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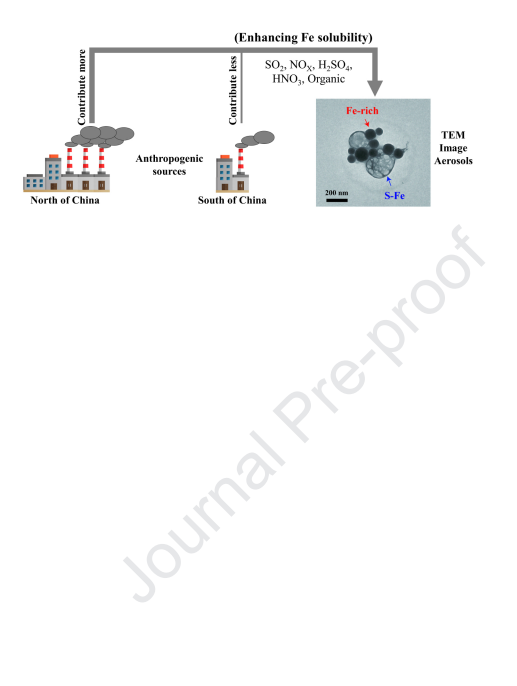
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Abstract

Soluble iron (Fe_s) in aerosols contributes to free oxygen radical generation with 21 implications for human health, and potentially catalyzes sulfur dioxide oxidation. It is 22 23 also an important external source of micronutrients for ocean ecosystems. However, factors controlling Fe_S concentration and its contribution to total iron (Fe_T) in aerosols 24 remain poorly understand. Here, Fe₅ and Fe_T in PM_{2.5} was studied at four urban sites 25 in eastern China from 21 to 31 December 2017. Average Fe_T (869-1490 ng m⁻³) and 26 Fe_s (24-68 ng m⁻³) concentrations were higher in northern than southern China cities, 27 but Fe solubility (%Fes, 2.7-5.0%) showed no spatial pattern. Correlation analyses 28 suggested %Fes was strongly correlated with Fes and PM_{2.5} instead of Fe_T 29 concentrations. Individual particle observations confirmed that more than 65% of 30 nano-sized Fe-containing particles were internally mixed with sulfates and nitrates. 31 Furthermore, there was a high correlation between sulfates or nitrates/Fe_T molar ratio 32 and %Fes. We also found that the sulfates/nitrates had weaker effects on %Fes at RH 33 < 50% than at RH > 50%, suggesting RH as indirect factor can influence %Fes in 34 PM_{2.5}. These results suggest an important role of chemical processing in 35 enhancing % Fe_s in the polluted atmosphere. 36

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Capsule abstract: Iron solubility related to sulfate and nitrate in fine particles in
polluted ambient air.

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41 Keywords: Polluted air; Bulk aerosol analysis; Individual particles analysis; Fe
42 solubility; Atmospheric acidification processing

44 1. Introduction

Iron (Fe) as an important aerosol component is an essential external source for 45 phytoplankton growth in large parts of the remote oceans; it indirectly modulates CO₂ 46 sequestration, and it thus has feedback effects on the global carbon cycle, and climate 47 (Martin and Fitzwater, 1988; De Baar et al., 1995; Jickells et al., 2005; Tagliabue et al., 48 2017; Matsui et al., 2018). Fe-containing fine particles can adversely affect human 49 health via reactive oxygen species (ROS) formation (Smith and Aust, 1997; Park et al., 50 51 2006; Abbaspour et al., 2014). In addition, Fe in aerosol particles or cloud droplets can convert S(IV) to S(VI) by catalytic oxidation, which is a substantial pathway for 52 atmospheric sulfate production (Alexander et al., 2009). These roles of Fe largely 53 depend on the fractional solubility of aerosol Fe (Shi et al., 2012), thus, crucial factors 54 and mechanisms that influence aerosol Fe solubility (%Fes) (the concentration ratio of 55 soluble Fe (Fe_S) and total Fe (Fe_T)) need to be better understood. 56

