

1 **Measurement Report: Abundance and fractional solubilities of aerosol metals in urban**  
2 **Hong Kong: Insights into factors that control aerosol metal dissolution in an urban site**  
3 **in South China**

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12 **Abstract**

13 Water-soluble metals are known to produce greater adverse human health outcomes than their  
14 water-insoluble forms. Although the concentrations of water-soluble aerosol metals are usually  
15 limited by atmospheric processes that convert water-insoluble metals to water-soluble forms,  
16 factors that control the solubilities of aerosol metals in different environments remain poorly  
17 understood. In this study, we investigated the abundance and fractional solubilities of different  
18 metals in size-fractionated aerosols collected at an urban site in Hong Kong, and identified the  
19 factors that modulated metal solubilities in fine aerosols. The concentrations of total and water-  
20 soluble metals in fine and coarse aerosols were the highest during the winter and spring seasons  
21 due to the long-range transport of air masses by northerly prevailing winds from emission sources  
22 located in continental areas north of Hong Kong. The study-averaged metal fractional  
23 solubilities spanned a wide range for both fine (7.8 % to 71.2 %) and coarse (0.4 % to 47.9 %)  
24 aerosols, but higher fractional solubilities were typically observed for fine aerosols. Sulfate  
25 was found to be strongly associated with both the concentrations of water-soluble Cr, Fe, Co,  
26 Cu, Pb, and Mn and their fractional solubilities in fine aerosols, which implied that sulfate-  
27 driven acid processing likely played an important role in the dissolution of the water-insoluble  
28 forms for these six metals. Further analyses revealed that these strong associations were due to  
29 sulfate providing both the acidic environment and liquid water reaction medium needed for the  
30 acid dissolution process. Thus, the variability in the concentrations of water-soluble Cr, Fe, Co,  
31 Cu, Pb, and Mn and their fractional solubilities were driven by both the aerosol acidity levels  
32 and liquid water concentrations, which in turn were controlled by sulfate. These results  
33 highlight the roles that sulfate plays in the acid dissolution of metals in fine aerosols in Hong  
34 Kong. Our findings will likely also apply to other urban areas in South China, where sulfate is  
35 the dominant acidic and hygroscopic component in fine aerosols.

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## 38 1. Introduction

39 Chronic exposures to atmospheric aerosols, especially those in the fine mode (PM<sub>2.5</sub>,  
40 aerosols with aerodynamic diameter  $\leq 2.5 \mu\text{m}$ ), have been linked to a myriad of deleterious  
41 effects on human health, including morbidity and excessive deaths through respiratory and  
42 cardiovascular diseases (Brook et al., 2010; Cohen et al., 2017). Some of the aerosol chemical  
43 species cause majority of the adverse human health outcomes even though they comprise a  
44 small fraction of the overall aerosol mass (Phalen, 2004; Lippmann, 2014). Metals are  
45 ubiquitous chemical species that contribute significantly to airborne aerosol toxicity even  
46 though they are typically present in aerosols in trace quantities (Costa and Dreher, 1997a;  
47 Frampton et al., 1999; Ye et al., 2018; Zhao et al., 2021). Natural sources, especially mineral  
48 dust and sea spray, dominate the global sources of aerosol metals (Nriagu, 1989; Garrett, 2000;  
49 Deguillaume et al., 2005; Mahowald et al., 2018). However, anthropogenic sources such as  
50 industrial activities and vehicular traffic contribute substantial quantities of aerosol metals in  
51 urban environments (Garg et al., 2000; Adachi and Tainosho, 2004; Deguillaume et al., 2005;  
52 Lough et al., 2005; Birmili et al., 2006; Lee et al., 2007; Cheng et al., 2009; Li et al., 2013;  
53 Jiang et al., 2015; Mahowald et al., 2018).

54 Metals exist in aerosols in water-insoluble and water-soluble forms. Compared to their  
55 water-insoluble forms, water-soluble metals have higher bioavailability, which reportedly  
56 allows them to produce adverse human health outcomes (Costa and Dreher, 1997b; Heal et al.,  
57 2009; Fang et al., 2015; Gao et al., 2020a; He et al., 2021). Some water-soluble transition metal  
58 ions (e.g., Fe(II), Fe(III), Cu(I), Cu(II)) are redox-active species and serve as catalysts in  
59 reaction cycles (e.g., Fenton-like reactions) to enhance the *in vivo* production of reactive  
60 oxygen species (ROS) (e.g.,  $\cdot\text{OH}$ ,  $\text{HO}_2\cdot$ ,  $\text{H}_2\text{O}_2$ ), which subsequently induce the physiological  
61 oxidative stress and inflammation involved in a variety of chronic and acute diseases (Bresgen  
62 and Eckl, 2015; Lakey et al., 2016; Bates et al., 2019). A recent epidemiologic study reported  
63 that water-soluble Fe concentrations in PM<sub>2.5</sub> showed strong correlations with cardiovascular-  
64 related emergency department visits in Atlanta (Ye et al., 2018). Less abundant water-soluble  
65 aerosol metals such as Cr and Pb are also known to exhibit both carcinogenic and  
66 noncarcinogenic risks to adults and children despite their small quantities (He et al., 2021).

67 Water-soluble metals also play important roles in ocean biogeochemistry and  
68 atmospheric processes. Atmospheric aerosol deposition is an important source of bioavailable  
69 dissolved metals in open oceans. The dissolved metals serve as nutrients, and in some cases  
70 toxins, for various aquatic species (De Baar et al., 2005; Boyd et al., 2007; Paytan et al., 2009;  
71 Jordi et al., 2012). Some transition metal ions such as Fe(III) and Mn(II) ions can facilitate the  
72 formation and aging of organic aerosols (Chu et al., 2013; Al-Abadleh, 2015; Slikboer et al.,  
73 2015; Chu et al., 2017; Al-Abadleh, 2021). The coupled redox cycling of Cu(I)/Cu(II) and  
74 Fe(II)/Fe(III) ions in aerosols has been proposed to be an important mechanism for the uptake  
75 of gas-phase HO<sub>2</sub> in aqueous aerosols, which has important implications for the tropospheric  
76 OH radical and O<sub>3</sub> budget (Mao et al., 2013; Mao et al., 2017). Mn(II)-catalyzed oxidation of  
77 SO<sub>2</sub> on aqueous aerosol surfaces reportedly contributes more than 90 % of the sulfate  
78 production during wintertime haze events in China (Wang et al., 2021).

79 Aerosol metals are primarily emitted into the atmosphere in water-insoluble forms  
80 (Nriagu, 1989). While water-soluble aerosol metals can be emitted directly into the atmosphere  
81 (Fang et al., 2015), the concentrations of water-soluble aerosol metals are likely limited by  
82 atmospheric processes that convert the water-insoluble metal forms to water-soluble forms  
83 (Mahowald et al., 2018). Given the important roles that water-soluble aerosol metals play in  
84 adverse human health outcomes and atmospheric processes, it is necessary to understand the  
85 factors that modulate the atmospheric processing, and hence the solubility, of aerosol metals.  
86 Aerosol Fe dissolution has been the focus of most previous studies. A wide range (<1 % to  
87 98 %) of fractional solubilities (ratio of the water-soluble metal mass concentration to the total  
88 metal mass concentration) has been reported for Fe in atmospheric aerosols (Mahowald et al.,  
89 2018). Anthropogenic-influenced aerosols generally have higher Fe solubility than fresh  
90 mineral dust (Sedwick et al., 2007; Schroth et al., 2009; Oakes et al., 2012). However, Fe  
91 solubility varies substantially in aerosols in different urban environments with high levels of  
92 anthropogenic activities (e.g., 1 % to 12 % in four cities in East China (Zhu et al., 2020) vs.  
93 around 20 % to 50 % in Hong Kong, South China (Jiang et al., 2014; Jiang et al., 2015).  
94 Although there are a number of atmospheric processes that can influence aerosol metal  
95 solubilities, acid processing and the formation of stable Fe-organic complexes are two key

96 chemical processes known to enhance aerosol Fe dissolution (Deguillaume et al., 2005; Ingall  
97 et al., 2018; Tao and Murphy, 2019; Giorio et al., 2022). At present, it remains difficult to  
98 explain the variability of aerosol Fe solubility in urban environments since the extent to which  
99 aerosol Fe dissolution is controlled by factors such as aerosol acidity and/or the presence of  
100 organic ligands (e.g., oxalate) in different urban environments is still not well understood. Even  
101 less is known about the factors that control the solubilities of other aerosol metals beyond Fe.

102 Hong Kong is a highly developed, densely populated city in the Guangdong-Hong  
103 Kong-Macau Great Bay Area (GBA) urban agglomeration, which is a large business and  
104 economic hub located in the southern part of China. While there have been some studies on the  
105 fractional solubilities of various aerosol metals in Hong Kong (Jiang et al., 2014; Jiang et al.,  
106 2015), to the best of our knowledge, there has not been a study that has investigated the factors  
107 that control the solubilities of aerosol metals in Hong Kong. In this study, we investigated the  
108 abundance and fractional solubilities of ten metals (Fe, Cu, Al, V, Cr, Mn, Co, Ni, Cd, and Pb)  
109 in aerosols at an urban site in Hong Kong. Our main goal is to identify the key factors that  
110 control the solubilities of metals in fine aerosols since they are believed to exert higher toxicity  
111 than coarse aerosols due to their small sizes. We focus primarily on aerosol metal dissolution  
112 through the acid processing and/or metal-organic complexation mechanisms. Hence, other  
113 aerosol species were also measured for comparisons to total and water-soluble metals. The  
114 measured aerosol inorganic ion composition was used as inputs for a thermodynamic model to  
115 determine the aerosol acidity levels, liquid water concentrations, and pH.

