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1 **The uptake of halogen (HF, HCl, HBr and HI) and nitric (HNO₃) acids**
2 **into acidic sulphate particles in quiescent volcanic plumes**

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9 **Abstract**

10 The uptake of halogen and nitric acids into acidic sulphate particles has important implications for
11 volcano monitoring and the environmental geochemistry of volcanic emissions. Using the Extended
12 Aerosol Inorganics Model (E-AIM) for HCl, HBr and HNO₃, combined with a simple three-parameter
13 model for HF and HI, we show that equilibrium partitioning of halogen and nitric acids into sulphate
14 particles is maximised at high relative humidity (RH), low temperature, low plume dilution (i.e., near-
15 source) and high SO₄²⁻/SO₂. The addition of metal chlorides (i.e., NaCl) enhances acidic gas
16 partitioning at high RH, but diminishes acidic gas partitioning at low RH due to a decrease in the
17 water content of particles and the formation of Na sulphate salts. We predict that acidic gas
18 partitioning is quantitatively significant (>1%) only in cool humid conditions (e.g., >80% RH at 298
19 K), adding confidence to spectroscopic and/or electrochemical determinations of gas ratios (e.g.,
20 SO₂/HCl, HCl/HF) for volcano monitoring. However, acidic gas partitioning remains environmentally
21 important over a wide range of conditions because of the significant variability in the amounts of
22 acidic gases in particle form (Cl⁻(aq), Br⁻(aq), I⁻(aq), NO₃⁻(aq), HF(aq)). This may result in diurnal and
23 seasonal changes in acidic gas deposition downwind of the volcano.

24 **Keywords:** volcanic plume, equilibrium model, masaya, E-AIM
25

26 1 Introduction

27 Active volcanoes are a major source of gases and particles to the atmosphere. A significant
28 contribution to global volcanic emissions comes from persistent degassing from volcanoes such as Mt
29 Etna (Sicily) and Masaya (Nicaragua). The activity at these volcanoes is characterised by the
30 sustained emission of volatile species (forming a quiescent volcanic plume) punctuated by sporadic
31 and typically minor eruptions. Volcanic gases are composed of > 90% H₂O and CO₂, and < 10% SO₂,
32 HCl and HF, with smaller and variable amounts of H₂S, H₂, CO, HBr and HI (Gerlach, 2004; Pyle and
33 Mather, 2009). A common feature of quiescent volcanic plumes is micron-sized sulphate particles,
34 with compositions between the acidic (e.g., H₂SO₄) and metal-rich (primarily (Na,K)₂SO₄) end-
35 members (e.g., Mather et al., 2003; Martin et al., 2011) at a molar abundance of SO₄²⁻/SO₂ ~ 0.01.

36 The uptake of halogen acids (HF, HCl, HBr, HI) into sulphate particles has wide implications in
37 volcanology. The magmatic SO₂/HCl gas ratio plays an important role in eruption forecasting and in
38 understanding volcanic degassing (e.g., Horrocks et al., 1999; Duffell et al., 2003) due to the SO₂/HCl
39 ratio decreasing as the magma becomes more degassed. However, the magmatic SO₂/HCl ratio may
40 be overprinted by preferential HCl uptake into sulphate particles. More generally, if uptake into
41 particles is significant, volcanic emissions of F, Cl, Br and/or I (Pyle and Mather, 2009) may be
42 underestimated if the gas and particle phases are not monitored together (as is often the case). Recent
43 measurements indicate ~1% of total HCl and HF are taken up into particles within the first few
44 minutes after emission (e.g., Mather et al., 2006a; Martin et al., 2008, 2010).

45 The uptake of halogen acids into particles is also environmentally important because particles have a
46 greater potential for deposition than gases, with the rate of sedimentation increasing with particle size.
47 In fact, uptake may control the extent of local deposition for halogen acids since volcanic plumes tend
48 to be emitted and transported at altitudes above the elevation of the surrounding land. At Mt Etna
49 (Sicily), significant airborne concentrations of particles were measured ~2 km below the plume (Allen
50 et al., 2006), consistent with a number of studies measuring or inferring deposition onto the volcano's
51 flanks (e.g., Aiuppa et al., 2006; Bellomo et al., 2007; Watt et al., 2009). At Masaya volcano

52 (Nicaragua), the environmental impacts of deposition include acidification of soils and a decrease in
53 vegetation health and diversity (e.g., Delmelle et al., 2002). Uptake may also influence local
54 atmospheric chemistry, due to the enhanced rates of chemical reactions within, or on the surface of,
55 particles. Of special interest is the autocatalytic bromine explosion, which results in the production of
56 reactive Br species (e.g., Br, BrO) and rapid O₃ depletion (Oppenheimer et al., 2006; von Glasow,
57 2010).

58 In contrast to the halogen acids, nitric acid (HNO₃) is not thought to be a primary magmatic emission
59 (Mather et al., 2004). Equilibrium models predict that NO is produced when atmospheric N₂ and O₂
60 are heated to > 1000°C at the vent (Gerlach, 2004; Martin et al., 2006). The minor amounts of NO₂
61 and HNO₃ formed under such conditions cannot explain measurements of high HNO₃/SO₂ (e.g., 0.02
62 at Masaya (Nicaragua), 0.05 at Etna (Sicily), Mather et al., 2004; 0.03 at Erebus (Antarctica),
63 Oppenheimer et al., 2010). Instead, it is supposed that NO is oxidised rapidly to HNO₃ at ambient
64 temperature in the volcanic plume (Roberts et al., 2009; Boichu et al., 2011), although the mechanism
65 for this process remains unclear. In any case, HNO₃ uptake will likely promote N deposition and this
66 may be important in N-limited ecosystems (Huebert et al., 1999).

67 The equilibrium limit of uptake is defined by “partitioning”, which may be calculated without
68 reference to any time-dependent (i.e., kinetic) processes. While the thermodynamic Extended Aerosol
69 Inorganics Model (E-AIM; Carslaw et al., 1995; Clegg et al., 1998; Wexler and Clegg, 2002) has been
70 used to predict the equilibrium composition of volcanic aerosol on a case-by-case basis (e.g., Mather
71 et al., 2004; Oppenheimer et al., 2006; Martin et al., 2008; Roberts et al., 2009; Martin et al., 2010),
72 the general features of acidic gas partitioning into acidic sulphate particles have yet to be established.
73 In this study we use E-AIM to investigate the extent of HCl, HBr and HNO₃ partitioning over a wide
74 range of atmospheric (i.e., relative humidity, temperature) and plume (i.e., dilution, composition)
75 conditions. A simple three-parameter model is used to extend these results to HF and HI partitioning.
76 The effects of dissolved Na⁺ ions on HCl partitioning are also investigated. These results offer new

77 insights into the uptake of halogen and nitric acids into acidic sulphate particles, and we discuss a
78 range of implications for volcanology and the environmental geochemistry of volcanic emissions.