Fe has natural (e.g., desert dust and soil dust) and anthropogenic (e.g., fossil fuel 57 combustion and steel industrial activities) sources (Mahowald et al., 2005; Jickells et 58 59 al., 2005; Sedwick et al., 2007). Different sources have different %Fes, spanning three orders of magnitude (0.04-81%) (Schroth et al., 2009). Natural emissions are the 60 major sources for Fe_T with a contribution of 70-80% in global air (Jickells et al., 61 2005), while their %Fes is less than 1% (Schroth et al., 2009). Although the 62 contribution of anthropogenic sources to Fe_T is small compared with that from natural 63 sources, their contribution to %Fe_S is much higher (0.06-81%) (Schroth et al., 2009; 64 Oakes et al., 2012). The anthropogenic Fe emissions are strongly associated with 65 anthropogenic combustion sources in regions afflicted with elevated air pollution 66 67 levels (Guieu et al., 2005; Lough et al., 2005; Sedwick et al., 2007; Zhang et al., 2019). Therefore, it is important to understand %Fes in continental air polluted by 68 various anthropogenic sources. 69

Chemical processing of aerosols during transport and aging in the atmosphere has been hypothesized to influence %Fe_s (Shi et al., 2011; Ito, 2015; Shi et al., 2015; Lin et al., 2019; Xie et al., 2020). Aerosol acidification involving anthropogenic pollutants was thought to be an important hypothesis: acids produced from anthropogenic

74 pollutants can dissolve aerosol Fe, thus increasing %Fe_s (Meskhidze et al., 2003; Rubasinghege et al., 2010; Zhang et al., 2018). Several studies have estimated aerosol 75 acidity during air polluted periods, but the results differ widely, pH ranging from close 76 to 2 (highly acidic) to about 7 (neutral) in North China based on the chemical 77 modelling calculations (Cheng et al., 2016; Wang et al., 2016; Shi et al., 2017; Guo et 78 al., 2017; He et al., 2018). Such a large pH discrepancy is still under debate because 79 no direct method has been used to measure pH value of individual particles until now. 80 81 As we know, Fe oxides can be dissolved into Fe_s in aerosol particles under pH < 4 (Shi et al., 2012). Recently, Li et al. (2017) confirmed at the first time that the Fe_s can 82 dissolve from Fe oxides mixed in acidic sulfate particles over East China Sea using 83 transmission electron microscopy (TEM) and nanoscale secondary ion mass 84 spectrometry (NanoSIMS) analysis methods. As this way, if the detailed information 85 of Fe_s can be obtained from bulk aerosol samples, we can provide direct evidence for 86 the fine particles are acidic in bulk sample level. Therefore, understanding mass 87 concentrations of Fe_T and Fe_S as well as the corresponding %Fe_S can be one direct 88 89 evidence to show aerosol acidity.

In this study, we collected PM_{2.5} and individual particle samples at four urban sites
of East China, and combined bulk aerosol and individual particle chemical analysis
techniques to investigate: (1) the concentrations of Fe_T, Fe_S, and corresponding %Fe_S;
(2) factors influencing %Fe_S, including Fe_S concentration, PM_{2.5} concentration, Fe_T
concentration, atmospheric acidification processing, mixing state of Fe-containing
particles, and relative humidity (RH).

96 2. Experimental methods

97 2.1. Sampling site

Four urban areas were selected to represent typical urban environments: Beijing, Handan, and Zhengzhou in the North China Plain (NCP), and Hangzhou in the Yangtze River Delta (YRD) of southern China (Fig. 1). A population in 2018 is about 21.5, 9.5, 10.1, and 9.8 million in Beijing, Handan, Zhengzhou, and Hangzhou city. The sampling sites in Beijing, Handan, Zhengzhou, and Hangzhou were located in China University of Mining and Technology (Beijing) (CUMTB), Hebei University of

Engineering (HUE), Zhongyuan University of Technology (ZUT), and Zhejiang University (ZU), respectively. The sampling instruments at each sampling site were installed on the rooftop of an academic building with a height of about 15 m above the ground. The surrounding environments of CUMTB, HUE, ZUT, and ZU are similar. They are all situated in the center of the corresponding city, and surrounded by intensive university and residential buildings, business offices and urban streets.