## 116 **2. Methods**

### 117 **2.1. Ambient sampling**

118 The sampling campaign took place at ground level next to a road in Kowloon Tong  
119 (22.3367° N, 114.1724° E). Kowloon Tong is located in the southern side of Hong Kong, and  
120 it is primarily a residential and commercial district which is close to Mongkok, one of the  
121 busiest commercial and most densely populated areas in Hong Kong with high density traffic  
122 flow. Weekly size-fractionated aerosol samples were collected on 7 March 2021 to 4 April 2021  
123 (spring season), 23 to 30 June 2021 and 7 to 14 July 2021 (summer season), 13 September

124 2021 to 11 October 2021 (fall season), and 15 December 2021 to 26 January 2022 (winter  
125 season). Back-trajectories calculations calculated by the Hybrid Split-Particle Lagrangian  
126 Integrated Trajectory (HYSPLIT) model using meteorological data from NCEP/NCAR  
127 Reanalysis (2.5° latitude-longitude grid) showed that the sampling site was under the influence  
128 of continental and marine air masses during the sampling periods, though the contributions of  
129 these air masses varied with the season (Figure S1).

130 An eleven stage Micro-Orifice Uniform Deposit Impactor (MOUDI) (Model 110, MSP  
131 Corp., USA) was used to collect and divide aerosols into different aerosol size bins under  
132 ambient conditions. Aerosols were collected on prebaked 47 mm diameter quartz filters  
133 (Tissuquartz 2500QAT-UP, Pall Corp., USA). The nominal cut points for the MOUDI eleven  
134 impactor stages were 0.056  $\mu\text{m}$ , 0.1  $\mu\text{m}$ , 0.18  $\mu\text{m}$ , 0.32  $\mu\text{m}$ , 0.56  $\mu\text{m}$ , 1.0  $\mu\text{m}$ , 1.8  $\mu\text{m}$ , 3.2  $\mu\text{m}$ ,  
135 5.6  $\mu\text{m}$ , 10  $\mu\text{m}$ , and 18  $\mu\text{m}$ . In the discussion below, for simplicity, we refer to aerosols  
136 collected on impactor stages with nominal cut points 0.056  $\mu\text{m}$ , 0.1  $\mu\text{m}$ , 0.18  $\mu\text{m}$ , 0.32  $\mu\text{m}$ ,  
137 0.56  $\mu\text{m}$ , 1.0  $\mu\text{m}$ , and 1.8  $\mu\text{m}$  as “fine aerosols”, while aerosols collected on impactor stages  
138 with nominal cut points 3.2  $\mu\text{m}$ , 5.6  $\mu\text{m}$ , 10  $\mu\text{m}$ , and 18  $\mu\text{m}$  were referred to as “coarse  
139 aerosols”. Aerosols were collected continuously for seven days (i.e., 24 hours  $\times$  7 days). This  
140 resulted in a total of four, two, four, and six weekly sets of aerosol filter samples collected  
141 during the spring, summer, fall, and winter seasons, respectively. After collection, the aerosol  
142 filter samples were immediately extracted for chemical analysis.

143 Thermodynamic model calculations used to determine the aerosol acidity levels, liquid  
144 water concentrations, and pH (Section 2.3) require gas-phase  $\text{NH}_3$  concentrations, ambient  
145 temperature and relative humidity (RH) as model inputs. Hence, weekly  $\text{NH}_3$  measurements  
146 were performed during each sampling period using four passive sampling devices (PSDs) and  
147 pre-coated collection pads (PS-100 and PS-154, Ogawa & Co., Pompano Beach, FL), except  
148 from 7 to 28 March 2021. The exposed PSD collection pads were extracted in purified  
149 deionized water (18.2 M $\Omega$ -cm) using the protocol recommended by the manufacturer. These  
150 aqueous extracts were subsequently analyzed by ion chromatography (Section 2.2) to  
151 determine the average  $\text{NH}_3$  concentration during the sampling period. A Vantage Vue Weather  
152 Station (Model 6250, Davis Instruments, USA) was used to measure ambient temperature and

153 RH during each sampling period.

## 154 **2.2. Chemical analysis**

155 Each aerosol filter sample was cut into four equal pieces for chemical analysis of  
156 different chemical components. One of the four pieces was extracted in purified deionized  
157 water via sonication (1 hour), followed by high speed vortexing at 3000 rpm (15 minutes). The  
158 resulting aqueous extract was filtered using 0.22  $\mu\text{m}$  pore size nylon filters (Jinteng Instrument  
159 Co., Tianjin, China) before it was analyzed by a Total Organic Carbon (TOC) analyzer (TOC-  
160 VCSH, Shimadzu, Japan) to determine the concentration of water-soluble organic carbon  
161 (WSOC). The TOC analyzer has a limit of detection (LOD) of 0.5  $\text{mg L}^{-1}$ . The second filter  
162 piece was similarly extracted in purified deionized water via sonication (1 hour) and high speed  
163 vortexing at 3000 rpm (15 minutes), and filtered using 0.22  $\mu\text{m}$  pore size nylon filters before it  
164 was analyzed by an ion chromatography (IC) system (Dionex ICS-1100, ThermoFisher  
165 Scientific, USA) using an isocratic method to determine the concentrations of water-soluble  
166 anions ( $\text{NO}_3^-$ ,  $\text{SO}_4^{2-}$ ,  $\text{Cl}^-$ , and  $\text{C}_2\text{O}_4^{2-}$ ) and cations ( $\text{NH}_4^+$ ,  $\text{Na}^+$ ,  $\text{K}^+$ ,  $\text{Ca}^{2+}$ , and  $\text{Mg}^{2+}$ ). Anion  
167 separation was achieved using a 4  $\times$  250 mm anion exchange column (Dionex IonPac AS18,  
168 ThermoFisher Scientific, USA) equipped with a 4  $\times$  50 mm guard column (Dionex IonPac  
169 AG18, ThermoFisher Scientific, USA). Cation separation was achieved using a 4  $\times$  250 mm  
170 cation exchange column (Dionex IonPac CS12A, ThermoFisher Scientific, USA) equipped  
171 with a 4  $\times$  50 mm guard column (Dionex IonPac CG12A, ThermoFisher Scientific, USA). 16  
172 mM potassium hydroxide and 31 mM methanesulfonic acid were used as eluents at a flowrate  
173 of 1.0  $\text{mL min}^{-1}$  for the anion and cation separations, respectively. The cation IC method was  
174 also used to analyze the aqueous extracts from the exposed PSD collection pads to determine  
175 the average  $\text{NH}_3$  concentration during each sampling week. The LODs for the cation IC method  
176 were 0.025  $\text{mg L}^{-1}$  for  $\text{NH}_4^+$ ,  $\text{Na}^+$ , and  $\text{Mg}^{2+}$ , and 0.025  $\text{mg L}^{-1}$  for  $\text{K}^+$  and  $\text{Ca}^{2+}$ . The LODs for  
177 the anion IC method were 0.125  $\text{mg L}^{-1}$  for  $\text{NO}_3^-$ ,  $\text{SO}_4^{2-}$ , and  $\text{C}_2\text{O}_4^{2-}$ , and 0.025  $\text{mg L}^{-1}$  for  $\text{Cl}^-$ .

178 The remaining two filter pieces were used for metal analysis. One filter piece was  
179 extracted with purified deionized water in metal-free centrifuge tubes via sonication (1 hour),  
180 followed by high speed vortexing at 3000 rpm (15 minutes). The resulting aqueous extract was

181 filtered using 0.22  $\mu\text{m}$  pore size nylon filters, and acidified with 2 %  $\text{HNO}_3$  prior to storage at  
182 4  $^\circ\text{C}$  before chemical analysis of water-soluble metals. The filterable metal fraction in the water  
183 extracts, defined in this study as water-soluble metals, will include all dissolved metal forms  
184 and any colloidal particles with diameters smaller than 0.22  $\mu\text{m}$ . This assumes that all colloidal  
185 particles with diameters smaller than 0.22  $\mu\text{m}$  can penetrate through the syringe filter, and that  
186 the syringe filter's retention efficiency of particles with diameters larger than 0.22  $\mu\text{m}$  is 100 %.  
187 The exact sizes and distribution of metal colloidal particles in the filterable metal fraction in  
188 water extracts in this study are not known. However, Yang et al. (2021) recently reported that  
189 around 84% of Fe and Cu colloidal particles that penetrated through 0.45  $\mu\text{m}$  syringe filters  
190 had nominal diameters smaller than 4 nm. The remaining 16% of Fe and Cu colloidal particles  
191 had nominal diameters between 4 nm and 0.45  $\mu\text{m}$ , and they may be in water-insoluble forms  
192 (e.g., Fe and Cu oxides) so they may not be "true" water-soluble species. Hence, analogous to  
193 observations made by Yang et al (2021), it is possible that the filterable metal fractions in water  
194 extracts in this study contain some metals in water-insoluble forms with diameters smaller than  
195 0.22  $\mu\text{m}$  that penetrated through the 0.22  $\mu\text{m}$  pore size filters. The last filter piece was extracted  
196 via acid digestion for chemical analysis of total metals. The acid digestion protocol we  
197 employed was adapted from published protocols (Jiang et al., 2014; Jiang et al., 2015). The  
198 filter piece was extracted in an acid digestion matrix (16 N  $\text{HNO}_3$  and 12 N  $\text{HCl}$  at a 3:1 volume  
199 ratio) placed in a glass microwave vial using a microwave synthesizer (Initiator+, Biotage,  
200 Sweden). The microwave synthesizer's digestion temperature was ramped up to 150  $^\circ\text{C}$ , and  
201 then held for 15 min. This was followed by cooling and ventilation for 30 minutes. An  
202 evaporation and recovery treatment was next performed to remove  $\text{Cl}^-$  from the matrix to  
203 reduce its interference during chemical analysis. The digestion solution was heated to 200  $^\circ\text{C}$   
204 on a hotplate. Once the solution was observed to be almost dry, 16 N  $\text{HNO}_3$  was added to the  
205 solution. When the solution was observed to be almost dry the second time, 2 %  $\text{HNO}_3$  was  
206 added to the solution. The resulting solution was filtered using 0.22  $\mu\text{m}$  pore size nylon filters,  
207 and then stored at 4  $^\circ\text{C}$  before chemical analysis of total metals. A standard reference material  
208 of San Joaquin soil (SRM 2709a, NIST) was digested and analyzed using the same protocols  
209 to evaluate the metal recoveries. Recoveries of 59.4 % for Cr, 67.0 % for Al, 93.7 % for Fe,

210 93.6 % for Ni, 100.2 % for Co, 98.6 % for Pb, 95.8 % for Cu, 99.6 % for Mn, 70.5 % for V,  
211 and 94.3 % for Cd were observed.

