an average of 2 ± 1 ppm S; the total sulfur 27 partition coefficient is 0.002 and the S2- partition coefficient is 28 0.007. Because the sulfur concentrations in the orthopyroxene 29 and olivine are very close to the detection limit for SXRF 30 (approximately 1 ppm), their partition coefficients should be 31 considered maximum values. 32

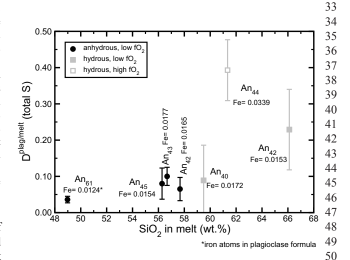


FIGURE 3. Total sulfur partition coefficients for plagioclase-melt 51 vs. silica concentration in the melt, at low and high oxygen fugacities 52 and with and without added water. Both the anorthite content and the 53 amount of iron in the plagioclase formula are displayed next to the 54 data points. The range in anorthite concentrations is too small to obtain 55 any meaningful relation between them and the partition coefficients, 56 and no clear dependence upon the silica concentration in the melt is observed. (Note that as discussed in the text the value for the one hydrous 57 58 experiment at high oxygen fugacity should be considered unreliable.) 59

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Partitioning of sulfur between melt and amphibole

2 Amphibole crystallized in two hydrous experiments, CS2014-3 20 (at 0.8 GPa, 1000 °C and high oxygen fugacity) and CS2014-30 4 (at 0.8 GPa, 1000 °C and low oxygen fugacity). In CS2014-30 the 5 average S concentration in amphibole was 87 ± 58 ppm (1 σ St.dev.) 6 and in CS2014-20 it was 29 ± 14 ppm, which yielded total sulfur 7 partition coefficients of 0.127 and 0.123, and S2- partition coef-8 ficients of 0.127 and 0.208, respectively. These values are larger 9 than measured in clinopyroxenes. However, the crystallization of 10 amphibole in CS2014-30 is surprising due to the measured water 11 concentration of only 1.1 wt% in the quenched glass, although 12 the presence of halogens in the melt of this experiment may be 13 responsible for the presence of amphibole (see review in Webster et 14 al. 2018). Amphibole crystallizing at such a low water concentra-15 tion might suggest the possibility of disequilibrium in CS2014-30; 16 we cannot discount this possibility, but we decided to include this 17 value because it is the only amphibole/melt partition coefficient 18 measured at high oxygen fugacity. 19

DISCUSSION

Influence of silicate mineral structure on *D*

Although clear trends in the clinopyroxene/melt partition coef-23 ficients as a function of melt composition can be seen in Figure 24 2, it is well known that the partition coefficients of trace elements 25 are controlled more by crystal chemistry and structure than by 26 melt composition (e.g., Blundy and Wood 1991). The total sulfur 27 partition coefficients and the S2- partition coefficients in Table 2 28 correlate with the average bond distance for the mean M(2)-O 29 distances in olivine, orthopyroxene, and clinopyroxene, with the 30 mean Ca-O distance in plagioclase, and with the mean M(4)-O 31 distance for a pargasitic hornblende (Fig. 4). 32