Beijing, the capital of China, is the national center for politics and culture. As a 110 megacity, Beijing mainly suffers from vehicular exhaust pollution. Emissions in 111 Beijing's neighboring regions also significantly influence its air quality due to 112 long-range transport of air pollutants. Zhang et al. (2016) suggested that the regional 113 transport of pollutants contributed 28-36% of PM_{2.5} in Beijing. The annual average 114 concentration of PM_{2.5} was 51 µg m⁻³ in 2018 (source from 2018 Beijing State of 115 Ecological Environment Bulletin), which exceeded the national standard (35 $\mu g \ m^{-3})$ 116 of China. 117

Handan in northern China is a heavy-industry city with principle industries for steel, coal, cement, coke and electric power generation, whose contribution to Handan GDP has now reached as high as 45% (Handan Statistical Yearbook, 2018). This high energy consumption has resulted in copious emissions of air pollutants. Handan city is among the most polluted cities in China with the annual average $PM_{2.5}$ concentration in 2018 at 69 µg m⁻³ (source from 2018 Hebei Province Ecology and Environment Condition Statement).

25 Zhengzhou in central China is a coal-driven energy consumption city, with coal 26 burning accounting for about 70% of energy consumption (Jiang et al., 2017). As a 27 hub of the country's major railway, motorway and aviation transportation, Zhengzhou 28 suffers from serious vehicular exhaust pollution. Zhengzhou is often ranked as among 29 the top ten most polluted cities in China with an annual average $PM_{2.5}$ concentration 30 of 63 µg m⁻³ in 2018 (source from 2018 Zhengzhou Environmental Quality Bulletin).

Hangzhou in southeastern China is the second largest city in the Yangtze River
Delta (YRD). As one of the most beautiful cities in China, industrial activities in
Hangzhou are minor. Traffic emission is one of the most important sources for

Hangzhou air pollution. In addition, pollutants emitted in northern China or in surrounding regions such as some heavy industries in Ningbo are transported into the city to significantly degrade its air quality. The annual average $PM_{2.5}$ concentration was 40 µg m⁻³ in 2018 (source from 2018 Hangzhou Environmental Status Bulletin).

138 **2.2. Sample collection**

PM_{2.5} and individual particle samples were collected at the four sampling sites from 139 21 to 31 December, 2017, only on days without rain. PM_{2.5} samples were collected on 140 141 90 mm diameter quartz filters for 11.5 h (daytime: 08:30-20:00; nighttime: 20:30-08:00 (next day)) using a TH-16A Intelligent PM_{2.5} sampler at a flow rate of 142 100 L min⁻¹ (Wuhan Tianhong Corporation, China). Before and after collection, the 143 flow rate was calibrated. Daytime and nighttime blank samples were collected using 144 the same method, but without pumping. Before sample collection, all quartz filters 145 were baked at 600 °C in a muffle furnace for 4 h to remove any possible contaminants. 146 After baking, the quartz filters were placed in a room with temperature of 20 ± 1 °C 147 and RH of $50 \pm 2\%$ for 24 h, then, they were weighed using a Sartorius analytical 148 balance (detection limit 0.001 mg). After sample collection, the loaded filters were 149 similarly conditioned and weighed. Difference value of the two weighed mass divide 150 by sample volume was PM_{2.5} concentration. 151

Individual particle samples were also collected on copper grids coated with carbon 152 film by a single-stage cascade impactor with a 0.3 mm diameter jet nozzle and a flow 153 of 1.0 L min⁻¹. Individual particle samples were collected four times each day at 8:00, 154 12:00, 18:00 and 0:00. The sampling duration spanned 30 s to 8 min depending on the 155 PM_{2.5} mass concentration. The collection efficiency of the single-stage cascade 156 impactor is 50% for aerodynamic diameter of 0.1 µm particles and a density of 2 g 157 cm⁻³. After sampling, the grids were placed in a sealed dry plastic tube and stored at 158 25° C and $20 \pm 3\%$ RH in a desiccator. 159