212 The concentrations of ten water-soluble and total metals ( $^{27}\text{Al}$ ,  $^{51}\text{V}$ ,  $^{52}\text{Cr}$ ,  $^{55}\text{Mn}$ ,  $^{57}\text{Fe}$ ,  
213  $^{59}\text{Co}$ ,  $^{60}\text{Ni}$ ,  $^{65}\text{Cu}$ ,  $^{111}\text{Cd}$ , and  $^{208}\text{Pb}$ ) were determined by an Inductively Coupled Plasma–Mass  
214 Spectrometry (ICP–MS) instrument (NexION 1000, PerkinElmer Inc., USA). The following  
215 parameters were used for the ICP-MS instrument:  $0.98\text{ L min}^{-1}$  nebulizer gas flow,  $1.2\text{ L min}^{-1}$   
216 auxiliary gas flow,  $15\text{ L min}^{-1}$  plasma gas flow,  $5\text{ mL min}^{-1}$  He gas flow, 1600 W RF power,  
217 35 rpm nebulizer pump rate, and 35 rpm sample pump rate. A multi-elemental calibration  
218 standard (IV-STOCK-13, Inorganic Ventures, USA) was used to quantify the ten water-soluble  
219 and total metals. An internal standard solution of  $^{115}\text{In}$  ( $10\text{ }\mu\text{g L}^{-1}$ ) was added to all samples  
220 and standards to monitor analytical drift. The LODs for  $^{27}\text{Al}$ ,  $^{51}\text{V}$ ,  $^{52}\text{Cr}$ ,  $^{55}\text{Mn}$ ,  $^{57}\text{Fe}$ ,  $^{59}\text{Co}$ ,  $^{60}\text{Ni}$ ,  
221  $^{65}\text{Cu}$ ,  $^{111}\text{Cd}$ , and  $^{208}\text{Pb}$  were 87, 0.8, 2.8, 1.6, 277, 0.7, 4.6, 6.7, 1, and  $0.4\text{ ng L}^{-1}$ , respectively.  
222 To identify the major sources of the aerosol metals, source apportionment was performed with  
223 positive matrix factorization (PMF) (Paatero and Tapper, 1994; Paatero, 1997) using the  
224 aerosol chemical components measured by the ICP-MS and IC. Details of the PMF method  
225 used can be found in Section S1 (SI).

### 226 **2.3. Thermodynamic modeling**

227 The thermodynamic model ISORROPIA-II was used to determine aerosol acidity levels,  
228 liquid water concentrations, and pH (Fountoukis and Nenes, 2007). Similar to the methodology  
229 employed by Fang et al. (2017), we ran ISORROPIA-II for each of the MOUDI impactor stages  
230 that collected fine aerosols. The measured water-soluble  $\text{NH}_4^+$ ,  $\text{SO}_4^{2-}$ ,  $\text{NO}_3^-$ ,  $\text{Cl}^-$ ,  $\text{Na}^+$ ,  $\text{Ca}^{2+}$ ,  $\text{K}^+$ ,  
231 and  $\text{Mg}^{2+}$  ions for the aerosols collected on the MOUDI impactor stage, gas-phase  $\text{NH}_3$ ,  
232 ambient temperature and RH were used as model inputs. Since gas-phase  $\text{NH}_3$  measurements  
233 were not available from 7 to 28 March 2021, we used  $\text{NH}_3$  measurements from 28 March to 4  
234 April 2021 as model inputs for the spring calculations. The measured  $\text{NH}_3$  concentrations  
235 during the study ranged from  $3.60\text{ }\mu\text{g m}^{-3}$  to  $8.18\text{ }\mu\text{g m}^{-3}$ , with a study-averaged concentration  
236 of  $5.01 \pm 1.25\text{ }\mu\text{g m}^{-3}$ . ISORROPIA-II was run in “forward” mode and under the assumption  
237 that the aerosols existed in a “metastable” equilibrium state (i.e., the aerosols only existed in



238 liquid form). These calculations assumed that the aerosols were in thermodynamic equilibrium  
239 with the gas phase. While fine aerosols satisfy this equilibrium condition, equilibrium between  
240 the gas and aerosol phases of coarse aerosols cannot be achieved due to kinetic limitations  
241 (Fountoukis et al., 2009). Thus, aerosol pH values were not calculated for coarse aerosols.

242 Fine aerosol pH values were calculated based on the molal definition (Pye et al., 2020):

$$243 \quad pH = -\log_{10} \gamma_{H^+} H_{aq}^+ = -\log_{10} \frac{1000 H_{air}^+}{W_i + W_o} \cong -\log_{10} \frac{1000 H_{air}^+}{W_i} \quad (1)$$

244 where  $\gamma_{H^+}$  is the hydronium ion activity coefficient,  $H_{aq}^+$  is the hydronium ion concentration  
245 within the ambient aerosol liquid water ( $\text{mol L}^{-1}$ ),  $H_{air}^+$  is the hydronium ion concentration per  
246 volume of air ( $\mu\text{g m}^{-3}$ ), and  $W_i$  and  $W_o$  are the aerosol liquid water concentrations ( $\mu\text{g m}^{-3}$ )  
247 associated with inorganic and organic species, respectively.  $H_{air}^+$  and  $W_i$  are the outputs  
248 provided by the ISORROPIA-II model, which assumes that  $\gamma_{H^+}$  is equals to unity.  $W_o$  can be  
249 estimated from the WSOC measurements using the approach described in Section S2 (SI).  
250 WSOC concentrations in the size-fractionated aerosols ranged from 0 to  $4.6 \mu\text{g m}^{-3}$ . The  
251 inclusion of  $W_o$  into calculations did not impact aerosol pH significantly (Figure S2). Thus,  
252 only aerosol pH values calculated using  $W_i$  will be reported here. Similar to Fang et al. (2017),  
253 lower pH values were typically calculated for aerosols collected on MOUDI impactor stages  
254 with smaller nominal cut points (i.e., these aerosols had smaller aerodynamic aerosol diameters)  
255 due to the higher mass concentrations of sulfate in these smaller aerosols. The fine aerosols  
256 were mostly acidic, with about 74 % of the calculated pH values lying between 2 and 4.

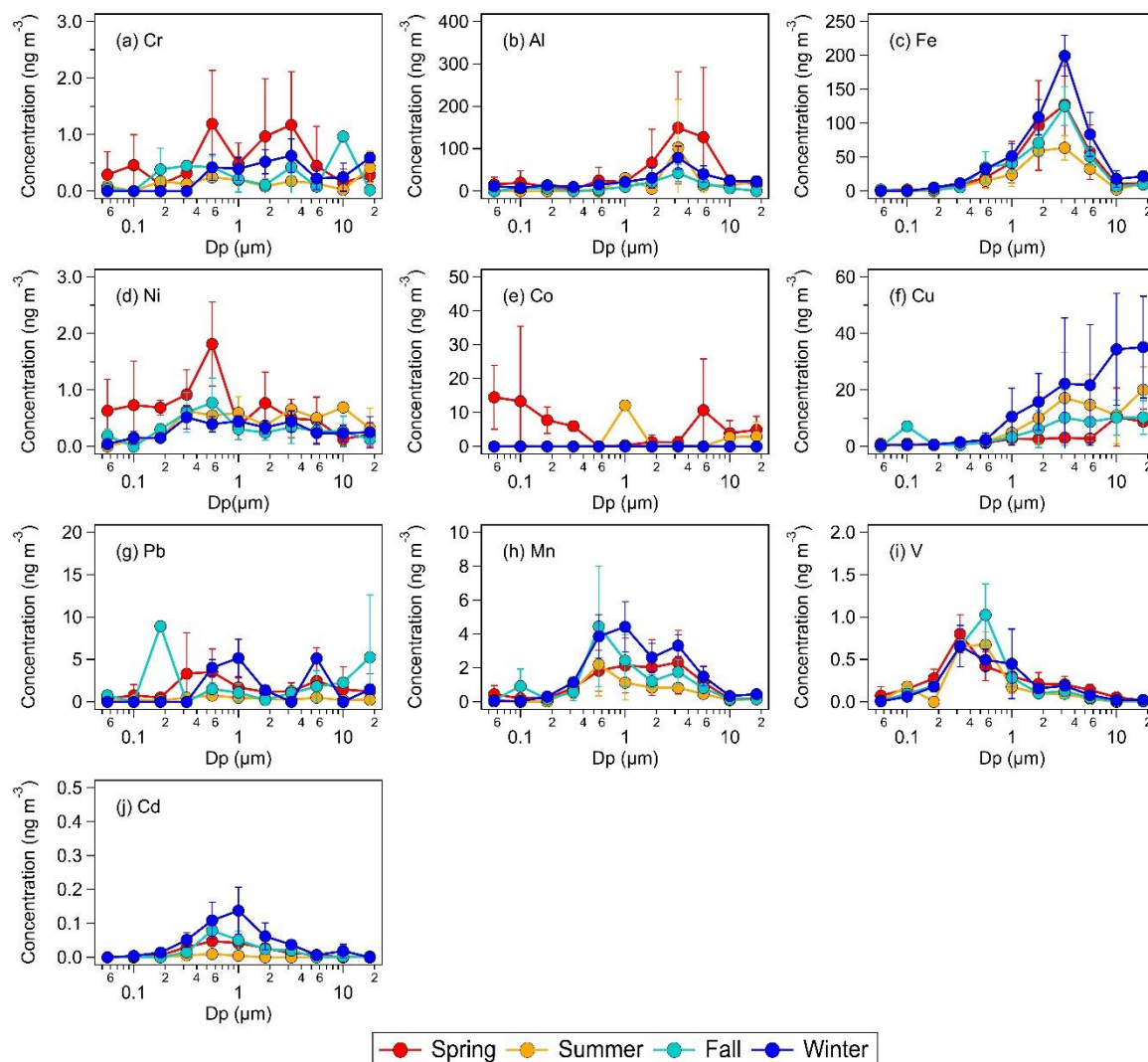
### 257 **3. Results and discussion**

#### 258 **3.1. Total metals**

259 Figure 1 shows the seasonal average mass concentrations of the ten measured total  
260 metals in size-fractionated aerosols. The size distributions of five of the metals (Al, Fe, Mn, V,  
261 and Cd) consistently exhibited a single mode. The modes for Mn, V, and Cd were  
262 predominantly found in the fine mode, while the modes for Fe and Al were predominantly  
263 found in the coarse aerosol mode. Figure 2a shows the seasonal average concentrations of the  
264 ten measured total metals in fine and coarse aerosols. For most of the metals, higher mass

