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# Fractional degassing of S, Cl and F from basalt magma in the Bárðarbunga rift zone, Iceland

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## Abstract

The composition of gas emitted from a volcano producing basalt magma can vary during an eruption and according to the volcano-tectonic setting of the degassing vents. Post-eruptive filter-pack gas samples from the 2014-2015 Holuhraun crater in the Bárðarbunga rift zone have lower ratios of S over halogens (Cl and F) and elevated F/Cl (~50 times lower S/Cl and ~5 times higher F/Cl; mass ratios) compared to samples of the syn-eruptive gas plume. The compositional changes are readily explained by Rayleigh distillation with decreasing sulphur concentrations and increasing concentrations of halogens and F relative to Cl in the final gas phase. For Cl the vapour-melt partition coefficient ( $\mathcal{D}^{V/M}$ ) decreased from 13-85 to 2.2 during residual degassing, whereas that of F remained uniform at approximately 1.8. Distinctly different degassing behaviour is observed for Cl and F. High  $\mathcal{D}$  for Cl may indicate an important influence of sulphur and water on Cl volatility in basaltic melt, whereas that of F remains unaffected. The primary gas of the Holuhraun magma had similar ratios of S over Cl and F as observed at the Kilauea rift zone which, together with lower S/halogens in the residual gas in both cases, suggests similar degassing mechanism. By inference, initial CO<sub>2</sub> degassing is likely to have occurred subglacially close to the Bárðarbunga central volcano before and during the 2014-2015 eruption on the rift-related fissure swarm.

## Introduction

Degassing of basaltic magma is known to impact the environment by emission of several different gaseous species but the physics of the degassing process is less well understood. At depth, where CO<sub>2</sub> is the dominant volatile species, gas segregation from basaltic or low-viscosity magma is thought to follow a cyclic behaviour, which may lead to alternating explosive and effusive activity (Sparks, 2003). The gas phase and the rising melt can evolve from a state of thermodynamic equilibrium to that of a progressive open-system fractional distillation, especially during the waning stage of an eruption (Aiuppa, 2009). Fractional degassing at shallow depth has been inferred from changing gas composition during single eruptions from craters at different altitude and during different types of volcanic activity, as in the case of the persistently degassing Mount Etna, Sicily (e.g. Aiuppa et al., 2002; Burton et

51 al., 2003; Allard et al., 2005). In addition, the syn-eruptive gas phase of a basaltic fissure  
52 eruption is expected to be different from the residual gas phase of a cooling lava during  
53 emplacement and crystallization (Swanson & Fabbi, 1973). A well-known example of a  
54 basaltic fissure eruption is that of Laki in Iceland which may have released up to 20% of the  
55 total emitted SO<sub>2</sub> and significant proportion of the halogens during lava field emplacement  
56 and crystallisation (Thordarson et al., 1996). The final degassing stage may thus lead to local  
57 hazes with different gas composition than the principal gas phase. Fluorine, for instance, with  
58 its relatively high solubility in basaltic melt compared to Cl and S (Carroll & Webster, 1994),  
59 will principally enter the final gas phase. Such degassing of F can lead to fluorosis in grazing  
60 animals, a well-known effect that caused the death of approximately two thirds of the  
61 Icelandic livestock during and soon after the Laki eruption (Thorarinsson, 1969; Pétursson et  
62 al., 1984; Thordarson and Self, 2003).

63  
64 Variations in the gas composition at basaltic volcanoes are also related to the global tectonic  
65 context, with hotspot volcanoes generally emitting gas of higher S/halogens than those in  
66 subduction-zone settings, despite several common factors as reviewed by Aiuppa (2009). At  
67 Kilauea, Hawaii, gas of different composition is released at summit craters than is emitted in  
68 flank eruptions in rift zones (e.g. gas type I and II of Gerlach & Graeber, 1985; Greenland et  
69 al., 1988; Symonds et al., 1994; Edmonds et al., 2009). Likewise, during the 2001 eruption of  
70 Etna, Burton et al. (2003) observed remarkably similar S/halogens in gas from the Northeast  
71 crater (NEC) and the Central craters as well as in the bulk gas plume, whereas gas from the  
72 Southeast crater (SEC) had considerably lower S/halogens and that of the flank crater, which  
73 fed the 2001 summer lava flow, had even lower ratios. Their results confirmed observations  
74 by Aiuppa et al. (2002), which in addition measured decreasing Cl/F in gas from the flank  
75 crater. These studies suggest a primary gas phase from single magma body degassing beneath  
76 NEC and Central craters, whereas SEC and the southern flank crater emitted a secondary gas  
77 phase. Aiuppa (2009) extended the open system Rayleigh-type degassing model for Etna to  
78 other basaltic volcanoes, a model that assumes uniform vapour-melt partition coefficients for  
79 S, Cl and F. A similar method was applied by Mather et al. (2012) when discussing degassing  
80 mechanism at Kilauea. However, neither Cl nor F volatility and their partition coefficients  
81 between vapour and melt phases are well understood. A comparison of the primary gas phase  
82 released during basaltic fissure eruption to that produced during the final degassing stages at  
83 the end of an eruption should provide better constraints on the mechanism, volatile solubility,  
84 and therefore the volatile partition coefficients between the silicate melt and the gas phase,  
85 needed for better understanding of the degassing processes.