We propose that the dominant dissolution mechanism of sulfur 33 is the replacement of some oxygen by S2-. This hypothesis is based 34 upon the similar sizes and same charge of S2-, 170 pm in sixfold 35 36 coordination and O2-, 121 pm in twofold coordination (Shannon 37 1976). This replacement also is suggested by the observation that 38 the total partition coefficient between clinopyroxene and melt is 39 significantly lower in experiments at high f_{02} , where most sulfur in the melt is present as an S⁶⁺ species, as opposed to the low f_{0_2} 40 41 experiments, where the sulfur in the melt is in an S²⁻ species (e.g., Fincham and Richardson 1954; Wilke et al. 2011). Furthermore, 42 the near-constant value of the clinopyroxene-melt S2- partition 43 coefficient is seen in Figure 2b, despite variations in melt com-44 position, water concentration, and oxygen fugacity, is more easily 45 explained if S²⁻ is exchanging between the crystals and the melts. 46 The correlation between the bond lengths and the sulfur crystal/ 47 melt partition coefficient (Fig. 4) is interpreted to indicate that 48 crystallographic sites with average cation-oxygen bond lengths 49 greater than 220 pm are necessary to accommodate substantial 50 51 amounts of sulfur, greater than a few parts per million, and that 52 sulfur replaces some of the oxygen coordinating the M(2) sites in olivine, orthopyroxene, and clinopyroxene, some oxygen co-53 ordinating the alkalis and alkali earths in plagioclase, and oxygen 54 coordinating the M(4) site in amphibole. 55 The sulfur partitioning (Fig. 4) also positively correlates with

The sulfur partitioning (Fig. 4) also positively correlates with
an increasing fraction of bridging O atoms in the crystal structure.
However, the replacement of bridging O atoms by sulfur seems

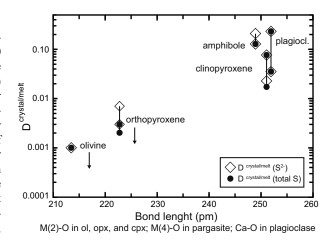


FIGURE 4. Correlation between the range of total sulfur and S^{2–} partition coefficients presented in Table 2 and the size of the M(2)-oxygen bond-length in olivine (Brown 1980{{auth: not listed in Ref list please add or fix}}), orthopyroxene (Cameron and Papike 1980{{auth: not listed in Ref list please add or fix}}), and clinopyroxene (Cameron and Papike 1980), the M(4)-oxygen bond-length in pargasite (Robinson et al. 1973), and the Ca-O bond-length in anorthitic plagioclase (Wainwright and Starkey 1971{{auth: not listed in Ref list please add or fix}}). The vertical lines connect the minimum and maximum partition coefficients measured for each mineral in this study. The arrows associated with olivine and orthopyroxene indicate that these are to be considered maximum values of the partition coefficients. See text for further discussion.

improbable because in this case the similar T-O bond lengths of the minerals would suggest similar partition coefficients, which is not seen in Figure 4.

Influence of clinopyroxene composition on D

When the total sulfur partition coefficients are plotted as a function of the Mg#, molecular Mg/(Mg+Fetotal) of clinopyroxenes crystallized in these experiments, two trends can be observed—one for low and one for high oxygen fugacity (Fig. 5a). However, when the S²⁻ partition coefficients are plotted against the clinopyroxene Mg#, only a weak correlation is visible (Fig. 5b). Most of that dependency is due to the results of Callegaro et al. (2014) at high Mg# values, where it appears that most clinopyroxene/melt S2- partition coefficients are within the uncertainty of each other (Fig. 2b), as was previously observed for the effect of melt composition. However, a small negative dependency of the sulfur partition coefficients (total and S²⁻) on the Mg# might be expected because as iron substitutes for magnesium in the structures of ferromagnesian minerals, the cation-oxygen bond distances get slightly longer (Cameron and Papike 1980). In contrast to the possible small effect of Mg# on partitioning, our results provide no evidence that ^{IV}Al plays a role on sulfur partitioning (Fig. 5c). Such an effect has been hypothesized and investigated for halogen partitioning between clinopyroxene and melt (O'Leary et al. 2010; Rosenthal et al. 2015; Urann et al. 2017; Bénard et al. (2017). Our observations are similar to those of Rosenthal et al. (2015) who found no significant effect of ^{IV}A1 in their partitioning measurements of halogens between clinopyroxene and melt.