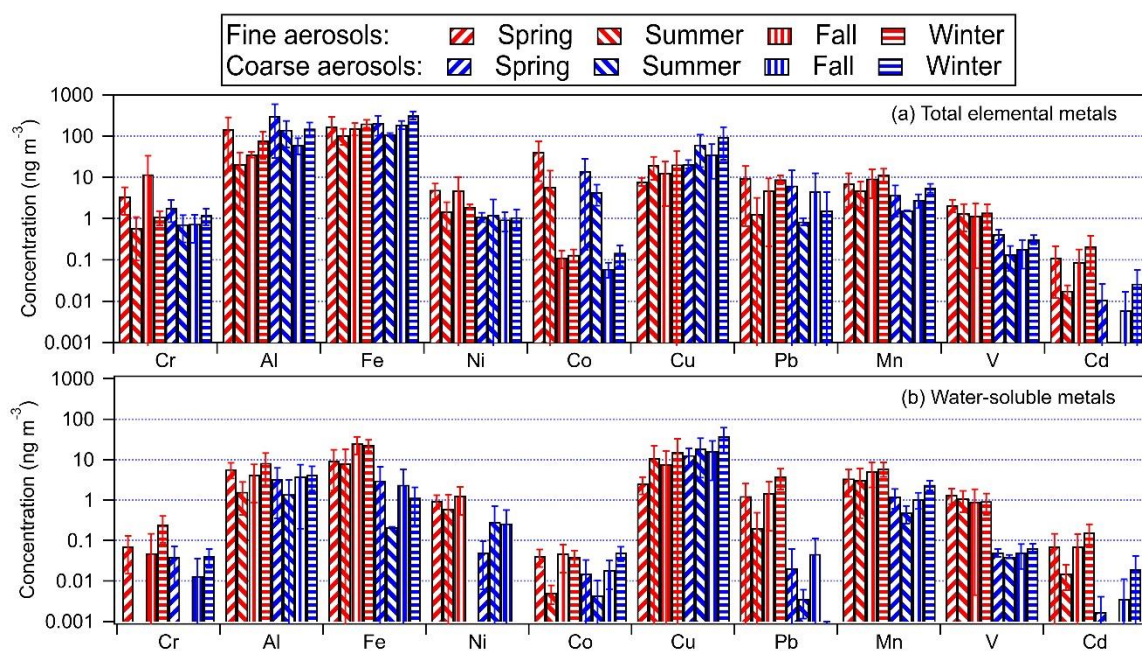
265 concentrations were measured during the winter and/or spring seasons. This could be attributed  
266 to the long-range transport of polluted air masses by northerly prevailing winds from emission  
267 sources located in continental areas north of Hong Kong (Figure S1). The metals could be  
268 arranged in the following order based on their abundances: Fe > Al > Cu > Co > Mn > Pb >  
269 Cr > Ni > V > Cd. This order of abundance was the same for both fine and coarse aerosols.

270 The mass concentrations of the two most abundant metals, Fe and Al, were usually  
271 higher than 10 ng m<sup>-3</sup> in both fine and coarse aerosols. Fe, Al, and Cu had substantially higher  
272 mass concentrations in coarse aerosols than in fine aerosols. The positive correlations between  
273 the mass concentrations of Al with the mass concentrations of Fe and Cu were the strongest  
274 among the nine metals ( $R = 0.62$  and  $R = 0.52$ , respectively) and statistically significant (Figure  
275 S3), which could be explained by large mass concentrations of Al, Fe, and Cu originating from  
276 similar sources. These three metals are known to originate mainly from dust sources (e.g.,  
277 mineral dust and road dust) (Hopke et al., 1980; Garg et al., 2000; Adachi and Tainosho, 2004;  
278 Lough et al., 2005; Chow et al., 2022). This is consistent with results from our PMF source  
279 apportionment analysis, which showed that the “dust” factor had large mass contributions from  
280 Al, Fe, and Cu (Figures S4 and S5). Mn, Ni, V, and Cd had higher mass concentrations in fine  
281 aerosols than in coarse aerosols. These four metals are known to be consistently found in  
282 aerosols from anthropogenic sources such as vehicle and ship emissions, combustion and  
283 industrial processes (Chow et al., 2022). Pb, Cr, and Co had mostly similar concentrations in  
284 the fine and coarse aerosols. Interestingly, the mass concentrations of Mn and Cr were  
285 positively correlated with the mass concentration of Al ( $R = 0.42$  and  $R = 0.33$ , respectively),  
286 and these correlations were statistically significant (Figure S3). Our PMF analysis apportioned  
287 Al to two factors, “dust” and “industrial factor 1”, though the Al contribution to “industrial  
288 factor 1” was substantially smaller compared to “dust” (Figures S4 and S5). The “dust” factor  
289 had a significant Mn contribution, which could explain the strong correlation between the mass  
290 concentrations of Al and Mn. Cr was apportioned primarily to “industrial factor 1”, which  
291 could explain the strong correlation between the mass concentrations of Al and Cr. The mass  
292 concentrations of Ni, V, Cd, Pb, and Co showed weak correlations with the mass concentration  
293 of Al (Figure S3).



294

295 **Figure 1:** Seasonal average concentrations of total elemental metals in size-fractionated  
 296 aerosols sampled by the MOUDI with the following nominal cut points (Dp): 0.056 μm (size  
 297 fraction 1), 0.1 μm (size fraction 2), 0.18 μm (size fraction 3), 0.32 μm (size fraction 4), 0.56  
 298 μm (size fraction 5), 1.0 μm (size fraction 6), 1.8 μm (size fraction 7), 3.2 μm (size fraction 8),  
 299 5.6 μm (size fraction 9), 10 μm (size fraction 10), and 18 μm (size fraction 11). The error bars  
 300 represent one standard deviation of the seasonal average value.



301  
 302 **Figure 2:** Seasonal average mass concentrations of (a) total metals and (b) water-soluble metals  
 303 in fine (red) and coarse (blue) aerosols. The error bars represent one standard deviation. The y  
 304 axes are on logarithm scales.

305 Jiang et al. (2015) previously measured the mass concentrations of various total metals  
 306 in PM<sub>2.5</sub> and PM<sub>2.5-10</sub> in Kowloon Tong. The authors carried out their measurements from 12  
 307 November 2012 to 10 December 2012 (winter) and from 8 April 2013 to 13 May 2013  
 308 (spring/summer). To gain some insights into how the aerosol metal concentrations at this urban  
 309 site have changed since 2012/2013, we compared the average mass concentrations of total  
 310 metals in fine and coarse aerosols measured in this study to those measured by Jiang et al.  
 311 (2015). As shown in Table S1, lower mass concentrations were measured in fine (21 % to 93 %  
 312 lower) and coarse (0.5 % to 92 % lower) aerosols for most of the metals in this study. While  
 313 the lower aerosol metal mass concentrations could be partly attributed to lower levels of  
 314 anthropogenic activities in 2021/2022 due to COVID-19, it is likely that the implementation of  
 315 numerous local and regional air pollution policies to reduce industrial and transport-related  
 316 emissions over the last decade contributed largely to this decrease. For instance, industrial  
 317 upgrades resulting from the implementation of the “double transfer” policy (industry and labor  
 318 transfer away from primary industries) in Guangdong likely caused the decline in the mass  
 319 concentrations of metals that are typically associated with industrial activities such as Cu and