86  
87 In this article syn- and post-eruptive gas phases of the Holuhraun eruption (29 August 2014 to  
88 27 February 2015) within the rift zone of the Bárðarbunga volcanic system in Iceland are  
89 compared. The eruption site was located approximately 40 km N of the Bárðarbunga central  
90 caldera, which is positioned above the presumed centre of the Iceland mantle plume. The  
91 eruptive fissure produced approximately 1.5 km<sup>3</sup> of basalt of uniform composition (ol-tholeiite)  
92 and liberated close to 10 Tg of SO<sub>2</sub> into the atmosphere (Sigmundsson et al., 2014; Gíslason et  
93 al., 2015; Gauthier et al., 2016; Bali et al., 2018; Halldórsson et al., 2018; Pfeffer et al., 2018).  
94 The large amount of gas emission from Holuhraun caused repeated air pollution events  
95 characterized by atmospheric SO<sub>2</sub> concentrations exceeding air quality standards, as well as  
96 high concentrations of fine-grained volcanogenic aerosols (Ilyinskaya et al., 2017; Stefánsson  
97 et al., 2017). Snow precipitated around the lava field, rain and the surrounding rivers were  
98 monitored during the eruption and had elevated F concentrations (Gíslason et al., 2015;  
99 Galeczka et al., 2017; Stefánsson et al., 2017) despite its very low abundance in the main gas  
100 plume (Gauthier et al., 2016). Understanding the origin and fate of fluorine during degassing

101 processes is thus of importance in order to assess its release to the atmosphere and subsequent  
102 environmental impact. Here, results are presented on S and halogen (Cl and F) concentrations  
103 in the post-eruptive gas phase released at Holuhraun in order to verify whether or not fluorine  
104 is ultimately degassed and if a simple Rayleigh distillation process occurring during the  
105 progressive basalt degassing explains the observed gas compositions. Halogen partition  
106 coefficients  $D^{\text{vapour/melt}}$  are estimated for Cl and F from S/Cl and S/F ratios and their variability  
107 towards the end of the eruption is assessed. The results are finally compared with those from  
108 Etna and Kilauea volcano (e.g., Gerlach & Graeber, 1985; Aiuppa et al., 2002; Burton et al.,  
109 2003; Allard et al., 2005; Edmonds et al., 2009; Mather et al., 2012) and show that the gas phase  
110 composition reflects the intrinsic parameters of the fractional degassing mechanism.

### 111 112 113 **Sample collection and analysis**

114 High-quality syn-eruptive gas samples are not easily collected during intense effusive activity  
115 such as the Holuhraun eruption. Two syn-eruptive gas plume samples could be collected in still  
116 weather on 2 October 2014, when the gas plume was grounded for a few hours because of a  
117 short-lived temperature inversion of the atmosphere (Gauthier et al., 2016; Fig. 1). A few  
118 months later, 21 January 2015, a third sample of the gas plume was collected from a helicopter  
119 (Ilyinskaya et al., 2017). This latter sample has an order of magnitude lower SO<sub>2</sub> concentration  
120 compared to the earlier samples, possibly reflecting greater air dilution on the plume's edge, or  
121 increasing distance from the vent.

122  
123 Recently published results (Stefánsson et al., 2017; Ilyinskaya et al., 2017) on ground-collected  
124 syn-eruptive gas composition can be compared with those given by Gauthier et al. (2016). Out  
125 of four filter-pack gas samples collected during the eruption by Stefánsson et al. (2017), three  
126 have very similar S/Cl mass ratios with an averaged value of  $41 \pm 7$ , characteristic of the syn-  
127 eruptive gas plume (Gauthier et al., 2016). In marked contrast, their samples exhibit much  
128 higher fluorine concentrations than reported by Gauthier et al. (2016), leading to lower Cl/F  
129 ratios, somewhat intermediate between primary vapour compositions (Gauthier et al., 2016)  
130 and residual vapour compositions (this work, Table 1). Marked enrichments in fluorine may  
131 arise though from other sources than the syn-eruptive gas phase, such as secondary vapour from  
132 the cooling lava field. The same holds true for ground-collected gas samples taken by Ilyinskaya  
133 et al. (2017), who also suggested that extreme variability in their own samples may be explained  
134 by a gas phase “partially sourced from the degassing lava”. In addition to the fortunate two  
135 samples from Gauthier et al. (2016), we therefore consider only their helicopter-aided airborne  
136 plume sample as representative of the principal syn-eruptive gas phase (Ilyinskaya et al., 2017).  
137 Despite being strongly diluted compared to the gas plume samples of Gauthier et al. (2016), the  
138 helicopter-aided sample from Ilyinskaya et al. (2017) has identical syn-eruptive S/Cl but  
139 significantly lower S/F and Cl/F.

140  
141 Ten days after the lava extrusion ended, the residual gas phase released from the cooling magma  
142 was collected. Three post-eruptive filter-pack samples were obtained in calm weather on 9  
143 March 2015 inside the main crater, approximately 50 m above the cooling lava lake (Fig. 1).  
144 Intensive degassing was observed from the lava-filled crater with visually estimated maximum  
145 gas emission from cracks inside the crater walls, one of which was selected for gas sampling.