79 **2 Methodology**

80 **2.1 The equilibrium assumption**

81 Particles in quiescent volcanic plumes are mostly secondary in origin (i.e., non-silicate) and are
82 formed in different temperature regimes (Óskarsson, 1980). The mixing of magmatic and atmospheric
83 gases at the vent ($> 700^{\circ}\text{C}$) results in cooling and condensation of metal salts (e.g., NaCl, KCl, CaCl₂,
84 MgCl₂, with smaller amounts of the corresponding fluorides; e.g., Symonds et al., 1992; Symonds and
85 Reed, 1993). Additionally, $\sim 1\%$ of SO₂ is oxidised to hygroscopic SO₃ by atmospheric O₂ at the vent
86 (Mather et al., 2006b). With further cooling ($< 338^{\circ}\text{C}$), SO₃ reacts with H₂O to form gas-phase H₂SO₄,
87 and subsequently droplets of H₂SO₄. The composition of particles measured at the crater-rim will
88 reflect not only the initial amounts of metal salts and H₂SO₄, but the extent of internal mixing between
89 these types of particles, and the interactions between particles and gases (e.g., H₂O, HF, HCl, HBr,
90 HI, HNO₃). The dynamics of this sequence are, of course, hugely complex, with significant potential
91 for non-equilibrium. Due to the extreme hygroscopicity of H₂SO₄, along with the high abundances of
92 gas-phase H₂SO₄ and H₂O, binary homogenous nucleation of H₂SO₄ and H₂O may occur, in addition
93 to the heterogeneous nucleation of H₂SO₄ onto metal salts. Single particle analyses (e.g., Ammann
94 and Burtscher et al., 1990; Ammann et al., 1993; Toutain et al., 1995; Martin et al., 2008; Moune et
95 al., 2010) show a range of compositions indicating that internal mixing is neither negligible nor
96 complete. Thus, while local equilibrium may be established between gases and particles over
97 timescales as short as a few seconds (Wexler and Seinfeld, 1992), the overall gas and particle
98 composition may remain out of equilibrium.

99 E-AIM (along with other aerosol thermodynamic models, e.g., ISORROPIA II, Fountoukis and
100 Nenes, 2007) assumes simply that particles are at equilibrium with their surroundings, exist as an
101 internal mixture, and are sufficiently large ($> 0.1 \mu\text{m}$) that they behave as bulk solids or liquids (i.e.,

102 the Kelvin effect is negligible). Despite these simplifications, thermodynamic calculations are
 103 valuable because they allow the influence of different variables (e.g., temperature, relative humidity,
 104 input composition) to be established. Furthermore, if equilibrium has not been attained at the crater-
 105 rim, equilibrium may be attained further downwind. It is in these older quiescent plumes that rapid
 106 heterogeneous oxidation processes (e.g., the bromine explosion) are thought to occur (e.g.,
 107 Oppenheimer et al., 2006, 2010; Bobrowski et al., 2007; Roberts et al., 2009; von Glasow, 2010) so
 108 our results have wider relevance.

109 2.2 Modelling

110 The variants of E-AIM used are E-AIM1 (Carslaw et al., 1995), which considers the H^+ - SO_4^{2-} - NO_3^-
 111 - Cl^- - Br^- - H_2O system over 200 to 330 K, and E-AIM3 (Clegg et al., 1998), which considers the H^+ -
 112 Na^+ - SO_4^{2-} - NO_3^- - Cl^- - H_2O system at 298.15 K. Both variants calculate the equilibrium
 113 composition for a given relative humidity (RH), temperature (T) and input composition.

114 Our “standard” input composition is based on a mean of 13 gas and particle analyses made at the
 115 crater rim of Masaya volcano (Nicaragua) during a period of quiescent degassing (Martin et al., 2010)
 116 (Table 2, n_i ; in $\mu\text{mol m}^{-3}$). SO_4^{2-} and HSO_4^- were not distinguished during the chemical analysis so the
 117 sum of these ions is represented by $n_{\text{SO}_4^{2-}(\text{aq},\text{s})}$ in Table 2. This data set was chosen as it is one of the
 118 few simultaneous measurements of halogen and nitric acid gases and particles in a quiescent volcanic
 119 plume. The total amount of each ion was fixed (i.e., mass balanced) using measurements of the most
 120 abundant species for Cl^- , Br^- , NO_3^- and SO_4^{2-} ($n_{\text{HCl}(\text{g})}$, $n_{\text{HBr}(\text{g})}$, $n_{\text{HNO}_3(\text{g})}$, $n_{\text{SO}_4^{2-}(\text{aq},\text{s})}$; in mol m^{-3} of air). K^+ ,
 121 Ca^{2+} and Mg^{2+} are not considered within E-AIM3 so these ions were entered as “equivalent” Na^+ (i.e.,
 122 $n_{\text{Na}^{+*}(\text{aq},\text{s})} = n_{\text{Na}^+(\text{aq},\text{s})} + n_{\text{K}^+(\text{aq},\text{s})} + 2n_{\text{Mg}^{2+}(\text{aq},\text{s})} + 2n_{\text{Ca}^{2+}(\text{aq},\text{s})}$; Moya et al., 2001). $n_{\text{H}^+(\text{aq},\text{s})}$ was entered to
 123 balance charge. E-AIM considers only the most major gas and particle species of each ion (i.e.,
 124 $\text{H}_2\text{O}(\text{g})$, $\text{HCl}(\text{g})$, $\text{HBr}(\text{g})$, $\text{HNO}_3(\text{g})$, $\text{Cl}^-(\text{aq})$, $\text{Br}^-(\text{aq})$, $\text{NO}_3^-(\text{aq})$, $\text{SO}_4^{2-}(\text{aq})$, $\text{HSO}_4^-(\text{aq})$, $\text{H}^+(\text{aq})$ in E-
 125 AIM1; $\text{H}_2\text{O}(\text{g})$, $\text{HCl}(\text{g})$, $\text{HNO}_3(\text{g})$, $\text{Cl}^-(\text{aq})$, $\text{NO}_3^-(\text{aq})$, $\text{SO}_4^{2-}(\text{aq})$, $\text{HSO}_4^-(\text{aq})$, $\text{H}^+(\text{aq})$, $\text{Na}^{+*}(\text{aq})$ and salts
 126 of Na^{+*} in E-AIM3). Minor species, such as the associated aqueous forms ($\text{H}_2\text{SO}_4(\text{aq})$, $\text{HCl}(\text{aq})$,
 127 $\text{HBr}(\text{aq})$, $\text{HNO}_3(\text{aq})$) and the extremely hygroscopic $\text{H}_2\text{SO}_4(\text{g})$, are not considered within E-AIM.

128 The variables to be investigated within E-AIM1 are: (1) relative humidity, (2) ambient temperature,
 129 (3) the amounts of SO_4^{2-} and halogen and nitric acids (reflecting differences in SO_3 production at the
 130 vent, and differences in the magmatic gas composition) and (4) plume dilution. The variables to be
 131 investigated within E-AIM3 are: (1) relative humidity and (2) the amount of Na^+ (as a proxy for a
 132 wider range of metal ions). Results are independent of the total atmospheric pressure because at fixed
 133 temperature RH defines the partial pressure of H_2O ($p_{\text{H}_2\text{O}(\text{g})}$; in atm) and the input composition defines
 134 the partial pressure of HCl, HBr and HNO_3 ($p_{\text{HCl}(\text{g})}$, $p_{\text{HBr}(\text{g})}$ and $p_{\text{HNO}_3(\text{g})}$; in atm).

135 We also use a three-parameter model for equilibrium partitioning (e.g., Mather et al., 2003), which
 136 considers non-dissociative dissolution of gas (HX) (i.e., Henry's law, $K_{\text{H:X}}$; E1A) and the subsequent
 137 dissociation into ions (i.e., acidity constants, $K_{\text{A:X}}$; E1B). The concentrations of aqueous species are
 138 defined by activity (a_i ; in mol dm^{-3}), while the concentrations of gaseous species are defined by partial
 139 pressure (p_i ; in atm). The total concentration of X in the particles is defined by $a_{\text{X}(\text{aq})}$ (i.e., $a_{\text{X}(\text{aq})} +$
 140 $a_{\text{HX}(\text{aq})}$) (E2). Literature values of $K_{\text{H:X}}$ and $K_{\text{H:A}}$ are given in Table 1. The three-parameter model
 141 shows that $a_{\text{X}(\text{aq})}$ increases with increasing $K_{\text{H:X}}$ and $K_{\text{H:A}}$, and decreases with increasing $a_{\text{H}^+(\text{aq})}$. In
 142 other words, acidic gas partitioning will decrease as the particles become more acidic.