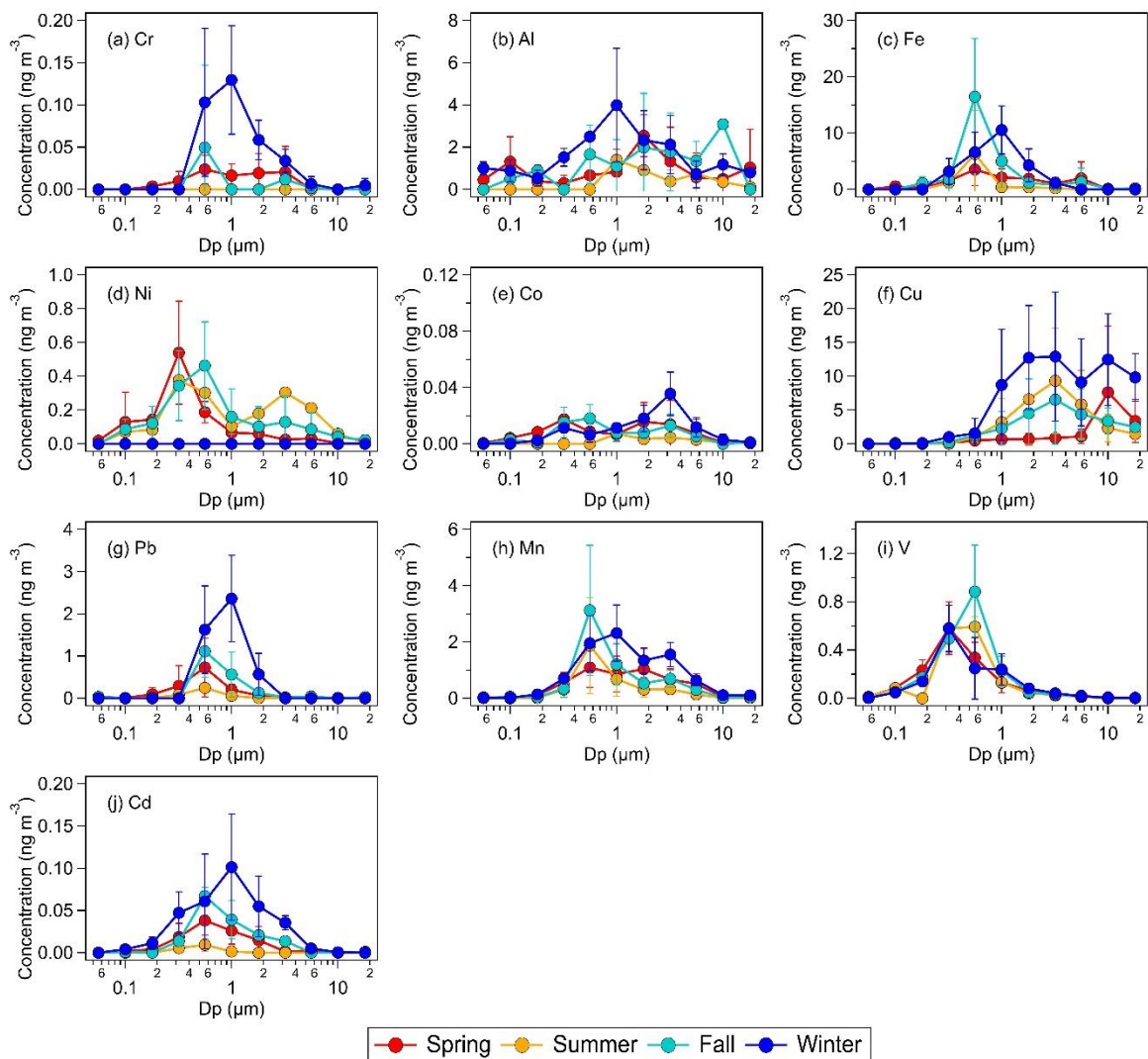
320 Mn (Zhong et al., 2013; Chow et al., 2022). In addition, government policies driving the switch  
321 to cleaner fuels for energy generation and transport in Hong Kong and the GBA likely caused  
322 the decline in the mass concentrations of metals such as Pb, Ni, V, and Fe. Interestingly, higher  
323 mass concentrations were measured for Fe and Cu in coarse aerosols in this study compared to  
324 those measured by Jiang et al. (2015). Fe and Cu in coarse aerosols have previously been linked  
325 to resuspended road dust from brake and tire wear (Garg et al., 2000; Adachi and Tainosho,  
326 2004; Lough et al., 2005). Based on publicly available government data (www.td.gov.hk), the  
327 number of registered motor vehicles in Hong Kong has increased by about 34 % over the last  
328 decade. It is possible that the higher Fe and Cu mass concentrations in coarse aerosols in this  
329 study were due to increased contributions from road dust as a result of increased vehicle fleet  
330 size at the urban site.

331 A PMF source apportionment analysis was performed to determine the major sources  
332 of aerosol metals measured in this study (Section S1). A five-factor solution was selected since  
333 it gave the most reasonable factor profiles and had high stability. The five factors were broadly  
334 classified as “sea salt”, “dust”, “ship emissions”, “industrial factor 1”, and “industrial factor 2”  
335 based on the tracer species with the highest mass loadings in each factor (Figure S4). A  
336 discussion on how these five factors were classified can be found in Section S1. Figure S5  
337 shows the seasonal mass contributions of each source to each metal species in coarse and fine  
338 aerosols. Metals with large fractions in the dust and sea salt source factor profiles generally had  
339 higher mass concentrations in coarse aerosols. Conversely, metals with large fractions in the  
340 ship emissions and industrial source factor profiles generally had higher mass concentrations  
341 in fine aerosols. Higher mass contributions were usually observed in the winter and/or spring  
342 seasons, which could be attributed to the long-range transport of polluted air masses by northerly  
343 prevailing winds from emission sources located in continental areas north of Hong Kong  
344 (Figure S1).

### 345 **3.2. Water-soluble metals**

346 Figure 3 shows the seasonal average mass concentrations of water-soluble metals in  
347 size-fractionated aerosols. The size distribution of six of the water-soluble metals (Cr, Fe, Pb,

348 Mn, V, and Cd) mostly exhibited a single mode, all of which were found in the fine aerosol  
 349 mode. Fe, Mn, V, and Cd exhibited a single mode for both their total and water-soluble  
 350 components (Figures 1 and 3). Of these four metals, only the modes of total and water-soluble  
 351 Fe showed obvious differences, with total Fe exhibiting a mode at around 3.2  $\mu\text{m}$  (size fraction  
 352 8) and water-soluble Fe exhibiting a mode at around 0.56  $\mu\text{m}$  to 1.0  $\mu\text{m}$  (size fractions 5 to 6).  
 353 The modes of total and water-soluble Cu also showed obvious differences. While the mode of  
 354 total Cu was at  $\geq 18 \mu\text{m}$  (Figure 1f), the modes of water-soluble Cu were found at substantially  
 355 small aerosol sizes (Figure 3f).



356

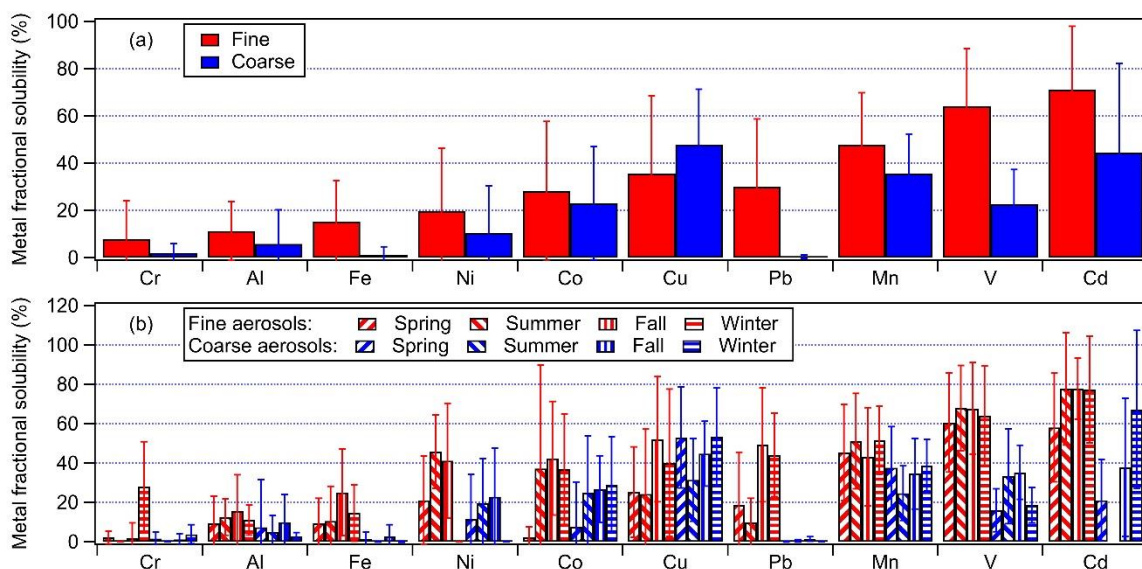
357 **Figure 3:** Seasonal average concentrations of water-soluble metals in size-fractionated aerosols  
 358 sampled by the MOUDI with the following nominal cut points (Dp): 0.056  $\mu\text{m}$  (size fraction  
 359 1), 0.1  $\mu\text{m}$  (size fraction 2), 0.18  $\mu\text{m}$  (size fraction 3), 0.32  $\mu\text{m}$  (size fraction 4), 0.56  $\mu\text{m}$  (size

360 fraction 5), 1.0  $\mu\text{m}$  (size fraction 6), 1.8  $\mu\text{m}$  (size fraction 7), 3.2  $\mu\text{m}$  (size fraction 8), 5.6  $\mu\text{m}$   
361 (size fraction 9), 10  $\mu\text{m}$  (size fraction 10), and 18  $\mu\text{m}$  (size fraction 11). The error bars represent  
362 one standard deviation of the seasonal average value.

363 Figure 2b shows the seasonal average mass concentrations of water-soluble metals in  
364 fine and coarse aerosols. Similar to the total metals, higher mass concentrations of water-  
365 soluble metals were usually measured during the winter and/or spring seasons. With the  
366 exception of Cu, the water-soluble metals usually had higher mass concentrations in fine  
367 aerosols than in coarse aerosols. The water-soluble metals generally had the same order of  
368 abundance as the total metals with some slight variations. The mass concentrations of water-  
369 soluble metals generally correlated with the mass concentrations of total metals (Table S2).  
370 This indicated that the water-soluble metals were largely derived from their total metals  
371 through atmospheric processing, and/or that water-soluble and water-insoluble metals have the  
372 same emission sources. For most of the metals, correlations between the mass concentrations  
373 of water-soluble and total metals were higher for fine aerosols than for coarse aerosols. This  
374 could be due to enhanced metal dissolution in fine aerosols via acid processing and/or the  
375 formation of stable metal-organic complexes, which are two atmospheric chemical processes  
376 that play key roles in influencing the solubilities of aerosol metals in many locations. This is  
377 because acidic inorganic species that promote acid processing and organic species that can  
378 serve as organic ligands are typically present in larger quantities in fine aerosols than in coarse  
379 aerosols. It is also possible that differences in metal mineralogy and atmospheric processing  
380 mechanisms in fine vs. coarse aerosols could have contributed to differences in the metal  
381 dissolution rates (Oakes et al., 2012; Longo et al., 2016; Ingall et al., 2018).

382 Figure 4 shows the study-averaged fractional solubilities for the ten metals in fine and  
383 coarse aerosols. The study-averaged metal fractional solubilities spanned a wide range for both  
384 fine (7.8 % to 71.2 %) and coarse (0.4 % to 47.9 %) aerosols. With the exception of Cu, the  
385 metals generally exhibited higher fractional solubilities in fine aerosols compared to coarse  
386 aerosols. The aerosol size-dependent metal fractional solubility could be explained by  
387 differences in the aerosol composition and metal mineralogy, which resulted in different metal  
388 dissolution rates and/or mechanisms for aerosols of different sizes. Our observations of mostly

389 higher metal fractional solubilities in fine aerosols are consistent with previous studies  
 390 conducted in Hong Kong and other locations worldwide (Baker et al., 2006; Jiang et al., 2014;  
 391 Jiang et al., 2015; Fang et al., 2017; Gao et al., 2019; Baker et al., 2020; Gao et al., 2020b;  
 392 Zhang et al., 2022). No season-dependent trend was observed for the metal fractional  
 393 solubilities.