146  
147 The same filter-pack technique as presented in Gauthier et al. (2016) was employed. Each filter  
148 pack was connected to a pump operating at a flow rate of 11 l/min for 23-72 min, collecting  
149 three gas and aerosol samples (HH1, HH2 and HH3; Table 1) from a total air volume of 0.25-  
150 0.79 m<sup>3</sup>. For comparison, the atmospheric composition free of volcanic gas (blank) was

151 collected approximately 500 m west of the lava field over one hour. Filters were subsequently  
152 analysed for S, Cl and F concentrations using ion chromatography at ICCF-Sigma, Université  
153 Clermont Auvergne, with analytical uncertainty of approximately 5%. For the consolidated and  
154 thus outgassed lava, S concentration was measured with a CHNS/O analyser at LMV-  
155 Université Clermont Auvergne, but those of Cl and F using absorption spectrophotometry and  
156 selective electrodes, respectively, at CRPG-Nancy, France. The  $2\sigma$  errors for S, Cl and F  
157 concentrations in the lava are less than 10%.

## 158 159 **Results**

160 The post-eruptive gas phase has S, Cl and F concentrations two to three orders of magnitude  
161 higher than the atmospheric background, which is therefore neglected. The gas samples are  
162 characterised by S, Cl and F concentrations ranging from 49-164, 53-104 and 27-46 (in  $\text{mg/m}^3$ ),  
163 respectively, and their composition significantly differs from the syn-eruptive gas plume. The  
164 S concentrations in the post-eruptive gas phase are lower than those emitted during the primary  
165 eruptive phase by a factor of 3 on average. In marked contrast, both Cl and F are in much higher  
166 concentrations in the post-eruptive gas: 11 times more Cl and 62 times more F. The S/Cl mass  
167 ratio is close to unity (0.92 - 1.58), substantially lower than the 40-52 of the primary gas phase  
168 (Gauthier et al., 2016). So is the Cl/F mass ratio, which is on average 2 in the post-eruptive gas  
169 phase against  $11 \pm 4$  in the syn-eruptive plume. Consequently, the S/F mass ratio (1.85 - 3.53)  
170 is two orders of magnitude lower in the final gas phase and its decline is already observed in  
171 the late January 2015 gas (Table 1). Although S remains the principal minor gas component,  
172 these observations suggest a much stronger halogen contribution to the vapour phase at the end  
173 of the eruption.

## 174 175 176 **Discussion**

### 177 *Arguments for almost complete S degassing*

178 The sulphur-rich gas plume from the Holuhraun eruption affected people over most part of  
179 Iceland because of the high volatility of S from the olivine tholeiitic basalt. Sulphur is known  
180 to degas from basaltic melt until melt quenching or solidification, which for example explains  
181 higher concentrations in water- vs air-quenched tephra, and tephra vs crystallized lava  
182 (Thordarson et al., 1996; Óladóttir et al., 2007; Haddadi et al., 2017). The very low S  
183 concentration in the degassed lava (97 ppm; Gauthier et al., 2016) compared to the initial  
184 amount preserved in melt inclusions (up to 1644 ppm S; Bali et al., 2018) suggests that 94% of  
185 sulphur was lost during magma ascent and evolution. Sulphide minerals could play a role in the  
186 sulphur budget of the Holuhraun products since a few globules were observed in the early lava  
187 and tephra in September 2014. However, most of the sulphides are present as irregular, partially  
188 dissolved particles and exhibit variable S isotope composition, whereas other isotope ratios  
189 remain mostly constant (Halldórsson et al., 2018). These sulphides are partially xenocrystic and  
190 most likely picked up during magma ascent. Smaller euhedral sulphide spheres are crystallized  
191 immiscible melt in the Holuhraun magma itself. However, they are scarce enough to ensure that  
192 their crystallization (acting as a S sink) and their destabilization (acting as a S source) does not  
193 drastically change the S budget at Holuhraun. It thus appears that most of the sulphur escaped  
194 the magma as a volatile species before the crystallisation of the lava, mostly as gaseous  $\text{SO}_2$   
195 since  $\text{H}_2\text{S}$  does not exceed a few percent in the gas phase (Gíslason et al., 2015). With  $\text{SO}_2$   
196 averaging *ca* 30% in the gas phase (e.g. Bali et al., 2018), the S liquid-gas partition coefficient  
197 ( $D^{l/g}(\text{S})$ ) can be estimated at  $97 \cdot 10^{-6} / 0.15 = 6 \cdot 10^{-4}$ . The figure must be considered as a maximum  
198 value for  $D^{l/g}(\text{S})$  (note that  $D^{l/g}$  is simply the reciprocal of  $\mathcal{D}^{V/M}$ ) because the final S  
199 concentration in the most degassed eruptive products could be lower than 97 ppm. Whichever  
200 the exact value, it appears reasonable to assume  $D^{l/g}(\text{S})$  close to nil. Finally, even if some S

201 remains in quenched groundmass glass or in the rare sulphides, the bearing on its partition  
202 coefficient between vapour and melt ( $\mathcal{D}^{V/M}$ ) is none, since that coefficient only measures the  
203 vapour-melt partition but neither vapour-glass nor melt-solid fractionation.