E1A
$$K_{\text{H:X}} = \frac{a_{\text{HX}(\text{aq})}}{p_{\text{HX}(\text{g})}}$$



E1B
$$K_{\text{A:X}} = \frac{a_{\text{H}^+(\text{aq})} a_{\text{X}^-(\text{aq})}}{a_{\text{HX}(\text{aq})}}$$

145 E2
$$\frac{a_{\text{X}(\text{aq})}}{p_{\text{HX}(\text{g})}} = K_{\text{H:X}} \left(1 + \frac{K_{\text{A:X}}}{a_{\text{H}^+(\text{aq})}} \right)$$

146 The complexity of the E-AIM model does not allow for a full uncertainty analysis although the
 147 uncertainty in the outputs will depend on the uncertainty of the thermodynamic database (Wexler and
 148 Clegg, 2002). In general, validation of E-AIM (and other aerosol thermodynamic models) is
 149 performed by comparison with field or laboratory measurements over a range of conditions (e.g., Yao
 150 et al., 1999; Wexler and Clegg, 2002; Clegg et al., 2008). The uncertainty within the three-parameter
 151 model (E2) can be considered within different regimes: (1) $K_{\text{A:X}}/a_{\text{H}^+(\text{aq})} < 1$ and (2) $K_{\text{A:X}}/a_{\text{H}^+(\text{aq})} > 1$.

152 The uncertainty in E2 is therefore equal to the uncertainty on $K_{H:X}$ or on the product $K_{A:X}K_{H:X}$
 153 (respectively). For strong acids, the product $K_{A:X}K_{H:X}$ is much better constrained than either $K_{A:X}$ and
 154 $K_{H:X}$ alone (i.e., there may be 1-2 orders of magnitude uncertainty on $K_{A:X}$ and $K_{H:X}$) because
 155 measurement of $K_{A:X}K_{H:X}$ does not require determination of $a_{HX(aq)}$ (only $p_{HX(g)}$, $a_{X-(aq)}$ and $a_{H+(aq)}$).
 156 Many more minor halogen species (e.g., $Br_2(aq)$, $Cl_2(aq)$, $I_2(aq)$, $BrCl(aq)$, $Br_2Cl^-(aq)$, $I_3^-(aq)$,
 157 $(HF)_n(aq)$) are not considered within the three-parameter model. Calculations of $a_{X(aq)}/p_{HX(g)}$ are
 158 therefore lower limits to the true $a_{X(aq)}/p_{HX(g)}$ since only $a_{HX(aq)}$ and $a_{X-(aq)}$ are considered.

159 3 Results and Discussion

160 E-AIM1 was used to estimate the equilibrium composition of volcanic aerosol as a function of RH for
 161 the standard input composition at 298 K. Results for equilibrium amounts (n_i ; in mol m⁻³ of air) and
 162 concentrations (c_i ; in mol dm⁻³ of solution) of aqueous species are shown in Figure 1. By absolute
 163 amount, the most important change with increasing RH is increasing $n_{H_2O(aq)}$, resulting in dilution of
 164 the aqueous phase and an increase in particle size. The responses of different chemical equilibria to
 165 dilution result in complex compositional changes. H^+ and HSO_4^- are the most abundant ions at low
 166 RH and exist in approximately equal amounts. Dilution of the aqueous phase promotes the
 167 dissociation of HSO_4^- into SO_4^{2-} and H^+ although the net effect is decreasing $c_{H+(aq)}$ (and thus
 168 increasing pH) with increasing RH.

169 The extent of HCl partitioning, which is represented by $n_{Cl-(aq)}$, is quantitatively minor (<1% of initial
 170 $n_{HCl(g)}$) below 80% RH but near-complete (>95%) above 95% RH (Figure 1). The increase in HCl
 171 partitioning above 80% RH (which we denote RH_{crit}) reflects $Cl^-(aq)$ becoming the most abundant
 172 anion (i.e., $n_{Cl-(aq)} > n_{SO_4^{2-}(aq)} + n_{HSO_4^-(aq)}$). The significance is that changes in $n_{Cl-(aq)}$ may now influence
 173 $n_{H_2O(aq)}$, or equivalently, that particle hygroscopicity is controlled by HCl rather than H_2SO_4 . In this
 174 regime, $n_{Cl-(aq)}$ and $n_{H_2O(aq)}$ both increase with RH, allowing a large change in $n_{Cl-(aq)}$ for only a small
 175 change in $c_{Cl-(aq)}$. Eventually, the increase in $n_{H_2O(aq)}$ exceeds the increase in $n_{Cl-(aq)}$ as the supply of
 176 $HCl(g)$ is exhausted, resulting in decreasing $c_{Cl-(aq)}$. The similarity between trends in HCl and HNO_3
 177 partitioning (i.e., $n_{NO_3-(aq)}$) above RH_{crit} reflects the influence of HCl on $n_{H_2O(aq)}$, while the differences

178 at low RH reflect greater $K_{H:X}$ for HNO_3 than HCl. HBr partitioning (i.e., $n_{\text{Br}-(\text{aq})}$) occurs more readily
179 than HCl or HNO_3 partitioning, being minor below 60% RH and near-complete above 85% RH. Thus,
180 at low RH, $n_{\text{Br}-(\text{aq})}/n_{\text{Cl}-(\text{aq})}$ is greater than initial $n_{\text{HBr}(\text{g})}/n_{\text{HCl}(\text{g})}$ (and, equivalently, initial $p_{\text{HBr}(\text{g})}/p_{\text{HCl}(\text{g})}$),
181 implying a relative enrichment of the particle phase with Br. These differences reflect the greater
182 acidity of HBr than either HCl or HNO_3 (Table 1), which allows for HBr partitioning into the highly
183 acidic particle phase present below RH_{crit} . The increase in $n_{\text{H}_2\text{O}(\text{aq})}$ at RH_{crit} has little effect on $n_{\text{Br}-(\text{aq})}$,
184 which is already close to its maximum value (defined by initial $n_{\text{HBr}(\text{g})}$).

185 E-AIM1 calculations were also made at 283 K and 313 K (Figure 1). The 283 K - 313 K temperature
186 range was chosen because it is centred on 298 K (allowing comparison of results between E-AIM1
187 and E-AIM3) and spans the range of ambient temperatures recorded at Masaya (e.g., Mather et al.,
188 2003, 2006). Below RH_{crit} , $n_{\text{Cl}-(\text{aq})}$, $n_{\text{Br}-(\text{aq})}$ and $n_{\text{NO}_3-(\text{aq})}$ decrease by a factor of 10 between 283 K and
189 313 K, with the direction of the trend being consistent with the exothermic nature of the process (i.e.,
190 the sum of R1A and R1B). $n_{\text{Cl}-(\text{aq})}$ and $n_{\text{NO}_3-(\text{aq})}$ are even more sensitive to temperature around RH_{crit} ,
191 because RH_{crit} itself increases with temperature.