394  
 395 **Figure 4:** (a) Study-averaged fractional solubilities of metals in fine and coarse aerosols. (b)  
 396 Seasonal average fractional solubilities of metals in fine and coarse aerosols. The error bars  
 397 represent one standard deviation.

398 Some studies have reported that aerosol metal fractional solubilities will exhibit inverse  
 399 relationships with the total metal concentrations as a result of atmospheric processing (Baker  
 400 and Jickells, 2006; Sholkovitz et al., 2012; Mahowald et al., 2018; Shelley et al., 2018; Zhang  
 401 et al., 2022). There was significant scatter in many of our datasets (Figure S6), which made it  
 402 difficult to discern some of the relationships between the metal fractional solubilities and total  
 403 metal concentrations. Inverse relationships between the fractional solubility and total metal  
 404 concentration were noticeable for Cr, Al, Fe, Ni, Cu, Pb, and Mn. However, inverse  
 405 relationships between the Co, V, and Cd fractional solubilities and their total metal  
 406 concentrations were less noticeable due to their low concentrations and scatter in their datasets.  
 407 A number of factors could have contributed to the scatter in the datasets. For instance, the  
 408 scatter could be a result of the total and water-soluble metal concentrations being substantially



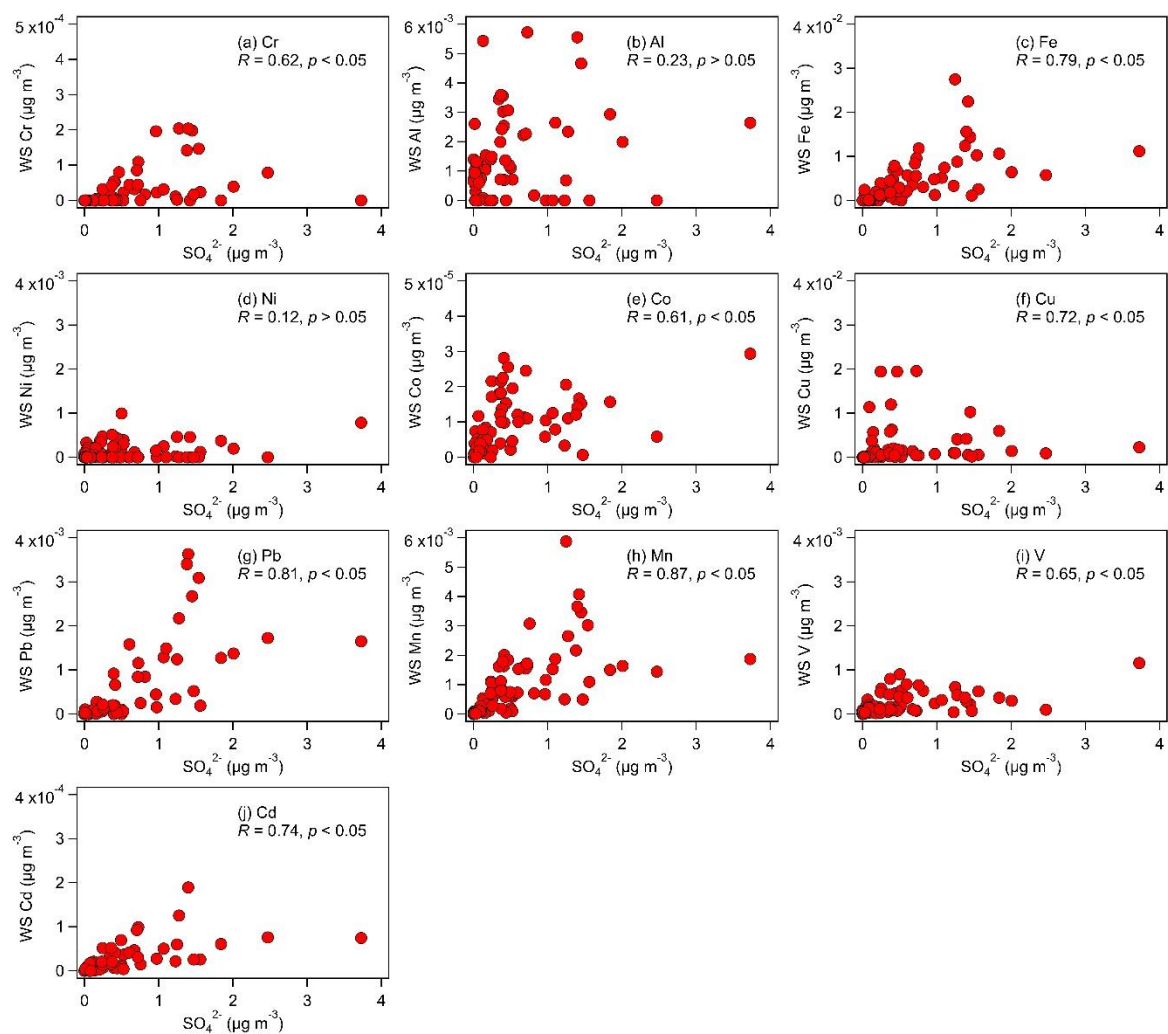
409 different in individual aerosol particles, which would not be captured by the bulk chemical  
410 analysis performed in this study (Oakes et al., 2012; Longo et al., 2016; Ingall et al., 2018).  
411 The metal dissolution rates in individual aerosol particles could also be significantly different  
412 due to differences in metal mineralogy, aerosol acidity levels, presence of organic ligands etc.  
413 in individual aerosol particles.

### 414 **3.3. Factors that control the aerosol metal solubilities**

415 Here, we identify the factors that control metal solubilities in fine aerosols since they  
416 are believed to exert higher toxicity than coarse aerosols due to their small sizes. Our analyses  
417 focus on aerosol metal dissolution via metal-organic complexation reactions and acid  
418 processing, which are two atmospheric chemical processes believed to drive aerosol metal  
419 dissolution in most environments. Laboratory studies have shown that the presence of organic  
420 ligands enhances Fe dissolution in aerosols (Paris et al., 2011; Chen and Grassian, 2013; Paris  
421 and Desboeufs, 2013; Wang et al., 2017). Water-soluble dicarboxylic acids, especially oxalate,  
422 form stable complexes with Fe ions, which will lower the energy barrier for Fe dissolution.  
423 While evidence of organic ligand-promoted metal dissolution in ambient aerosols has been less  
424 conclusive, recent field studies compared the oxalate and water-soluble Fe concentrations to  
425 show that the presence of organic ligands could contribute to aerosol Fe solubility. For instance,  
426 strong positive correlations between oxalate and water-soluble Fe mass concentrations were  
427 observed for PM<sub>2.5</sub> collected at six urban and rural sites in Canada (Tao and Murphy, 2019).  
428 The Fe fractional solubility was also observed to be positively correlated with the molar ratio  
429 of oxalate and Fe for PM<sub>2.5</sub> collected at a suburban site in Qingdao, China (Zhang et al., 2022).

430 To investigate whether organic ligands influenced aerosol metal solubilities in this study,  
431 we attempted to measure oxalate in the size-fractionated aerosol samples using IC. However,  
432 we could not detect oxalate, which indicated that the concentrations of oxalate (if present) were  
433 below the detection limits of our IC instrument. It should be noted that although a recent study  
434 reported the copresence of Fe and oxalate in individual aerosol particles at a suburban site in  
435 Hong Kong using single particle mass spectrometry (Zhou et al., 2020), organic ligand-  
436 promoted metal dissolution is a slow process, and it plays a minor role in metal dissolution

437 under low pH conditions (Zhu et al., 1993). The fine aerosols collected in this study were  
 438 mostly acidic, with about 60 % of the calculated pH values being less than 3. This suggested  
 439 that organic ligand-promoted dissolution may have played a minor role in enhancing aerosol  
 440 metal solubilities in this study due to the acidic nature of the aerosols.



441  
 442 **Figure 5:** Relationships between the mass concentrations of water-soluble (WS) metals and  
 443 sulfate in fine aerosols. Only data with non-zero total metal concentrations were used in the  
 444 figures. Also shown are the spearman correlation coefficients for each relationship.

445 The acidic nature of the aerosols raises the possibility that acid processing played a  
 446 major role in enhancing aerosol metal solubilities. A previous study that utilized nanoscale  
 447 single-particle mass spectrometric and imaging techniques to analyze the mixing states of Fe-  
 448 containing aerosols collected over the East China Sea provided insights into the mechanism of

449 Fe dissolution by acidic species that condensed onto atmospheric aerosols (Li et al., 2017). The  
450 authors reported that Fe oxide-rich aerosols emitted from steel plants and coal combustion were  
451 coated with thick layers of acidic sulfate after 1 to 2 days of atmospheric aging. These sulfate  
452 coatings originated from the condensation of sulfuric acid, which were formed from reactions  
453 of anthropogenic SO<sub>2</sub>. While the fresh aerosols were composed primarily of insoluble Fe oxide,  
454 the aged aerosols contained soluble Fe sulfate that were internally mixed in the sulfate coatings.  
455 Although the mechanism proposed by Li et al. (2017) focused on explaining how sulfate-driven  
456 acid processing leads to the dissolution of the water-insoluble forms Fe, this mechanism likely  
457 applies to the other aerosol metals as well. During acid processing, acidic species have to  
458 overcome the buffering capacity of the aqueous aerosol particle to raise the aerosol acidity level  
459 to the point where the dissolution of metal species is thermodynamically favored. Sulfate was  
460 the most abundant aqueous-phase acidic species in our size-fractionated aerosol samples. The  
461 concentrations of nitrate (another aqueous-phase acidic species) were very low (about 18 times  
462 lower than sulfate, on average), while aqueous-phase organic acids were not detected. Hence,  
463 we first analyzed the relationships between the concentrations of water-soluble metals and  
464 sulfate. Figure 5 shows that despite the scatter in the datasets, the concentrations of water-  
465 soluble metals were positively correlated with the concentration of sulfate, though the  
466 correlations between the concentrations of sulfate and water-soluble Al and Ni were not  
467 statistically significant. These positive correlations could be due, in part, to the water-soluble  
468 metals and sulfate precursor (i.e., SO<sub>2</sub>) being emitted from the same sources. However, the  
469 masses of primary water-soluble aerosol metals are not known. The positive correlations could  
470 also be due to the role that sulfate plays in aerosol metal dissolution during acid processing.