204

#### 205 *Estimation of vapour/melt partition coefficients of Cl and F*

206 In contrast to S, which is severely depleted in the final solid product, Cl and F concentrations  
207 in the crystallized lava reach 110 and 190 ppm, respectively, which is within the range observed  
208 in melt inclusions and the lava groundmass (Gauthier et al., 2016; Bali et al., 2018). Moreover,  
209 given their higher solubility in basaltic melt (Carroll & Webster, 1994), both Cl and F are  
210 expected to degas principally at or very close to the surface. The important decreases in S/Cl,  
211 S/F and Cl/F from the syn-eruptive gas plume to the post-eruptive gas phase clearly indicate  
212 decreasing availability of S for the gas phase in the crystallizing basalt magma and increasing  
213 relative concentrations of Cl and especially F in the final gas phase. This concentration pattern  
214 strongly suggests a Rayleigh distillation process, with most of the S extracted from the melt.  
215 The Rayleigh law,  $C = C_0 \times f^{D-1}$ , where C denotes the measured concentration of a given gas  
216 species,  $C_0$  its initial gas concentration, f the fraction of gas remaining in the magma (vapour  
217 plus melt) and D the concentration in liquid divided by that of the gas (i.e. the partition  
218 coefficient,  $D^{l/g}$ ) can be linearized as:

219

$$220 \log Cl = \log Cl_0 + (D_{Cl}^{l/g} - 1) \times \log f$$

$$221 \log S = \log S_0 + (D_S^{l/g} - 1) \times \log f$$

222

223 When  $D^{l/g}(S)$  is close to zero and can be neglected, the equations can be combined to eliminate  
224 f, which leads to:

225

$$226 \log Cl = (1 - D_{Cl}^{l/g}) \times \log S - [(1 - D_{Cl}^{l/g}) \times \log S_0] + \log Cl_0.$$

227

228 The  $D^{l/g}(Cl)$  can thus be determined from the slope of the linear relationship between log Cl  
229 and log S. The vapour/melt partition coefficient ( $\mathcal{D}^{V/M}$ ) for Cl, being simply the reciprocal of  
230  $D^{l/g}(Cl)$ , thus can readily be calculated. The  $\mathcal{D}^{V/M}$  for F is obtained in the same way. Note that  
231 if  $D^{l/g}(S)$  were slightly higher than 0, these calculated  $\mathcal{D}^{V/M}$  for halogens would be the maximum  
232 possible values.

233

234 The results are plotted in Figure 2 where Cl and S concentrations in gas samples collected  
235 during the eruption form a strong linear correlation ( $R^2 = 0.990$ ). From the slope a surprisingly  
236 low  $D^{l/g}(Cl)$  is calculated that leads to unusually high value for  $\mathcal{D}^{V/M}$  ( $= 1/D^{l/g} = 85$ ), meaning  
237 that 85 times more mass of Cl went into the gas phase compared to what remained in the melt  
238 phase. The results from Stefánsson et al. (2017) fall on the lower concentration end of the  
239 correlation. Taken together with the present results, a somewhat lower regression coefficient  
240 ( $R^2 = 0.987$ ) is obtained and a slope that yields less extreme  $\mathcal{D}^{V/M}$  of 13. At first glance, our  
241 values might seem extreme, but in fact, they are on the same order of magnitude as the highest  
242 experimentally determined  $\mathcal{D}^{V/M}$  by Stelling et al. (2008) and Alletti et al. (2009;  $\mathcal{D}^{V/M}$  for Cl  
243 as high as 24). Given the difficulties in determining  $\mathcal{D}^{V/M}$  for Cl, such as the presence or absence  
244 of a third phase (i.e. brine) in the experimental products (Webster et al., 1999), we consider the  
245 value of 13 to 85 realistic, albeit possibly a maximum value. Chlorine most likely formed gas  
246 species and outgassed at very low pressure, close to or at the surface, aided by extensive water  
247 and sulphur degassing. In fact, evidence for sulphur-aided degassing of Cl does exist. Sulphur  
248 chloride species such as  $S_2Cl$  can form in a gas phase, depending on the H/C and oxygen

249 fugacity (Zolotov & Matsui, 2002), and complex ligands such as  $S_2Cl_2^{4-}$ ,  $S_2Cl^{3-}$ ,  $SCl_2^{3-}$ , and  
250  $S\cdot Cl_2^{3-}$  have been proposed for volatile transport of platinum-group elements (Fleet & Wu,  
251 1993). Clearly experimental results are needed to improve understanding of sulphur degassing  
252 effect on the volatility of Cl in mafic magma. The slope of the trend line for the three post-  
253 eruptive gas samples ( $R^2 = 0.84$ ) is significantly lower leading to an order of magnitude smaller  
254  $\mathcal{D}^{V/M}$  for Cl, or only 2.2. The low  $\mathcal{D}^{V/M}$  in the post-eruptive gas phase could reflect the decreased  
255 role of a carrier gas for Cl such as  $H_2O$  or  $SO_2$ , which concentrations can be assumed as  
256 relatively low in the post-eruptive gas phase. Interestingly, similar  $\mathcal{D}^{V/M}$  has been determined  
257 in several experimental studies on Cl solubility in basalt without much sulphur present (Webster  
258 et al., 1999; Stelling et al., 2008; Alletti et al., 2009). Such a low  $\mathcal{D}^{V/M}$  for Cl may thus be  
259 characteristic for residual degassing of basalts with a gas phase with relatively low S/Cl as  
260 observed at Holuhraun.

261  
262 Fluorine concentrations in all syn-eruptive samples show more scatter, possibly due to  
263 contributions from different sources as suggested by Ilyinskaya et al. (2017). Therefore, we  
264 again calculate best fit only for the two plume samples of Gauthier et al. (2016) and the  
265 helicopter-aided sample of Ilyinskaya et al. (2017). The resulting  $\mathcal{D}^{V/M}$  for F in the syn-eruptive  
266 gas phase is close to that obtained for the three post-eruptive gas samples: 1.9 vs 1.7,  
267 respectively. The much lower  $\mathcal{D}^{V/M}$  for F compared to Cl during the eruption illustrates its  
268 higher solubility in the basaltic melt, whereas in the residual gas phase similar volatility is  
269 obtained for both halogens. Knowledge of  $\mathcal{D}^{V/M}$  for F is limited and experimental results are  
270 needed to test the accuracy of our approach. The halogen concentration in the residual gas phase  
271 is thus approximately twice that of the final crystallizing melt.