Comparison between S partitioning and H, C, F, and Cl partitioning between nominally volatile-free crystals and melts

The partition coefficients of total sulfur and S^{2–} between nominally volatile-free minerals (NVFMs) and silicate melts are typically lower than those of fluorine, higher than those of carbon, and similar to those of chlorine and hydrogen (Fig. 6). Hydrogen, fluorine, and chlorine display similar trends in the value of the partition coefficient as seen for sulfur in ferromagnesian crystals (Figs. 6). Indeed, the crystal/melt partition coefficients for each of these elements increase in the order: olivine < orthopyroxene < clinopyroxene.

The plagioclase/melt fluorine partition coefficients are similar to those of clinopyroxene and amphibole, whereas H and Cl plagioclase/melt partition coefficients are more than one order of magnitude lower (Fig. 6). The hydrous mineral amphibole displays significantly higher partition coefficients of Cl, H, and F than the other ferromagnesian crystals and plagioclase because of the structural role of these volatiles in the amphibole crystal lattice (Fig. 6).

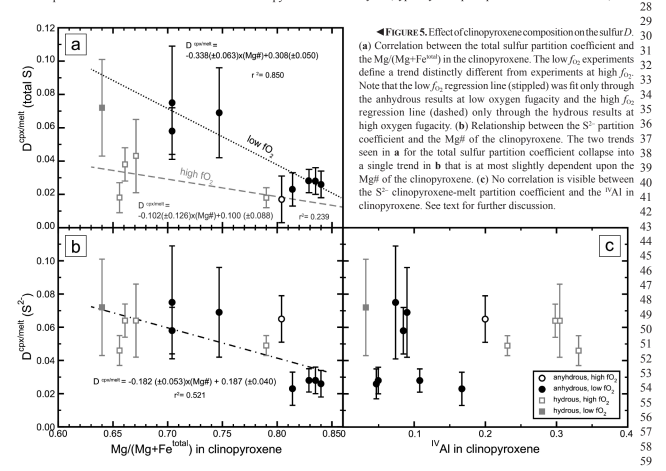
The similar crystal/melt partitioning of H, F, S, and Cl supports the hypothesis that the dissolution mechanism of these elements into silicate minerals is similar and occurs probably either as a replacement of an oxygen atom (F, Cl, and S) or by association with an oxygen atom (H).

In contrast to these volatiles, the partition coefficient of carbon between melt and olivine, orthopyroxene, or clinopyroxene appears to be approximately constant, although orthopyroxene may have a lower partition coefficient than either olivine or clinopyroxene (Fig. 6). However, this behavior needs further investigation because 1 of the relatively large uncertainties in the partition coefficients 2 derived from the very low S concentrations in these crystals. The 3 unique behavior of carbon in comparison to the other elements in 4 Figure 6 suggests a different dissolution mechanism, but discussion 5 of this mechanism is far beyond the scope of this contribution. 6 7

IMPLICATIONS: SULFUR PARTITIONING APPLIED TO NATURAL MAGMATIC SYSTEMS

The proposed crystal/melt partition coefficients for sulfur combined with previously published ones for hydrogen, carbon, fluorine, and chlorine can be used to provide insights into the concentrations of the igneous quintet of major volatiles in magmatic systems.

Estimating the pristine volatile budget of a magma that has 14 15 already solidified is challenging, and this is particularly true for sulfur, carbon, and water, which are degassed earlier than fluorine 16 17 or chlorine in the eruptive history of magmas (e.g., Spilliaert et al. 18 2006). Therefore, quantitative estimates of gas budgets in melts from 19 past eruptions are still scarce because of the rarity of melt inclusions, which are the primary means of determining pre-eruptive volatile 20 concentrations in magmatic melts (e.g., Devine et al. 1984; Johnson ²¹ 22 et al. 1994; Cannatelli et al. 2016). In the absence of melt inclusions, the magma volatile budget may be determined by a combination of 2324 the measurement of volatile concentrations in natural minerals with 25 experimentally determined partition coefficients (mineral/melt D). 26 The challenge in this case is set by the low concentration of volatiles 27 in the crystals, typically at the parts-per-million level in NVFMs, such