192 To investigate the different influences of (input) SO_4^{2-} and HCl on acidic gas partitioning, E-AIM1
193 calculations were made for the standard input composition at 298 K, with initial $n_{\text{SO}_4^{2-}(\text{aq,s})} = 0, 0.2, 2,$
194 $20 \mu\text{mol m}^{-3}$ (Figure 2A), and separately, with initial $n_{\text{HCl}(\text{g})} = 0, 40, 400, 4000 \mu\text{mol m}^{-3}$ (Figure 2B).
195 For simplicity, only $n_{\text{Cl}-(\text{aq})}$ and $n_{\text{H}_2\text{O}(\text{aq})}$ are shown in each Figure. Figure 2A (i.e., variable SO_4^{2-})
196 shows that below RH_{crit} , both $n_{\text{H}_2\text{O}(\text{aq})}$ and $n_{\text{Cl}-(\text{aq})}$ are proportional to SO_4^{2-} , hence RH_{crit} is independent
197 of SO_4^{2-} . Above RH_{crit} , $n_{\text{Cl}-(\text{aq})}$ is independent of SO_4^{2-} , consistent with particle hygroscopicity being
198 controlled by HCl rather than H_2SO_4 . Figure 2B (i.e., variable HCl) shows that decreasing HCl results
199 in increasing RH_{crit} , since a higher RH must be reached before $\text{Cl}(\text{aq})$ becomes the most abundant
200 anion. Consistent with our interpretation, $n_{\text{H}_2\text{O}(\text{aq})}$ increases with HCl above RH_{crit} , but is independent
201 of HCl below RH_{crit} . An additional calculation was made for the standard input composition with
202 initial $n_{\text{HCl}(\text{g})} = n_{\text{HBr}(\text{g})} = n_{\text{HNO}_3(\text{g})} = 0 \mu\text{mol m}^{-3}$. Here, $n_{\text{H}_2\text{O}(\text{aq})}$ was similar to $n_{\text{H}_2\text{O}(\text{aq})}$ calculated for the

203 standard input composition with initial $n_{\text{HCl(g)}}=0 \mu\text{mol m}^{-3}$, indicating that neither HBr nor HNO_3 have
 204 a significant influence on $n_{\text{H}_2\text{O(aq)}}$.

205 Plume dilution has the effect of decreasing SO_4^{2-} and HCl together. To investigate this influence on
 206 acidic gas partitioning, E-AIM1 calculations were made with $D/D_0 = 0.1, 1, 10, 100$ (where D_0 is the
 207 dilution at the crater rim) and the standard input composition at 298 K (Figure 2C). Input
 208 concentrations are proportional to D^{-1} , except $p_{\text{H}_2\text{O(aq)}}$, which is fixed by RH. We assume that the
 209 aerosol volume within the background atmosphere is negligible. Below RH_{crit} , $n_{\text{H}_2\text{O(aq)}}$ is proportional
 210 to SO_4^{2-} (Figure 2A) and hence proportional to D^{-1} . In contrast, $n_{\text{Cl-(aq)}}$ is proportional to D^{-2} , because
 211 both $n_{\text{H}_2\text{O(aq)}}$ and $n_{\text{HX(g)}}$ are proportional to D^{-1} . Thus, a 10-fold plume dilution ($D/D_0=10$) results in a
 212 10-fold decrease in $n_{\text{H}_2\text{O(aq)}}$ and a 10-fold decrease in $n_{\text{HCl(g)}}$, resulting in a 100-fold decrease in $n_{\text{Cl-(aq)}}$.
 213 These factors affect RH_{crit} so $n_{\text{Cl-(aq)}}$ is highly sensitive to D around RH_{crit} . A limitation of the model is
 214 that as D/D_0 increases, SO_4^{2-} tends to zero, rather than the true atmospheric background, resulting in
 215 an underestimate of partitioning at high D/D_0 .

216 E-AIM3 was used to estimate the equilibrium composition of volcanic aerosol as a function of RH
 217 and equivalent Na^{*+} ($n_{\text{Na}^{*+}(\text{aq,s})}$, i.e., the total amount of Na^{*+} in the aqueous and solid phases) for the
 218 standard input composition at 298 K. Calculations were made by introducing equal amounts of Na^{*+}
 219 and Cl, simulating the addition of NaCl to acidic sulphate particles (e.g., Óskarsson, 1980). Results
 220 for equilibrium amounts of aqueous species and salts are shown in Figure 3. Equilibrium amounts of
 221 Na-sulphate salts ($\text{NaHSO}_4 \cdot \text{H}_2\text{O}$, $\text{Na}_3\text{H}(\text{SO}_4)_2$, Na_2SO_4) are summed and shown as “other $\text{Na}^*(\text{s})$ ” on
 222 Figure 3. The system shows complex behaviour because of salt formation. At low RH (e.g., 50%),
 223 increasing $n_{\text{Na}^{*+}(\text{aq,s})}$ results in decreasing $n_{\text{H}_2\text{O(aq)}}$, $n_{\text{H+}(\text{aq})}$, $n_{\text{Cl-(aq)}}$ and $n_{\text{NO}_3\text{-(aq)}}$. Further increases in
 224 $n_{\text{Na}^{*+}(\text{aq,s})}$ result in the crystallisation of less hygroscopic Na-sulphate salts rather than aqueous
 225 solutions. The net result is that HCl partitioning decreases with increasing $n_{\text{Na}^{*+}(\text{aq,s})}$ due to a decrease
 226 in the water content of the particle, and only increases when all SO_4^{2-} has been associated with Na^{*+}
 227 and NaCl forms (i.e., the stable assemblage is $\text{Na}_2\text{SO}_4 + \text{NaCl}$). HNO_3 partitioning does not increase
 228 with increasing $n_{\text{Na}^{*+}(\text{aq,s})}$ because no Na-nitrates are formed (although these are considered within E-

229 AIM3). In contrast, at high RH (e.g., 75%), increasing $n_{\text{Na}^+(\text{aq},\text{s})}$ results in increasing $n_{\text{H}_2\text{O}(\text{aq})}$, $n_{\text{Cl}^-(\text{aq})}$,
 230 $n_{\text{NO}_3^-(\text{aq})}$ and decreasing $n_{\text{H}^+(\text{aq})}$. No salts are formed under these conditions as RH exceeds the
 231 deliquescence RH (DRH) of all salts formed at lower RH (i.e., Na-sulphates, NaCl). The decreasing
 232 $n_{\text{H}^+(\text{aq})}$ reflects the volatilisation of HCl, shown by the initially greater increase of $n_{\text{Na}^+(\text{aq})}$ than $n_{\text{Cl}^-(\text{aq})}$
 233 with increasing $n_{\text{Na}^+(\text{aq},\text{s})}$. We suggest that HBr (not considered within E-AIM3) and other acidic gases
 234 will show similar trends to HCl and HNO₃.