#### 272 273 *Comparison with Etnean gas*

274 Halogen degassing has been extensively studied at Mount Etna, where persistent open-conduit  
275 degassing presents ample opportunities for gas sampling. Fractional degassing was proposed  
276 for the 2000 eruption at the SE crater where variations of S/Cl correlated with eruptive  
277 behaviour (Allard et al., 2005). Moreover, temporal decline of S/Cl and Cl/F was observed from  
278 the central craters to the flank lava-forming eruption in 2001 (Aiuppa et al., 2002; Burton et al.,  
279 2003). Despite the obvious difference between the subduction zone-related Etna stratovolcano,  
280 where gas was sampled from several craters at different altitude, and Holuhraun in the  
281 Bárðarbunga rift zone, where both syn- and post-eruptive gas was sampled from the same  
282 fissure crater, an interestingly similar degassing patterns can be observed. In Figure 2, the S, Cl  
283 and F gas concentrations from Aiuppa et al. (2002) are plotted and the slope of the best-fit lines  
284 used to infer the vapour-melt partition coefficients for Cl and F during the 2001 flank eruption.  
285 Severe S depletion relative to the halogens was observed and the calculated  $\mathcal{D}^{V/M}$  for Cl is 2.3,  
286 and 1.8 for F, indistinguishable from that of the Holuhraun post-eruption gas phase. Flank  
287 eruptions at Etna thus bear similarities with fractional degassing at Holuhraun and, as suggested  
288 by Aiuppa et al. (2002, 2009), with that of Kilauea volcano, Hawaii. However, Etna produces  
289 gas especially enriched in Cl, as expected from its subduction-zone settings. The much higher  
290 gaseous S concentrations and S/Cl at the hotspot-related Bárðarbunga and Kilauea is worth  
291 further discussion.

#### 292 293 *Comparison with Kilauea*

294 In Figure 3, we compare the S/Cl and S/F mass ratios measured in syn- and post-eruptive gas  
295 phases of Holuhraun, as well as in the degassed lava. The decreasing trend observed from the  
296 syn-eruptive gas plume through the residual gas phase to the degassed lava is consistent with a  
297 simple Rayleigh fractionation. The simplest explanation for the observed behaviour is

298 progressive distillation of S from the rift-zone magma. When the eruption ceases, lava cools  
299 down and residual volatiles form a gas phase causing the melt to crystallise. The flux of carrier  
300 gas (mostly H<sub>2</sub>O and SO<sub>2</sub>) is still important enough to flush halogens out of the cooling lava  
301 but the low S concentration in the residual melt leads to a drastic decrease in Cl volatility, and/or  
302 increase in Cl solubility. Because fraction f of volatiles remaining in the melt decreases as the  
303 outgassing progresses, the Rayleigh equation implies increasing halogens, and notably fluorine,  
304 concentrations, not only in gas collected close to the degassing lava but also in the near-  
305 environment. The volcanic haze of basaltic fissure eruptions consequently comprises, in  
306 addition to water and sulphur dioxide, chlorine and fluorine gas species that may carry many  
307 metal and metalloid species out of the final melt of crystallizing basaltic lava. Accordingly,  
308 plume samples collected close to advancing lava field, like those sampled by Stefánsson et al.  
309 (2017) and Ilyinskaya et al. (2017) are prone to be mixed with F-enriched vapour, as  
310 exemplified in Figure 3 where their samples lie in between the two end members.

311  
312 Gauthier et al. (2016), and more recently Edmonds et al. (2018), showed that hotspot-related  
313 volcanoes exhibit common trace-element degassing patterns. One of the best-studied hotspot  
314 basaltic volcanoes is Kilauea, Hawaii. Although the geodynamics of Bárðarbunga volcano is  
315 under the influence of both the Iceland mantle plume and extensional tectonics, while Kilauea  
316 is merely related to the Hawaii hotspot, their volcano-tectonic context is surprisingly  
317 comparable. Type II SO<sub>2</sub>-rich gas is characteristic of the Kilauea east rift zone (Gerlach and  
318 Graeber, 1985), and also of the gas plume released from the Holuhraun craters in the  
319 Bárðarbunga fissure swarm (= rift zone). In addition, both volcanoes share common physical  
320 parameters such as a magma velocity of ~1 m.s<sup>-1</sup> (Swanson et al., 1979; Eaton et al., 1987;  
321 Parfitt & Wilson, 1994; Gauthier et al., 2016). Therefore, the comparison of S/Cl and S/F mass  
322 ratios at both volcanoes is relevant for better understanding degassing processes at basaltic  
323 volcanoes. It can be seen in Figure 3 that syn-eruptive gas samples from the rift zone of Kilauea  
324 and the Bárðarbunga fissure swarm (Holuhraun) have similar S/Cl and S/F ratios. The same  
325 holds true for residual gases as well as the most degassed products collected at Kilauea. In fact,  
326 a continuous trend of decreasing S concentrations is observed from Type II gas of the Kilauea  
327 rift zone (Symonds et al., 1994) towards the Pele's tears of one of the rift-zone craters (Edmonds  
328 et al., 2009). Gas burst and lava spatter data extend this trend to even lower S/halogens. The  
329 simplest explanation is a progressive distillation of S from the rift-zone magma leading to  
330 decreasing solubility and increasing volatility of the halogens.