- Meteorological data were measured and recorded every 5 min by an automatedweather instrument (Kestrel 5500, USA).
- 162 **2.3. Fe extraction procedure**
- **163 2.3.1 Fe_T fraction**

The microwave acid digestion was employed to digest the quartz fiber-filter 164 samples into liquid solution for Fe analysis. Firstly, the digestion vessels were cleaned 165 by ultrasonification with ultra-pure water (18.2 M Ω) for 15 min, then with 5% HNO₃ 166 for 15 min, and finally with ultra-pure water for 15 min. Then, one quarter of the 167 sample filters were placed in the digestion vessel with a mixed-acid solution 168 consisting of 6 ml nitric acid (65%, Merck, Germany), 2 ml hydrogen peroxide (> 8%, 169 Beijing Institute of Chemical Reagents, China) and 0.6 ml hydrofluoric acid (40%, 170 171 Merck, Germany). After closing the vessels, the samples were digested by a microwave digestion system (MARS 5, CEM Corporation, Matthews, NC, USA) on 172 the basis of a temperature-controlled procedure, increasing to 120 °C in 8 min and 173 holding for 3 min, then increasing to 160 °C in 10 min and holding for 10 min, and 174 finally increasing to 190 °C for 10 min and holding for 55 min. After cooling to room 175 temperature, the digested materials was transferred to cleaned brown PTFE bottles 176 and diluted to 100 ml using ultra-pure water. Three blank filters for each sampling site 177 were treated in the same manner as the samples. 178

179 **2.3.2 Fes fraction**

Ultrasonification was used to extract the water-soluble fraction of the samples 180 filters for Fe_s analysis following the procedure described by Kanai et al. (2003). One 181 quarter of the sample filters were placed in clean tubes with 15 ml ultra-pure water. 182 183 Then, the tubes were placed in an ultrasonic bath in ultra-pure water for 60 min. The extracts filtered through a 0.22 PTFE water were μm pore size 184 (polytetrafluoroethylene) syringe filter into cleaned brown PTFE bottles, and 185 subsequently acidified with ultra-pure concentrated HNO₃ to 0.4% v/v HNO₃. Three 186 187 blank filters for each sampling site were treated in the same manner as the samples. All solutions were stored at 4 °C until instrumental analysis. 188

189 2.4. Analytical procedures of Fe

The concentrations of the total and water-soluble fractions of Fe were determined by inductively coupled plasma mass spectrometry (ICP-MS, Agilent 7500ce). Detailed descriptions of the procedure were given in Pan et al. (2013). Briefly, according to the standard procedures and criteria specified in the manufacturer's

manual, the ICP-MS was optimized daily by a tuning solution containing Li, Y, Tl, Ce 194 and Co. External calibration standards (Agilent Technologies, Environmental 195 Calibration Standard) were employed to quantify the Fe, and an internal standard 196 (containing ⁴⁵Sc, ⁷²Ge, ¹⁰³Rh, ¹¹⁵In, ¹⁵⁹Tb, ¹⁷⁵Lu and ²⁰⁹Bi) was added online during Fe 197 analysis. The two certified materials (soil: GBW07401, fly ash: GBW08401) were 198 digested and analyzed in the same manner as the samples for recovery calculation. 199 The recovery of Fe was greater than 95%. Moreover, no significant Fe was found in 200 201 the field and reagent blank samples. The detection limits of Fe_T and Fe_S were 0.15 and 2.43 μ g l⁻¹, respectively. 202

203 2.5. Analysis of water-soluble ions, organic carbon, and elemental carbon

Water-soluble ions were analyzed by ion chromatography (Dionex ICs-90, Dionex Corporation, USA). Detailed descriptions about the analytical method were given in Zhang et al. (2017).

Organic carbon (OC) and elemental carbon (EC) were analyzed by a Sunset Laboratory carbon analyzer with the thermal-optical transmittance method. Organic matter (OM) concentrations were obtained by multiplying the OC concentration by 1.91, as reported in Xing et al. (2013).