235 Figure 4 shows calculations for HCl, HBr and HNO₃ partitioning, expressed as $a_{\text{X}(\text{aq})}/p_{\text{HX}(\text{g})}$, from the
 236 three-parameter model (E2). Also shown are calculations for HCl, HBr and HNO₃ partitioning from
 237 E-AIM1. A good agreement is found between results from the three-parameter model and E-AIM1,
 238 and the non-consideration of HBr(aq) and HCl(aq) within E-AIM1 appears justified in the case of
 239 highly acidic sulphate particles. However, HNO₃ partitioning into acidic sulphate particles may be
 240 underestimated within E-AIM1 because HNO₃(aq) is not considered (only NO₃⁻(aq)). In the three-
 241 parameter model, $a_{\text{X}(\text{aq})}/p_{\text{HX}(\text{g})}$ for HNO₃ stabilises above $K_{\text{A}:\text{NO}_3}$ (corresponding to below ~60% RH)
 242 because HNO₃(aq) is present in greater abundance than NO₃⁻(aq). We may speculate that HNO₃(aq)
 243 was not included within E-AIM1 because HNO₃(aq) is unstable in other less acidic atmospheric
 244 aerosols. Hence, while the uncertainties on $a_{\text{X}(\text{aq})}/p_{\text{HX}(\text{g})}$ within E-AIM1 for HCl and HBr are small and
 245 only relate to the uncertainties on the product $K_{\text{A}:\text{X}}K_{\text{H}:\text{X}}$, the uncertainties on $a_{\text{X}(\text{aq})}/p_{\text{HX}(\text{g})}$ for HNO₃ (at
 246 low RH) are potentially much larger. However, since NO₃ is a minor component of the particle phase
 247 in volcanic emissions (e.g., Table 2), underestimation of $a_{\text{X}(\text{aq})}/p_{\text{HX}(\text{g})}$ for HNO₃ is unlikely to
 248 compromise E-AIM results for other species.

249 HBr and HI partitioning (Figure 4) are predicted to be similar in terms of $a_{\text{X}(\text{aq})}/p_{\text{HX}(\text{g})}$ due to the
 250 similarity in the product $K_{\text{H}:\text{X}}K_{\text{A}:\text{X}}$ (Table 1). This result suggests that an estimate of $n_{\text{I}^-(\text{aq})}$ can be made
 251 by substituting HI for HBr within the E-AIM1 model and calculating $n_{\text{I}^-(\text{aq})}$ as $n_{\text{Br}^-(\text{aq})}$. This
 252 approximation is justified because the differences in the activity coefficients (γ) of Br⁻(aq) and I⁻(aq)
 253 are small over a wide range of conditions (e.g., Hamer and Wu, 1972; Brimblecombe and Clegg,
 254 1988) and because HBr has no influence on $n_{\text{H}_2\text{O}(\text{aq})}$ (Figure 2B). Within the three-parameter model,
 255 HF partitioning is predicted to be independent of a_{H^+} over a wide range of RH (<99%) because

256 HF(aq) is more abundant than F(aq) in acidic particles. $n_{\text{HF(aq)}}$ can be estimated from E1A,
 257 substituting $a_{\text{HF(aq)}} = \gamma n_{\text{HF(aq)}}/V$, where V is the aerosol volume, and $p_{\text{HF(g)}} = \beta(n_{\text{HF(g):0}} - n_{\text{HF(aq)}})$, where
 258 $n_{\text{HF(g):0}}$ is initial $n_{\text{HF(g)}}$ (Table 2) and β is the conversion factor between 1 atm and 1 mol m^{-3} at 298 K.
 259 This approximation (E3) assumes that HF partitioning influences neither V nor the activity of other
 260 components of the particle phase, and is justified if $n_{\text{HF(aq)}}$ remains a minor component of the particle
 261 phase. HCl, HBr and HI have large γ at low RH (> 100), while HF has a small γ at low RH (e.g., < 1),
 262 which would promote uptake according to E3.

263 E3
$$n_{\text{HF(aq)}} = \frac{(\beta K_{\text{HF}} V)}{(\gamma + \beta K_{\text{HF}} V)} n_{\text{HF(g):0}}$$

264 Figure 4 shows estimates for HI partitioning (assuming $\gamma_{\text{HI}} = \gamma_{\text{HBr}}$) and HF partitioning (assuming
 265 $\gamma_{\text{HF}} = 1$) into volcanic aerosol as a function of RH for the standard input composition at 298 K. HCl,
 266 HBr and HNO_3 partitioning are shown for comparison. At low RH, the extent of HF partitioning is
 267 greater than that of the other halogens, which have small $K_{\text{H:X}}$ but large $K_{\text{A:X}}$. Increasing RH has little
 268 influence on HF partitioning until RH_{crit} is reached, when V increases significantly, causing an
 269 increase in $n_{\text{HF(aq)}}$. Within the approximation, HI and HBr partitioning are proportional over all RH.
 270 This simply reflects the constancy of the ratio $a_{\text{X(aq)}}/p_{\text{HX(g)}}$ (E2) with fixed $a_{\text{H+ (aq)}}$. These results
 271 indicate that at low RH, the particle phase is enriched in F, and to a lesser extent Br and I, relative to
 272 the gas-phase composition (Table 2).

273 3.3 Implications for volcanology and environmental geochemistry

274 In summary, the equilibrium partitioning of acidic gases into acidic sulphate particles is maximised at
 275 high RH and low temperature, in near-source (i.e., less dilute) plumes with high $\text{SO}_4^{2-}/\text{SO}_2$. The
 276 addition of NaCl enhances acidic gas partitioning at high RH, but diminishes acidic gas partitioning at
 277 low RH due to the formation of Na-sulphate salts.

278 HCl partitioning is predicted to exceed 1% of total HCl at $\sim 80\%$ RH and be near-complete ($>95\%$)
 279 above 95%, indicating that spectroscopic or electrochemical measurements of magmatic gas-phase

280 SO₂/HCl from the crater-rim are robust below 80% RH. At higher RH, HCl partitioning may result in
281 erroneously high measured SO₂/HCl ratios. This effect is intensified by the fact that high RH is often
282 associated with lower temperatures (e.g., at night-time). Burton et al., (2001) reported an apparent loss
283 of 38% HCl to the particle phase, by comparison of SO₂/HCl ratios in the day-time (302 K, 42% RH,
284 using solar occultation FTIR spectroscopy) and the night-time (292 K, 93% RH, using lunar
285 occultation FTIR spectroscopy). While their model for HCl partitioning was incomplete (i.e., H₂SO₄
286 was not considered), our modelling results support their conclusions that the difference in SO₂/HCl
287 may be explained in terms of HCl partitioning. HNO₃, HBr and HI are not measured routinely in the
288 field, although partitioning is also predicted to be significant above ~80% RH. These results should be
289 taken into account when planning measurement strategies or when interpreting data from continuous
290 gas monitoring.

291 Over a wide range of atmospheric conditions acidic gas partitioning has negligible influence on gas-
292 phase concentrations, e.g., $p_{\text{HCl(g)}}$, $p_{\text{HBr(g)}}$, $p_{\text{HI(g)}}$, $p_{\text{HNO}_3\text{(g)}}$ and $p_{\text{HF(g)}}$. However, a small change in
293 temperature and/or RH may significantly affect the amounts of particle forms of acidic gases, e.g., $n_{\text{Cl-}}$
294 $_{\text{(aq)}}$, $n_{\text{Br-}}_{\text{(aq)}}$, $n_{\text{I-}}_{\text{(aq)}}$, $n_{\text{NO}_3\text{-}}_{\text{(aq)}}$ and $n_{\text{HF(aq)}}$. This would result in increased acidic gas partitioning in the night-
295 time, when temperatures are typically lower and RH higher, consistent with clear diurnal trends in $n_{\text{Cl-}}$
296 $_{\text{(aq)}}$ at Masaya (Mather et al., 2003). Furthermore, since particle size also increases at high RH due to
297 water uptake (e.g., Figure 1), the rate of acidic gas deposition would be expected to increase during
298 the night-time. Equally, as RH is high during the wet season (April - October), acidic gas deposition
299 may be enhanced at certain times of the year, which may impact negatively on seasonal crop growth
300 (e.g., Delmelle et al., 2002). This seasonal influence is potentially shown by measurements of $n_{\text{Cl-}}_{\text{(aq,s)}}$
301 at Masaya, which were lower than background in March/April (in 2007, 2009 and 2010; Martin et al.,
302 2011), and higher than background in December (2001; Mather et al., 2003). Our results indicate that
303 the relative rates of deposition of HF, HCl, HBr, HI and HNO₃ (in particle form) may differ from the
304 relative abundances of the gases, due to increasing partitioning of HF, and to a lesser extent HBr and
305 HI, relative to HCl at low RH. In contrast, at high RH the relative rates of deposition will more

306 closely mimic the relative abundances of the gases. This is exploited by studies using rainwater
307 chemistry as a direct proxy for magmatic gas composition (e.g., Aiuppa et al., 2006).