331  
332 Within this framework, it might seem surprising not to detect at Bárðarbunga-Holuhraun the  
333 equivalent of CO<sub>2</sub>-enriched Type I gas released from the Halema'uma'u crater inside the  
334 Kilauea caldera (Gerlach and Graeber, 1985). Indeed, the gas from Kilauea's summit crater  
335 exhibits the highest recorded S/Cl and S/F ratios (Mather et al., 2012) and it likely represents  
336 the very first step of the Rayleigh distillation process (Figure 3). Whether such primary gas is  
337 absent or hidden within the Bárðarbunga volcanic system is worth considering. Gauthier et al.  
338 (2016) observed that <sup>210</sup>Po, a short-lived isotope from the highly volatile element polonium,  
339 was at unusually low level in the main gas plume released from the Holuhraun craters. Their  
340 preferred interpretation was that earlier degassing of Po, and of CO<sub>2</sub> which was also depleted  
341 in the main gas plume, took place somewhere under the glacier-covered Bárðarbunga caldera  
342 and the magma pathway towards the Holuhraun fissure craters. Because of the ice cover of  
343 Bárðarbunga, such a first-stage CO<sub>2</sub>-enriched gas phase remains obscure and hard to detect.  
344 Nevertheless, ice-cauldrons formed 1-5 km SE of Bárðarbunga during the rifting event in the  
345 second-half of August 2014 (Reynolds et al., 2019), and shortly before the eruption started,  
346 could be the surface manifestations of CO<sub>2</sub>-rich gas burst during magma ascent. Alternatively,  
347 the chimney of seismicity detected 12 km SE of Bárðarbunga caldera and interpreted to be

348 caused by vertical melt migration from 24 km depth towards the brittle-ductile crustal transition  
349 at 9 km could be another candidate for the site of early CO<sub>2</sub> degassing (Hudson et al., 2017).

350

## 351 **Conclusion**

352

353 The different compositions of the syn- and post-eruptive gas emitted during the Holuhraun  
354 2014-2015 eruption is explained by fractional degassing following the Rayleigh distillation law.  
355 Since S is extremely volatile, its partition into the gas phase can be assumed close to  
356 quantitative. In that case, the vapour-melt partition coefficient  $\mathcal{D}^{V/M}$  can be calculated for both  
357 Cl and F from their correlation with S concentrations. Elevated  $\mathcal{D}^{V/M}$  for Cl during the eruption  
358 may reflect the effect of sulphur degassing on Cl concentration, but experimental verification  
359 is needed to test this proposition. In the post-eruptive gas phase,  $\mathcal{D}^{V/M}(\text{Cl})$  is significantly lower,  
360 most likely because S is strongly depleted at the end of the Rayleigh process, and Cl and F reach  
361 similar vapour-melt partition coefficients. Improved knowledge of  $\mathcal{D}^{V/M}$  for minor volatile  
362 species is required for the use of halogens in gas monitoring at active basaltic volcanoes (e.g.,  
363 Aiuppa, 2009) but our study suggests a complex behaviour, partly dependent on sulphur  
364 degassing. Further constraints on  $\mathcal{D}^{V/M}(\text{Cl})$  will also aid interpretation of Cl isotopes, which are  
365 known to fractionate during degassing (Sharp et al., 2010).

366

367 An open-system Rayleigh-type outgassing of the lava field explains elevated F concentrations  
368 measured in the environmental samples close to the lava field. Sustained halogen, and notably  
369 fluorine, degassing from cooling lavas may play a major role in the transportation of volatile  
370 trace elements. The results of this study underline the importance of integrating post-eruptive  
371 degassing processes when quantifying the entire volcanic gas emissions and their impact on the  
372 environment.

373

374 The mass ratios of sulphur over halogens (S/Cl and S/F) in the gas phase illustrate a strong  
375 analogy between degassing processes operating at both Kilauea and Bárðarbunga volcanoes. In  
376 contrast to Kilauea, CO<sub>2</sub>-enriched primary gases were not released into the atmosphere during  
377 the Holuhraun eruption. However, we speculate that the ice cauldrons generated before and  
378 during lateral magma transfer beneath the Vatnajökull ice-sheet resulted from CO<sub>2</sub> gas burst  
379 rather than small sub-glacial eruptions.

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381

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545 **Table 1.** Volatile species concentrations and element mass ratios in post- (prefix HH) and syn-  
 546 eruptive (prefix BARB) gas samples from the 2014-2015 Holuhraun eruption, Bárðarbunga  
 547 volcanic system, Iceland. ‘Blank’ refers to the local atmosphere. Samples BARB-A and -B  
 548 are taken from Gauthier et al. (2016). Residual concentration in whole-rock sample of the degassed  
 549 Holuhraun lava is given as  $\mu\text{g/g}$ .