211 **2.6. Individual particle analysis**

The copper grids were analyzed by a JEOL JEM-2100 transmission electron 212 microscope (TEM) combined with an energy-dispersive X-ray spectrometer (EDS). 213 1613 particles in Beijing samples, 1667 particles in Handan samples, 1523 particles in 214 Zhengzhou samples and 1833 particles in Hangzhou samples were analyzed by the 215 TEM/EDS at 200 kV. TEM does an excellent job of determining the morphology and 216 mixing state of individual particles; EDS detects the main elements above carbon (\geq 217 12). Copper was not included in the analyses due to the interferences of the copper 218 TEM grids. EDS collection duration was limited to 15 s to reduce beam damage. Five 219 areas from center to periphery of the grids were chosen for analysis to ensure their 220 representativeness. Equivalent circle diameters (ECDs) of the particles were identified 221 222 by iTEM software (Olympus Soft Imaging Solutions GmbH, Germany).

223 3. Results and discussion

224 **3.1. Overview of PM_{2.5} pollution**

 $PM_{2.5}$ concentrations were $155 \pm 60 \ \mu g \ m^{-3}$ in Beijing, $237 \pm 71 \ \mu g \ m^{-3}$ in Handan, 225 $179 \pm 90 \ \mu g \ m^{-3}$ in Zhengzhou, and $93 \pm 18 \ \mu g \ m^{-3}$ in Hangzhou (Table 1) during 226 21-31 December, 2017, which were all higher than the national daily PM_{2.5} standard 227 of 75 μ g m⁻³. Even the lowest PM_{2.5} concentrations in Beijing (74 μ g m⁻³), Handan 228 (117 µg m⁻³), Zhengzhou (51 µg m⁻³), and Hangzhou (71 µg m⁻³) were close or higher 229 than 75 μ g m⁻³. The day number that PM_{2.5} concentration exceeded 75 μ g m⁻³ to total 230 observation days were 10/12, 13/13, 13/14, and 17/18 in Beijing, Handan, Zhengzhou, 231 and Hangzhou city, respectively. In general, PM_{2.5} concentrations were 1.7-2.6 times 232 higher in Beijing, Handan, and Zhengzhou cities in the NCP than in Hangzhou city in 233 the YRD. 234

OM was the most abundant chemical component in $PM_{2.5}$ in Beijing, Handan, Zhengzhou and Hangzhou cities with contributions of 40%, 31%, 29%, and 31%, respectively (Fig. S1). The next most abundant components in $PM_{2.5}$ were nitrate (NO₃⁻), sulfate (SO₄²⁻), and ammonium (NH₄⁺) with contributions of 13%, 8%, and 7% in Beijing, 14%, 9%, and 7% in Handan, 17%, 8%, and 7% in Zhengzhou, and 20%, 9%, and 8% in Hangzhou.

241 **3.2. Overview of individual particles data**

A total of 1613, 1667, 1523, and 1833 individual aerosol particles collected in 242 Beijing, Handan, Zhengzhou, and Hangzhou cities, were analyzed by TEM/EDS 243 (Table S1). Based on elemental composition and morphology of individual particles, 244 the internally mixed sulfate particles (e.g., S-OM, S-rich, S-soot, S-fly ash, and S-Fe) 245 were dominant in all the analyzed particles, which were 68% in Beijing, 62% in 246 247 Handan, 63% in Zhengzhou, and 73% in Hangzhou (Fig. S2). All internally mixed sulfate particles contain S. Because of the detection limitation of TEM/EDS for 248 ammonium nitrate, the technique could not quantify nitrates in individual particles. 249 However, some studies already confirmed that sulfate particles normally contained 250 secondary nitrates in individual secondary particles in urban air (Li et al. 2016; 251 Riemer et al., 2019). 252