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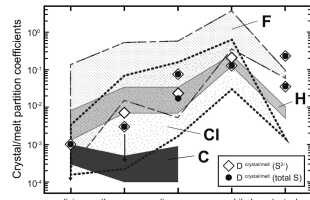
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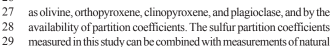
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olivine orthopyroxene clinopyroxene amphibole plagioclase

FIGURE 6. Partition coefficients (Table 2) for total sulfur (black circles) 16 and S²⁻ (open diamonds) measured in this study and in Callegaro et al. (2014). 17 compared to those obtained for carbon and hydrogen as water (Hauri et al. 18 2006; Hamada et al. 2013; Rosenthal et al. 2015; Lloyd et al. 2016) and for 19 fluorine and chlorine (Hauri et al. 2006; O'Leary et al. 2010; Dalou et al. 2012, 20 2014; Guggino 2012; Beyer et al. 2012{{auth: not listed in Ref list please 21 add or fix}}, 2016; Van den Bleeken and Koga 2015; Bénard et al. 2017; 22 Lloyd et al. 2016). Minimum and maximum sulfur partition coefficients are 23 plotted for clinopyroxene, amphibole, and plagioclase. The arrows below the 24 sulfur partition coefficients for olivine and orthopyroxene indicate that the plotted values are considered maxima. 25

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- crystals to determine the concentration of sulfur in coexisting melts 31
- from which the minerals crystallized. Given the measurement of 32
- sulfur in a crystal and knowledge of the oxygen fugacity of crystal-33
- lization, the S2- partition coefficients determined in this study can be used to calculate the S2- concentrations in the melt. This value can 34
- be combined with the S^{6+}/S^{2-} ratio (Wilke et al. 2011) to calculate 35
- 36 total sulfur in the melt. This technique allows determination of sulfur
- 37 in coexisting melts ranging in oxygen fugacity from the FMQ buf-
- 38 fer to more reduced conditions, such as the dominant magmas on
- 39 Earth, MORBs, and those associated with Large Igneous Provinces.
- 40 However, our newly established partition coefficients can be applied
- 41 also to magmas with oxidation states near FMQ+2, typically found
- 42 at convergent margins (Carmichael 1991).
- 43 On the other hand, if the oxidation state is not known, the total 44 sulfur partition coefficients for reduced or oxidized conditions can 45 be used.
- 46 These partition coefficients will allow calculation of the sulfur 47 budget of ancient natural basalts, particularly those constituting 48 Large Igneous Provinces (LIPs), whose timing often coincides with 49 mass extinction events (Wignall 2001; Bond and Wignall 2014). 50 Quantifying gas loads and rates of degassing for LIP magmas is 51 fundamental to understand this causal relationship, and particular 52 attention, through analyses or models, has been recently directed to 53 sulfur (e.g., Self et al. 2008, 2014; Callegaro et al. 2014; Jones et al. 54 2016; Schmidt et al. 2016).
- 55 We stress however that we are working with S concentrations 56 very close to the detection limits of the SXRF technique; therefore 57 the uncertainties involved are very large. A further characterization 58 of standards will help reduce the uncertainties in the future, but 59

at present we advise that the here proposed partition coefficients should be applied only to analyses carried out by the same analytical technique (SXRF), as well as the same data reduction routine (PYMCA). Thanks to fast acquisition times and small spot size (few square micrometers), SXRF provides the opportunity to map sulfur in crystals and to potentially discover evidence of degassing episodes or magma mixing events (i.e., S loss or S uptake by the system) during crystal growth, an application of great interest in the study of active volcanic systems.

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- 53 MANUSCRIPT HANDLED BY BRUCE WATSON
- 54
- **Endnote:**

¹Deposit item AM-20-57188, Supplemental Material. Deposit items are free to 56 all readers and found on the MSA website, via the specific issue's Table of Con-57 tents (go to http://www.minsocam.org/MSA/AmMin/TOC/2020/May2020_data/ 58 May2020_data.html). 59