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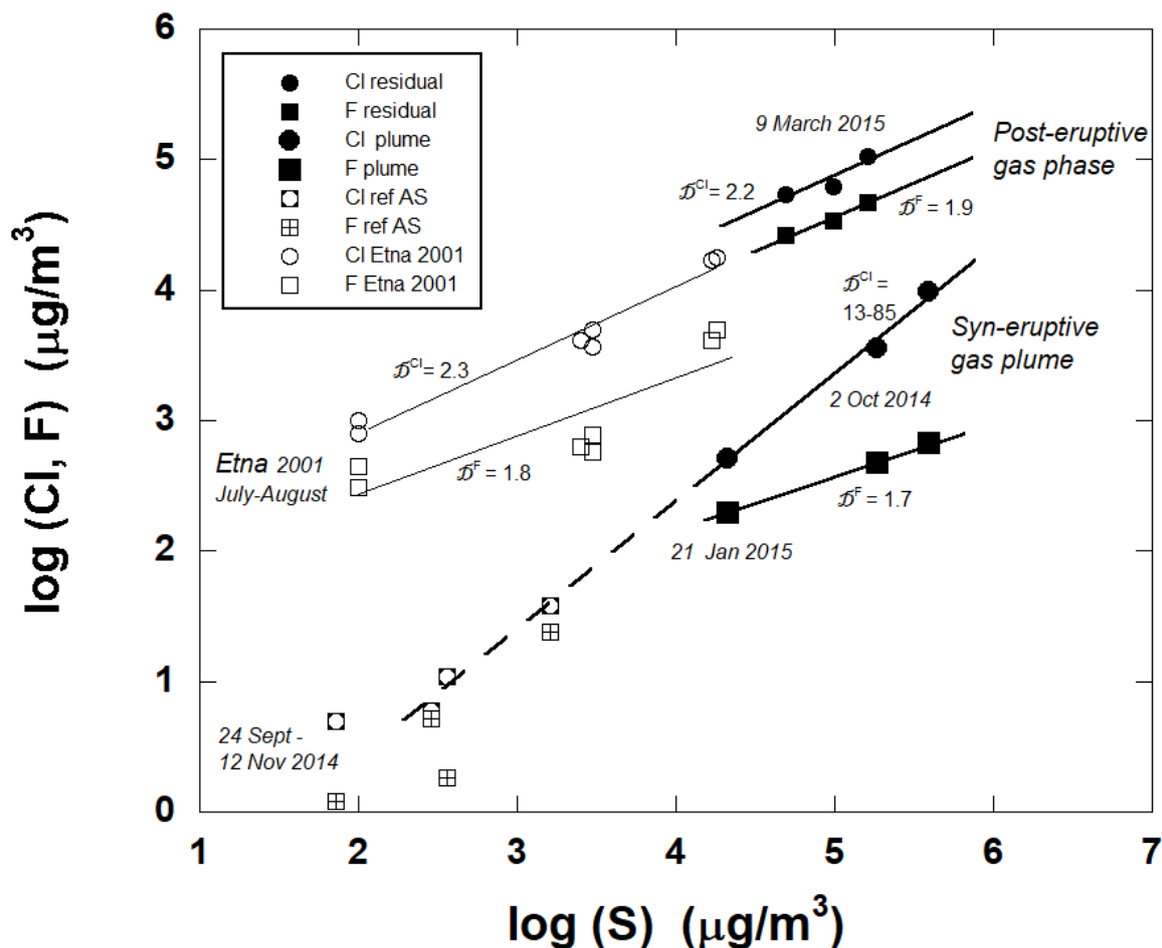
	Lava	BARB-A	BARB-B	Loc2 FP*	HH1	HH2	HH3	Blank
Sampling date	29/08/2014	02/10/2014	02/10/2014	21/01/2015	09/03/2015	09/03/2015	09/03/2015	09/03/2015
Latitude ( $^{\circ}\text{N}$ )	64.85	64.87	64.87	64.89	64.85	64.85	64.85	64.86
Longitude ( $^{\circ}\text{W}$ )	16.86	16.86	16.86	16.8	16.83	16.83	16.83	16.86
S ( $\text{mg/m}^3$ )	97.0	185	391	21.5	97.6	49.0	164	0.323
Cl ( $\text{mg/m}^3$ )	110	3.54	9.72	0.515	61.8	53.2	104	0.720
F ( $\text{mg/m}^3$ )	190	0.475	0.674	0.199	33.7	26.5	46.3	0.517
S/Cl mass	0.88	52	40	42	0.45	1.58	0.92	1.57
Cl/F mass	0.58	7.5	14.4	2.6	1.39	1.83	2.01	2.25
S/F mass	0.51	390	580	108	0.62	2.90	1.85	3.53
SO <sub>2</sub> /HCl molar	0.97	58	44	46	1.74	1.02	1.74	0.50
HCl/HF molar	0.31	4.0	7.7	1.4	0.98	1.08	1.20	0.75
SO <sub>2</sub> /HF molar	0.30	230	340	88	1.72	1.10	2.09	0.37

\*airborne plume from Ilyinskaya et al. (2017)