308 Within the internal mixtures considered in this study, particles are dominated by SO_4^{2-} or HSO_4^-
309 below 80% RH and by Cl^- above 80% RH. HCl partitioning below 80% RH is inhibited due to the
310 acidity of the aerosol. Alternatively, particles in different size fractions may be compositionally
311 distinct (i.e., an external mixture), with only particles in the same size fraction forming an internal
312 mixture (e.g., Mather et al., 2004; Martin et al., 2011). This may enhance acidic gas partitioning as
313 acidic gases interact preferentially with the less acidic particles within the size distribution. While
314 halogen and nitric acids do not react directly with water to form Cl^- -rich aqueous particles (except
315 above 80% RH, although even this may be hindered by the slow rate of homogenous nucleation), Cl^- -
316 rich particles may be stable at any RH by salt formation. Requirements for a “stable” size distribution
317 with enhanced acidic gas partitioning (i.e., greater overall $\text{Cl}^-/\text{SO}_4^{2-}$ than predicted for a fully internal
318 mixture) are therefore a size separation of H^+ and Cl^- , and a size association of Cl^- and metal ions.
319 These conditions appear to be met in the case of Masaya, where particle size measurements indicate a
320 fine mode of $\text{Na}^+\text{-K}^+\text{-H}^+\text{-SO}_4^{2-}$ and a coarse mode of $\text{Mg}^{2+}\text{-Ca}^{2+}\text{-Cl}^-\text{-F}^-$ (e.g., Mather et al., 2003;
321 Martin et al., 2011), with overall $n_{\text{Cl}^-(\text{aq,s})}/n_{\text{SO}_4^{2-}(\text{aq,s})} = 0.2$ (Table 2). In contrast, while particle size
322 measurements at Villarrica volcano (Chile) (Mather et al., 2004) indicate a size association between
323 H^+ , SO_4^{2-} and Cl^- , overall $n_{\text{Cl}^-(\text{aq,s})}/n_{\text{SO}_4^{2-}(\text{aq,s})} (= 0.006)$ is much lower than at Masaya.

324 Our results also have implications for physical and chemical processing in volcanic plumes.
325 Entrainment of air into the plume and dilution results in decreased aerosol volume (assuming that the
326 background aerosol volume is negligible) and decreased gas concentrations, which would promote
327 evaporation of halogen and nitric acids from the particles. Further complexity is introduced by the
328 dependence of partitioning on temperature and relative humidity. These factors may inhibit or
329 promote heterogeneous chemistry under certain conditions. In any case, it is likely that the
330 partitioning of halogen and nitric acids may have a strong spatial and temporal variation. Our results
331 are important because they establish the intrinsic thermodynamic influences on plume evolution.
332 Furthermore, as only a few processes are considered, the results offer an insight into plume evolution

333 in the absence of the more complex chemistry considered within ambient temperature model studies
334 (e.g., Roberts et al., 2009; von Glasow, 2010), or suggested from field measurements (e.g., SO₂
335 oxidation at ambient temperature; Oppenheimer et al., 2010).

336 4 Conclusions

337 The uptake of halogen and nitric acids into acidic sulphate particles is a fundamental process with
338 wide implications in volcanology and the environmental geochemistry of volcanic emissions.

339 Using the thermodynamic Aerosol Inorganics Model (E-AIM; Wexler and Clegg, 2002) for HCl, HBr
340 and HNO₃, combined with a simple three-parameter model for HF and HI (e.g., Mather et al., 2003),
341 we show that the equilibrium partitioning of halogen and nitric acids into acidic sulphate particles is
342 maximised at high relative humidity and low temperature, in near-source (i.e., less dilute) volcanic
343 plumes with high SO₄²⁻/SO₂. The addition of NaCl enhances acidic gas partitioning at high RH, but
344 diminishes acidic gas partitioning at low RH due to a decrease in the water content of the particles and
345 the formation of Na-sulphate salts. We predict that acidic gas partitioning into acidic sulphate
346 particles is quantitatively significant (>1%) only in cool humid conditions (e.g., > 80% RH at 298 K),
347 adding confidence to spectroscopic and/or electrochemical determinations of gas ratios (e.g.,
348 SO₂/HCl, HCl/HF) for volcano monitoring. Over a wider range of atmospheric conditions, acidic gas
349 partitioning remains environmentally important because of significant changes in the amounts of
350 particle forms of acidic gases, e.g., HF(aq), Cl⁻(aq), Br⁻(aq), I⁻(aq) and NO₃⁻(aq). This influence may
351 result in increased rates of acidic gas deposition during the times of increased RH, such as the night-
352 time and during the wet season.

353 Despite the inherent limitations of the equilibrium assumption (i.e., is equilibrium attained over an
354 appropriate timescale?) and parametric modelling (i.e., are variables independent?), our results are
355 important because they establish the intrinsic thermodynamic influences on plume evolution. These
356 influences are complementary to the kinetic influences (i.e., chemical reactivity, dispersion)
357 considered within more complex models of ambient temperature plume evolution.

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- 469

470 **TABLE 1**

471 Henry's law ($K_{H:X}$) and acidity ($K_{A:X}$) constants for the halogen and nitric acids. K_H was calculated
 472 from values of $K_{H:X}K_{A:X}$ (Brimblecombe and Clegg, 1988) and $K_{A:X}$ (Greenwood and Earnshaw,
 473 1997).

Species	$K_H^{(1)}$ (mol dm ⁻³ atm ⁻¹)	$K_A^{(2)}$ (mol dm ⁻³)	$K_{H:X}K_{A:X}^{(1)}$ (mol ² dm ⁻⁶ atm ⁻¹)
HNO ₃	9.6×10^4	2.5×10^1	2.5×10^6
HF	9.6×10^3	1×10^{-3}	9.6×10^0
HCl	2.0×10^{-1}	1×10^7	2.0×10^6
HBr	1.3×10^0	1×10^9	1.3×10^9
HI	2.5×10^{-2}	1×10^{11}	2.5×10^9

474

475 **TABLE 2**

476 Gas and aerosol composition for Masaya volcano (Nicaragua) (Martin et al., 2010). The standard
 477 composition for E-AIM was defined by the values in bold (i.e., HNO₃, HCl, HBr, SO₄²⁻ and Na^{*+})