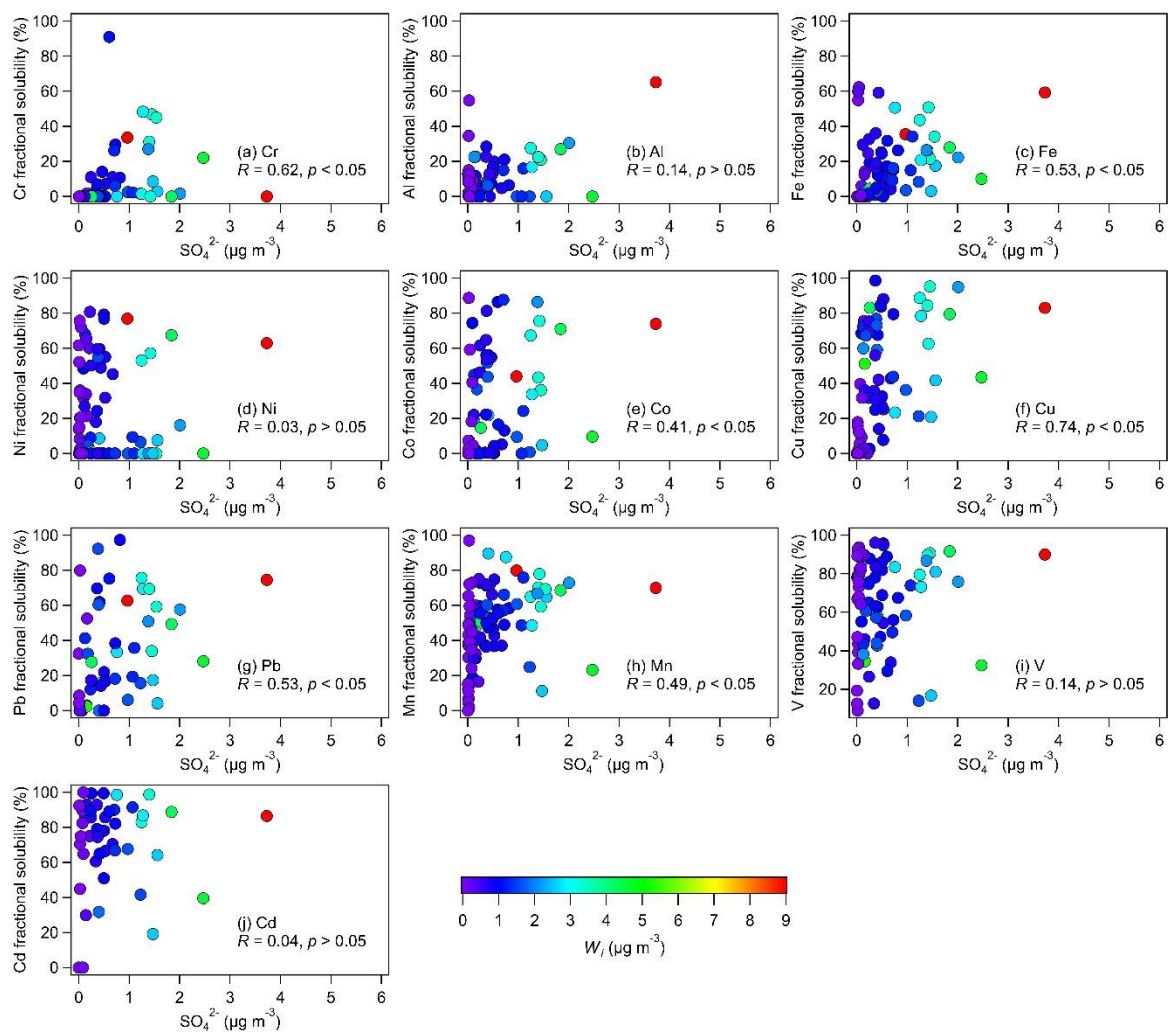
471 To investigate the roles that sulfate and nitrate played in controlling aerosol metal  
472 solubilities, we analyzed the relationships between the metal fractional solubilities and sulfate  
473 and nitrate concentrations. In general, the correlations between the metal fractional solubilities  
474 and sulfate concentration (Table 1) were substantially higher than the correlations between the  
475 metal fractional solubilities and nitrate concentration (Table S3). This implied that sulfate likely  
476 plays a more important role than nitrate in controlling aerosol metal solubilities, which is not  
477 surprising given the low concentrations of nitrate detected (about 18 times lower than sulfate,

478 on average). Analyses of the correlations between the metal fractional solubilities and sulfate  
479 concentration (Table 1 and Figure 6) indicated that the Cr, Fe, Co, Cu, Pb, and Mn fractional  
480 solubilities were positively correlated with the sulfate concentration, and these correlations  
481 were statistically significant. This implied that sulfate played a key role in the formation of  
482 water-soluble Cr, Fe, Co, Cu, Pb, and Mn, likely through sulfate-driven acid dissolution of their  
483 water-insoluble forms. Conversely, the positive correlations between the sulfate concentration  
484 and the Al, Ni, V, and Cd fractional solubilities were weak and not statistically significant.  
485 Interestingly, the V and Cd fractional solubilities showed weak correlations with the sulfate  
486 concentration ( $R = 0.14$  and  $R = 0.04$ , respectively), whereas their water-soluble concentrations  
487 showed strong correlations with the sulfate concentration ( $R = 0.65$  and  $R = 0.74$ , respectively).  
488 It is possible that the strong correlations of sulfate concentration with water-soluble V and Cd  
489 concentrations but not with V and Cd fractional solubilities were due to a large fraction of  
490 water-soluble V and Cd having the same sources as sulfate and its precursor (i.e.,  $\text{SO}_2$ ). For  
491 instance, Celo et al. (2015) reported that substantial concentrations of water-soluble aerosol  
492 metals (including V), sulfate, and  $\text{SO}_2$  are present in exhaust emissions from the main engines  
493 of commercial marine vessels.

494 **Table 1:** Spearman rank correlations between the metal fractional solubilities and  $W_i$  and  $H_{air}^+$   
495 in fine aerosols<sup>a</sup>

Metal	Sulfate	$W_i$	$H_{air}^+$	pH
Cr	<b>0.62</b>	<b>0.42</b>	<b>0.48</b>	-0.16
Al	0.14	0.08	0.14	-0.06
Fe	<b>0.53</b>	<b>0.31</b>	<b>0.50</b>	<b>-0.33</b>
Ni	0.03	0.01	0.18	<b>-0.26</b>
Co	<b>0.41</b>	<b>0.41</b>	<b>0.23</b>	-0.05
Cu	<b>0.74</b>	<b>0.72</b>	<b>0.24</b>	-0.07
Pb	<b>0.53</b>	<b>0.41</b>	<b>0.34</b>	-0.13
Mn	<b>0.49</b>	<b>0.43</b>	<b>0.23</b>	-0.01
V	0.14	0.01	0.21	-0.20
Cd	0.04	0.10	0.13	0.22

496 <sup>a</sup> Bold: statistically significant ( $p < 0.05$ )



497

498 **Figure 6:** Relationships between the metal fractional solubilities and sulfate mass  
 499 concentration in fine aerosols. Only data with non-zero total metal concentrations were used in  
 500 the figures. Also shown are the spearman correlation coefficients for each relationship. The  
 501 symbols are colored by the corresponding  $W_i$  concentrations calculated by ISORROPIA-II.  
 502 The  $W_i$  concentrations increased with sulfate concentrations.

503 High levels of aerosol acidity and liquid water are generally needed for the acid  
 504 dissolution of metals in an aqueous aerosol particle. In addition to being the main contributor  
 505 to aerosol acidity levels (i.e.,  $H_{air}^+$ ), sulfate is a highly hygroscopic species that will influence  
 506 the overall aerosol water uptake behavior, which will drive  $W_i$ . Sulfate was the main driver of  
 507  $W_i$  in fine aerosols in our study since the mass concentrations of nitrate (another highly  
 508 hygroscopic species) were very low (about 18 times lower than sulfate, on average). Both  $W_i$   
 509 and  $H_{air}^+$  were controlled primarily by sulfate (sulfate and  $W_i$   $R = 0.90, p < 0.05$ ; sulfate and

510  $H_{air}^+$   $R = 0.63$ ,  $p < 0.05$ ). Thus, we analyzed the relationships between the aerosol metal  
511 fractional solubilities and  $W_i$  and  $H_{air}^+$  (Figures S7 and S8). Table 1 shows that correlations  
512 between the Al, Ni, V, and Cd fractional solubilities and  $W_i$  and  $H_{air}^+$  were weak. Together, the  
513 weak correlations between the fractional solubilities of Al, Ni, V, and Cd and sulfate,  $W_i$ , and  
514  $H_{air}^+$  implied that acid processing may have played a minor role in enhancing the solubilities  
515 of these four metals. Other atmospheric processes beyond acid processing (e.g., cloud  
516 processing, photoreduction) could have played more important roles in enhancing the  
517 solubilities of these four metals (Zhu et al., 1993; Spokes et al., 1994; Kuma et al., 1995). It is  
518 possible that these four metals had slow acid dissolution rates as a result of their mineralogy  
519 and oxidation states. The impacts of mineralogy and oxidation states on the susceptibilities of  
520 water-insoluble Al, Ni, V, and Cd to acid dissolution are currently not known. However,  
521 previous studies showed that different aerosol Fe mineralogy and oxidation states have  
522 different susceptibilities to acid dissolution that will occur at different timescales (Ingall et al.,  
523 2018). Hence, analogous to Fe, it is possible that the mineralogy and oxidation states of Al, Ni,  
524 V, and Cd in the collected aerosols may have resulted in these four metals being less susceptible  
525 to acid processing, which in turn caused them to undergo slow sulfate-driven acid dissolution  
526 from water-insoluble forms to water-soluble forms.