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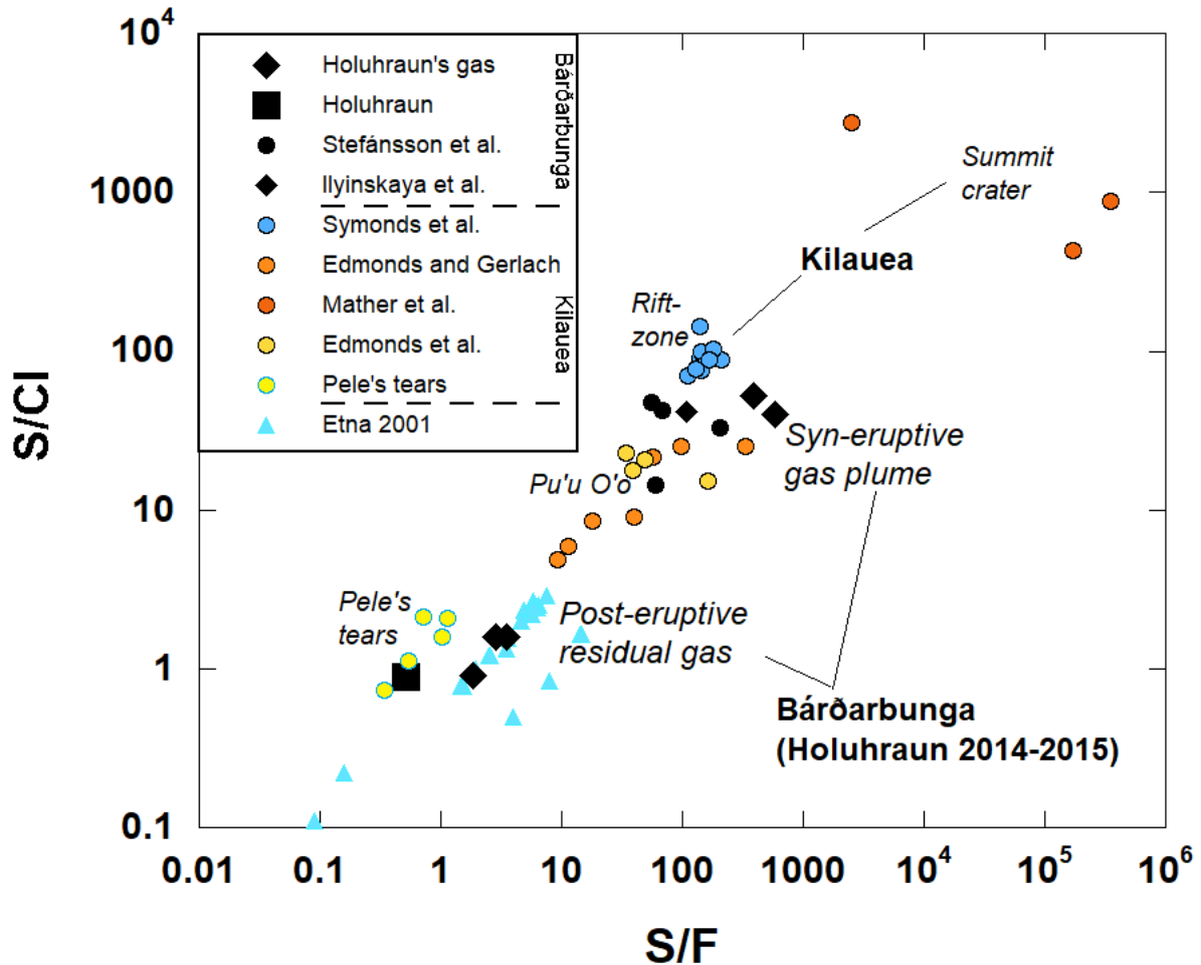


555 **Fig. 1** Grounded gas plume at Holuhraun on 2 October 2014 caused by atmospheric  
 556 temperature inversion. Top left and right insets show, respectively, sampling conditions  
 557 within the syn-eruptive gas plume and of the post-eruptive gas phase inside the eruptive  
 558 fissure on 9 March 2015.  
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**Fig. 2** Halogen vs sulphur concentrations in syn- and post-eruptive gas (plume and residual gas) from Holuhraun. Circles and squares denote Cl and F concentrations, respectively. The post-eruptive gas phase data are from the present study. Filled symbols for the syn-eruptive gas plume data are from Gauthier et al. (2016) and Ilyinskaya et al. (2017), whereas results from Stefánsson et al. (2017; ref AS on plot) are shown with open symbols. The squared correlation coefficient,  $R^2$ , ranges from 0.837 for post-eruptive Cl vs S to 0.999 for syn-eruptive F vs S concentrations. Also plotted are results from Aiuppa et al. (2002) for the 2001 flank eruption on the south slope of Mt. Etna, Sicily. The vapour-melt partition coefficient ( $\mathcal{D}^{\text{V/M}}$ ) shown is simply the reciprocal of  $D^{\text{I/g}}$  calculated from the slope of the linear best-fit and by assuming  $D^{\text{I/g}}$  for  $S = 0$ . See text for further discussion.



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**Fig. 3** Mass ratios of S/Cl versus S/F illustrating the decreasing sulphur concentration from the syn-eruptive gas plume (Gauthier et al., 2016) to the post-eruptive residual gas phase at Holuhraun, Bárðarbunga volcanic system, Iceland. Also displayed are syn-eruptive results from Ilyinskaya et al. (2017) and Stefánsson et al. (2017). The Kilauea summit, rift zone and Pu'u O'o gas and Pele's tears compositions are respectively from Mather et al. (2012), Symonds et al. (1994), Edmonds and Gerlach (2006) and Edmonds et al. (2009). Kilauea rift zone and Pu'u O'o gas are of similar composition as that of Holuhraun, whereas the Pele's tears record the residual gas phase comparable to that of the post-eruptive gas and outgassed lava of Holuhraun. Gas collected during the Etna 2001 eruption displays similar exhaustion of S during fractional degassing at significantly lower S/halogens (Aiuppa et al., 2002).