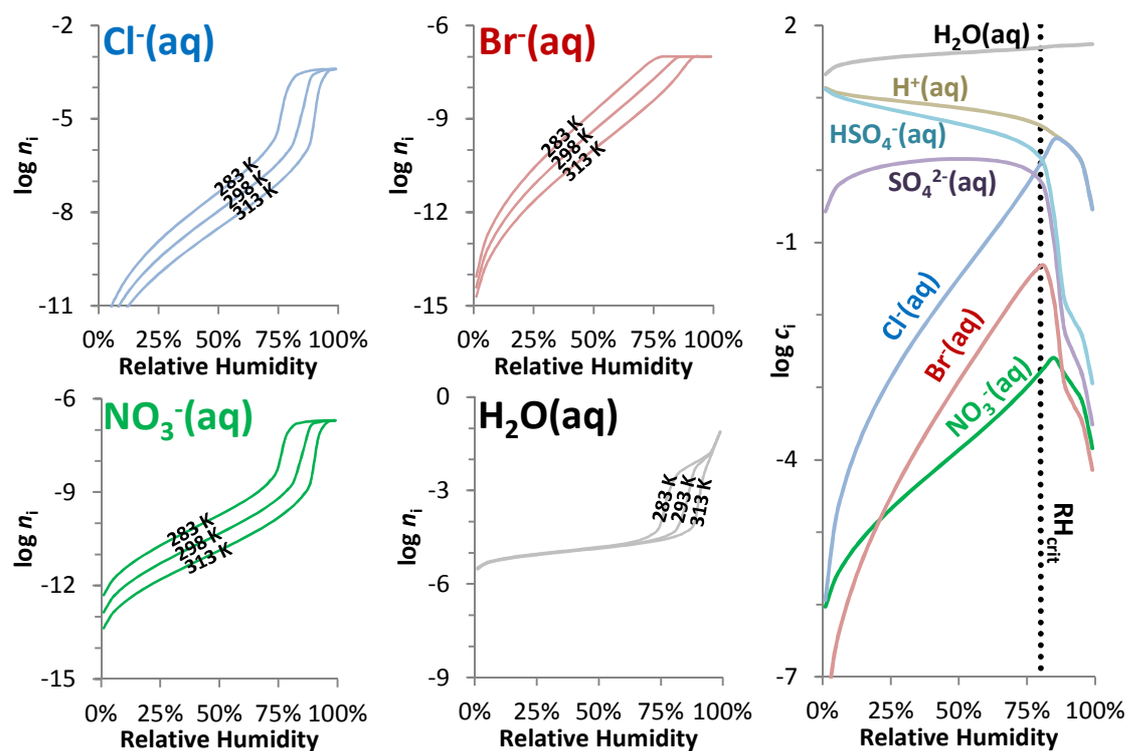
Gas phase	Concentration ($\mu\text{mol m}^{-3}$)	Particle phase	Concentration ($\mu\text{mol m}^{-3}$)
$n_{\text{SO}_2(\text{g})}$	400	$n_{\text{SO}_4^{2-}(\text{aq,s})}$	2
$n_{\text{HNO}_3(\text{g})}$	0.2	$n_{\text{NO}_3^-(\text{aq,s})}$	0.03
$n_{\text{HF}(\text{g})}$	60	$n_{\text{F}^-(\text{aq,s})}$	0.1
$n_{\text{HCl}(\text{g})}$	400	$n_{\text{Cl}^-(\text{aq,s})}$	0.4
$n_{\text{HBr}(\text{g})}$	0.1	$n_{\text{Na}^+(\text{aq,s})}$	2
$n_{\text{HI}(\text{g})}$	0.005	$n_{\text{Na}^{*+}(\text{aq,s})}$	3

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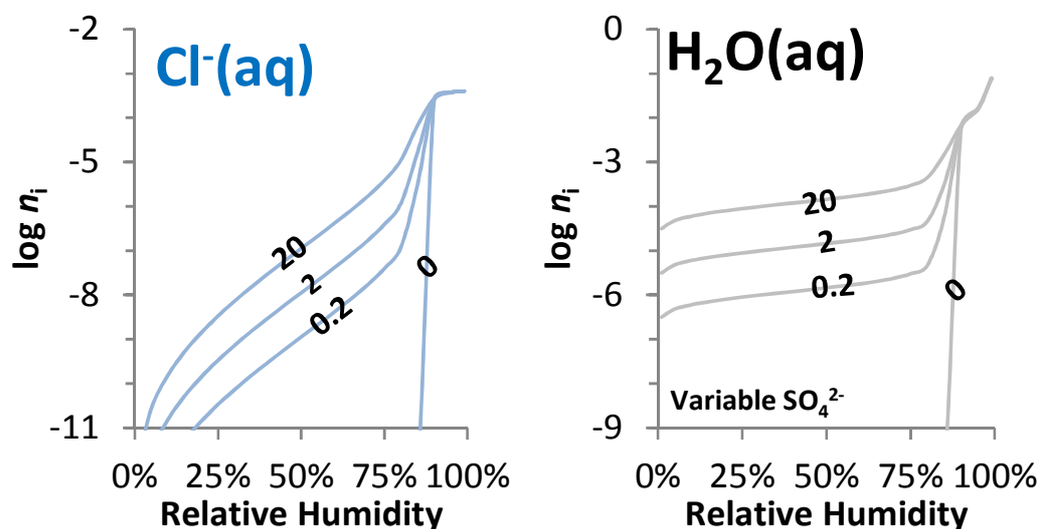
480 **FIGURE 1**

481 E-AIM1 results for the equilibrium amounts of selected aqueous species (n_i ; in mol m⁻³ of air) for the
 482 standard input composition at 283 K, 298 K and 313 K. Also shown are the equilibrium
 483 concentrations (c_i ; in mol dm⁻³ of solution) of all aqueous species considered at 298 K. RH_{crit} is
 484 indicate by a vertical dashed line.

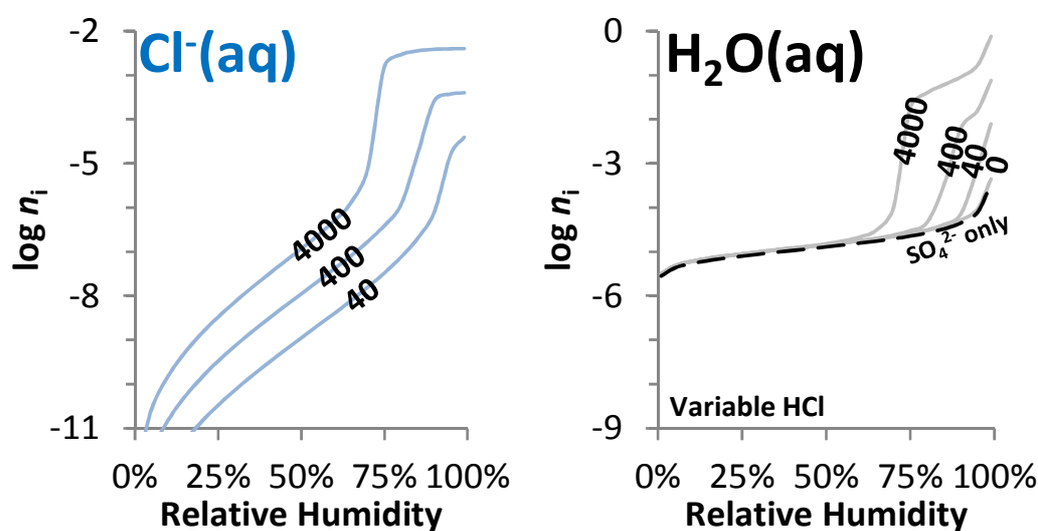


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487 **FIGURE 2A**488 E-AIM1 results for $n_{\text{Cl}^-(\text{aq})}$ and $n_{\text{H}_2\text{O}(\text{aq})}$ (mol m⁻³) for the standard input composition with initial $n_{\text{SO}_4^{2-}}$ 489 $n_{\text{SO}_4^{2-}} = 0, 0.2, 2, 20 \mu\text{mol m}^{-3}$ at 298 K.