527 Table 1 shows that the Cr, Fe, Co, Cu, Pb, and Mn fractional solubilities were positively  
528 correlated with  $W_i$  and  $H_{air}^+$ , and these correlations were statistically significant. Together, the  
529 statistically significant positive correlations between the fractional solubilities of Cr, Fe, Co,  
530 Cu, Pb, and Mn and sulfate,  $W_i$ , and  $H_{air}^+$  indicated that acid processing likely played an  
531 important role in enhancing the solubilities of these six metals. The fractional solubilities of  
532 Co, Cu, Pb, and Mn were more strongly correlated with the  $W_i$  concentration than with the  
533  $H_{air}^+$  concentration. This suggested that  $W_i$  had a stronger influence on the acid dissolution of  
534 Co, Cu, Pb, and Mn. The strong influence that  $W_i$  has on the metal fractional solubility could  
535 be explained by the role of aerosol water as a reaction medium for the acid dissolution of metals  
536 in an aqueous aerosol particle. Wong et al. (2020) previously showed that at a relatively  
537 constant aerosol pH, a decrease in  $W_i$  will lead to a decrease in the reaction medium volume,  
538 which in turn will lead to decreases in the overall formation rates of water-soluble metals.

539 Conversely, the fractional solubilities of Cr and Fe were more strongly correlated with the  $H_{air}^+$   
540 concentration than with the  $W_i$  concentration. This suggested that the aerosol acidity levels had  
541 a stronger influence on the acid dissolution of Cr and Fe. Despite the statistically significant  
542 positive correlations between the fractional solubilities of Cr, Fe, Co, Cu, Pb, and Mn and  
543 sulfate,  $W_i$ , and  $H_{air}^+$  (Table 1), there was significant scatter in the datasets (Figures 6, S7, and  
544 S8). This scatter could be a result of the sulfate,  $W_i$ ,  $H_{air}^+$ , total and water-soluble metal  
545 concentrations being substantially different in individual aerosol particles, which would not be  
546 captured by the bulk chemical analysis and thermodynamic modeling performed in this study.  
547 The metal dissolution rates in individual aerosol particles could also be significantly different  
548 due to differences in metal mineralogy, aerosol acidity levels, etc. in individual aerosol particles.  
549 In addition, a recent study by Yang et al. (2021) reported that the filterable metal fractions in  
550 the water extracts may contain some metals in water-insoluble forms with small diameters that  
551 allowed them to pass through the pores of syringe filters. This would result in over-estimated  
552 metal fractional solubilities, which could explain why some of the data points in Figure 6  
553 showed high metal fractional solubilities at low sulfate concentrations.

554 Interestingly, variability in the aerosol pH did not appear to be a key driver of the  
555 variability in the solubilities of Cr, Fe, Co, Cu, Pb, and Mn. It was difficult to discern aerosol  
556 pH-dependent fractional solubility trends for these six metals, and their fractional solubilities  
557 were not highly correlated with aerosol pH (Table 1 and Figure S9). This could be attributed  
558 partly to the scatter in the datasets caused by differences in the metal solubilities and pH in  
559 individual aerosol particles that would not be captured by the bulk chemical analysis and  
560 thermodynamic modeling performed in this study. The absence of obvious aerosol pH-  
561 dependent fractional solubility trends could also be due to the insensitivity of aerosol pH to the  
562 variability of sulfate ( $R = -0.22$ ,  $p < 0.05$ ). Based on Equation (1), the aerosol pH could be  
563 viewed simply as the ratio of  $H_{air}^+$  and  $W_i$ . Both  $W_i$  and  $H_{air}^+$  were highly variable in this study,  
564 and both were controlled primarily by sulfate. As a result, the ratio of  $H_{air}^+$  and  $W_i$ , or the  
565 aerosol pH, would be fairly insensitive to sulfate even though it was driven primarily by sulfate.  
566 Previous studies have similarly reported weak or the absence of aerosol pH-dependent metal  
567 fractional solubility trends despite evidence of aerosol metal dissolution being enhanced by

568 acid processing (Shi et al., 2020; Wong et al., 2020).

#### 569 **4. Conclusions**

570 In this study, we investigated the abundance and fractional solubilities of ten metals (Fe,  
571 Cu, Al, V, Cr, Mn, Co, Ni, Cd, and Pb) in size-fractionated aerosols collected at an urban site  
572 in Hong Kong. Weekly aerosol samples were collected for a month during different seasons  
573 from March 2021 to January 2022. The main objective of this study was to identify the key  
574 factors that controlled metal solubilities in fine aerosols, with a focus on aerosol metal  
575 dissolution via the acid processing and metal-organic complexation mechanisms. Hence, other  
576 aerosol chemical species were measured in addition to the total and water-soluble metals.

577 Higher mass concentrations of total metals were usually measured during the winter  
578 and/or spring seasons. This was likely due to the long-range transport of polluted air masses by  
579 northerly prevailing winds from emission sources located in continental areas north of Hong  
580 Kong. The total metals could be arranged in the following order based on their abundances:  
581  $Fe > Al > Cu > Co > Mn > Pb > Cr > Ni > V > Cd$ . This order of abundance was the same for  
582 both fine and coarse aerosols. The major sources of the total metals were sea salt, dust, ship  
583 emissions, and industrial activities. Higher mass concentrations of water-soluble metals were  
584 also usually measured during the winter and/or spring seasons. With the exception of Cu, the  
585 water-soluble metals had higher mass concentrations in fine aerosols than in coarse aerosols.  
586 The mass concentrations of water-soluble metals generally correlated with the mass  
587 concentrations of total metals, which implied that the water-soluble metals were largely derived  
588 from their total metals through atmospheric processing and/or that water-soluble and water-  
589 insoluble metals have the same emission sources. The study-averaged metal fractional  
590 solubilities spanned a wide range for both fine (7.8 % to 71.2 %) and coarse (0.4 % to 47.9 %)  
591 aerosols. With the exception of Cu and Co, the metals exhibited higher fractional solubilities  
592 in fine aerosols compared to coarse aerosols. The aerosol size-dependent metal fractional  
593 solubility could potentially be attributed to differences in the composition and metal  
594 mineralogy which resulted in different metal dissolution rates and/or mechanisms for aerosols  
595 of different sizes.



596 The fine aerosols collected in this study were mostly acidic, with about 60 % of the  
597 calculated pH values below 3. The acidic nature of the fine aerosols combined with oxalate  
598 (which forms metal-organic complexes easily) not being detected in our aerosol samples  
599 suggested that organic ligand-promoted dissolution likely played a minor role in enhancing  
600 aerosol metal solubilities. This is because organic ligand-promoted metal dissolution is a slow  
601 process, and it plays a minor role in metal dissolution under low pH conditions. Our analyses  
602 showed that sulfate, which is the dominant fine aerosol acidic species, exhibited statistically  
603 significant positive correlations with both the water-soluble concentrations of Cr, Fe, Co, Cu,  
604 Pb, and Mn and their fractional solubilities. In addition, sulfate controlled  $W_i$  and  $H_{air}^+$ , both of  
605 which are needed for acid dissolution of metals in an aqueous aerosol particle. The water-  
606 soluble concentrations of Cr, Fe, Co, Cu, Pb, and Mn and their fractional solubilities exhibited  
607 statistically significant positive correlations with both  $W_i$  and  $H_{air}^+$ . Together, the statistically  
608 significant positive correlations between the fractional solubilities of Cr, Fe, Co, Cu, Pb, and  
609 Mn and sulfate,  $W_i$ , and  $H_{air}^+$  indicated that acid processing likely played an important role in  
610 enhancing the solubilities of these six metals. The fractional solubilities of Co, Cu, Pb, and Mn  
611 were more strongly correlated with the  $W_i$  concentration than with the  $H_{air}^+$  concentration,  
612 which implied that  $W_i$  had a stronger influence on the acid dissolution of these four metals.  
613 The fractional solubilities of Cr and Fe were more strongly correlated with the  $H_{air}^+$   
614 concentration than with the  $W_i$  concentration, which implied that the aerosol acidity levels had  
615 a stronger influence on the acid dissolution of these two metals. Conversely, our analyses  
616 suggested that acid processing played a minor role in enhancing the solubilities of Al, Ni, V,  
617 and Cd. It is possible that the mineralogy and oxidation states of these four metals made them  
618 less susceptible to acid processing.

619 In conclusion, this study highlights the key role that sulfate plays in controlling the  
620 solubilities of a host of metals in fine aerosols (in this case, Cr, Fe, Co, Cu, Pb, and Mn). This  
621 is mostly due to sulfate's ability to both strongly acidify the aerosol particle and provide the  
622 liquid reaction medium needed for the acid dissolution of metals. Although this study was  
623 performed at an urban site in Hong Kong, we expect our findings to broadly apply to other  
624 urban areas in Hong Kong and South China, where sulfate is the dominant acidic and

625 hygroscopic component in fine aerosols. Results from this study can also provide insights into  
626 how the solubilities of different aerosol metals will change with the decrease in sulfate as Hong  
627 Kong and other cities in South China transition away from coal combustion as their main  
628 energy source to improve local and regional air quality and combat climate change.

629 **Data availability:** The data used in this publication is available to the community and can be  
630 accessed at: <https://doi.org/10.5281/zenodo.7013770> (Yang et al., 2022).

631 **Author contributions:** J.Y. and T.N. designed the study. J.Y. collected the field samples. J.Y.,  
632 L.M., and W.C.A. performed chemical analysis of the field samples. J.Y., X.H., Y.M., and T.N.  
633 analyzed the data. J.Y. and T.N. prepared the manuscript with contributions from all co-authors.

634 **Competing interests:** One of the authors is a member of the editorial board of *Atmospheric*  
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