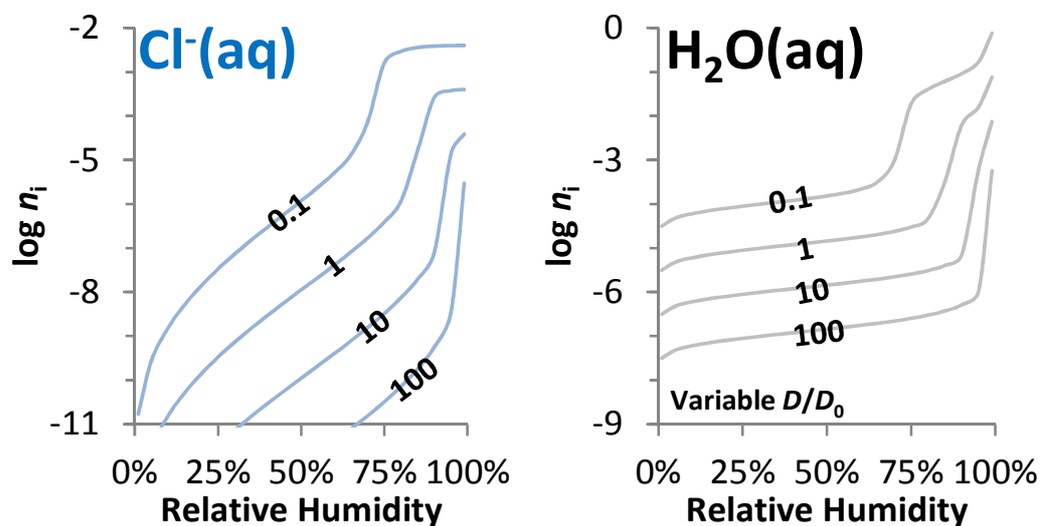
490

491 **FIGURE 2B**492 E-AIM1 results for $n_{\text{Cl}^-(\text{aq})}$ and $n_{\text{H}_2\text{O}(\text{aq})}$ (mol m⁻³) for the standard input composition with initial493 $n_{\text{HCl}(\text{g})} = 0, 40, 400, 4000 \mu\text{mol m}^{-3}$ at 298 K. An additional calculation was made with initial $n_{\text{HCl}(\text{g})} =$ 494 $n_{\text{HBr}(\text{g})} = n_{\text{HNO}_3(\text{g})} = 0 \mu\text{mol m}^{-3}$ at 298 K (i.e., SO₄²⁻ only).

495

496 **FIGURE 2C**

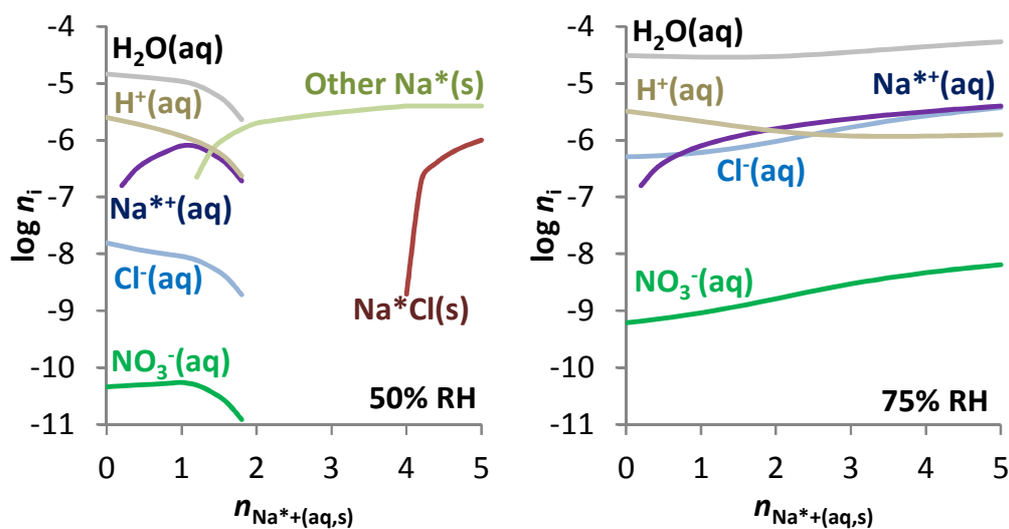
497 E-AIM1 results for $n_{\text{Cl}^-(\text{aq})}$ and $n_{\text{H}_2\text{O}(\text{aq})}$ (mol m⁻³) for the standard input composition with dilutions
 498 (D/D_0) of 0.1, 1 and 10 at 298 K.



499

500 **FIGURE 3**

501 E-AIM3 results for equilibrium amounts various species (n_i ; in mol m⁻³ of air) for the standard input
 502 composition with varying $n_{\text{Na}^+(\text{aq},\text{s})}$ at 50% RH and 75% RH, at 298 K.

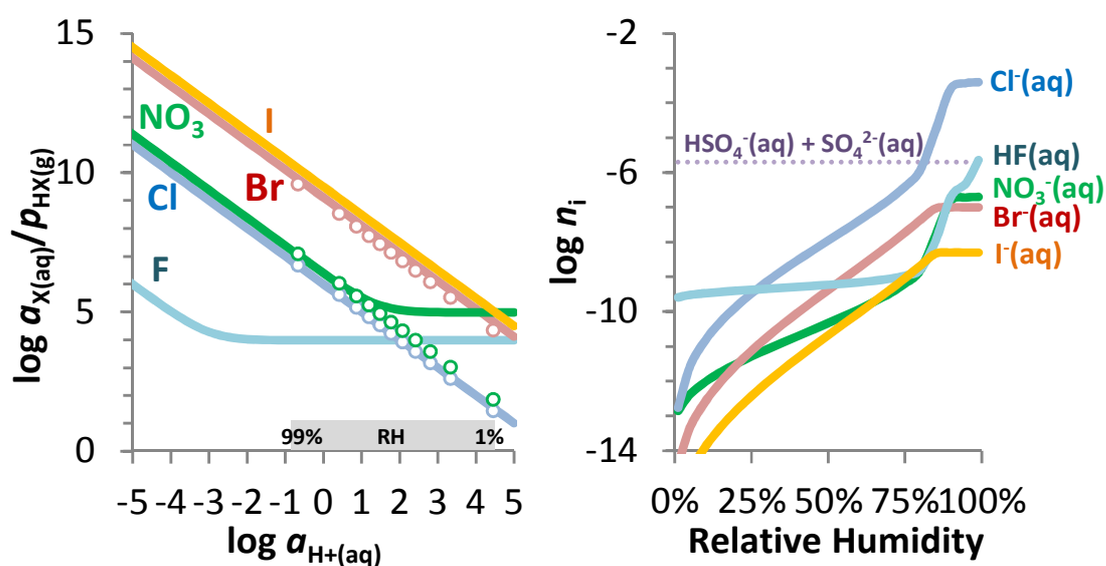


503

504

505 **FIGURE 4**

506 Three-parameter model results for equilibrium $a_{X(aq)}/p_{HX(g)}$ (with $a_{X(aq)}=a_{X-(aq)}+ a_{HX(aq)}$) for the standard
 507 input composition at 298 K, using parameters in Table 1. Equivalent results from E-AIM1 for Cl, Br
 508 and NO_3 are shown with the approximation that $a_{HX(aq)}=0$. Also shown are results for $n_{\text{Cl-(aq)}}$, $n_{\text{Br-(aq)}}$,
 509 $n_{\text{NO}_3\text{-(aq)}}$ (in mol m^{-3} of air) from E-AIM1 for the standard input composition at 298 K, as a function of
 510 the relative humidity. $n_{\text{HF(aq)}}$ and $n_{\text{I-(aq)}}$ are estimated as described in the text.



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