253 **3.3. Fe solubility**

Table 1 presents the concentrations of Fe_T and Fe_S in PM_{2.5} as well as %Fe_S at the 254 four sites. The average Fe_T concentration was 1490 ± 428 ng m⁻³ in Beijing, $1310 \pm$ 255 271 ng m⁻³ in Handan, 1132 \pm 467 ng m⁻³ in Zhengzhou, and 869 \pm 215 ng m⁻³ in 256 Hangzhou during 21-31 December, 2017, accounting for $1.14 \pm 0.60\%$, $0.60 \pm 0.21\%$, 257 $0.90 \pm 0.58\%$, and $0.95 \pm 0.31\%$ of PM_{2.5}, respectively. The average Fe_S concentration 258 was 68 \pm 46 ng m^-3 in Beijing, 59 \pm 33 ng m^-3 in Handan, 32 \pm 20 ng m^-3 in 259 Zhengzhou, and 24 ± 8.5 ng m⁻³ in Hangzhou (Table 1). Fe_T and Fe_S concentrations in 260 the cities of NCP were 1.3-1.7 and 1.3-2.8 times higher than that in the city of YRD, 261 respectively. Here, we calculated %Fes as Fes concentration/Fe_T concentration \times 262 100%. The results showed that the average %Fes was $5.0 \pm 3.8\%$ in Beijing, $4.5 \pm 2.6\%$ 263 in Handan, $2.7 \pm 1.5\%$ in Zhengzhou, and $3.0 \pm 1.1\%$ in Hangzhou (Table 1). %Fe_S in 264 Zhengzhou was lower than that in Hangzhou, although Fe_T and Fe_S concentrations in 265 the former were higher than the latter. 266

We compared the measurements of Fe_T and %Fe_S with those in the marine 267 atmosphere. Table 2 shows that Fe_T concentrations (869-1490 ng m⁻³) in this study are 268 much higher than those in the marine atmosphere, ranging from 28.4 ng m^{-3} over the 269 Pacific Ocean (Buck et al., 2013), 218 ng m⁻³ in the North Atlantic Ocean (Buck et al., 270 2010), 590 ng m⁻³ at the Bay of Bengal (Srinivas et al., 2012), to 761 ng m⁻³ at the 271 East China Sea (Hsu et al., 2010). In contrast, %Fes (2.7%-5.0%) in this study is 272 273 1.2-3.3 times lower than those in the marine atmosphere, ranging from 6.0% in the Bay of Bengal, 7.7% in the East China Sea, 8.1% in the Pacific Ocean, to 9.0% in the 274 North Atlantic Ocean. These results indicate that long-range transport of 275 Fe-containing particles significantly increases the %Fe_s in fine particles. 276

277 **3.4. Factors influencing Fe solubility**

278 3.4.1 Correlations between %Fe_s and PM_{2.5}, Fe_s, Fe_T

% Fe_S had strong correlations with Fe_S at all four urban sites with correlation coefficients of 0.81-0.96 (Fig. 2). %Fe_S and PM_{2.5} also had high correlations with the correlation coefficients at 0.58-0.93, but %Fe_S had no obvious correlations with Fe_T except the Hangzhou site. In addition, Figure S3 shows that %Fe_S generally displays similarly variation trend with PM_{2.5} and Fe_S concentrations, but different from Fe_T.

3.4.2 Potential chemical processing in enhancing Fe solubility 284

To understand what controled the solubility of Fe, we compared %Fes on 285 non-haze, light haze, intermediate haze and heavy haze days (Fig. S4). Here we 286 defined non-haze days as daily PM_{2.5} concentration \leq 75 µg m⁻³, light haze days as 287 $75 < PM_{2.5} \le 150 \ \mu g \ m^{-3}$, intermediate haze days as $150 < PM_{2.5} \le 250 \ \mu g \ m^{-3}$, and 288 heavy haze days as $> 250 \ \mu g \ m^{-3}$. Figure S4 shows that %Fe_s range from 0.9% to 1.2% 289 (average: $1.0\% \pm 0.2\%$) on non-haze days, 1.4% to 6.3% (average: $2.9\% \pm 1.2\%$) 290 291 on light haze days, 1.5% to 10.6% (average: $4.4\% \pm 2.7\%$) on intermediate haze days and 3.8% to 11.4% (average: $6.5\% \pm 2.6\%$) on heavy haze days. In a word, the %Fes 292 significantly increased from non-haze to heavy haze days at each sampling site. 293 There are two possible reasons to explain the increased %Fe_S following the heavy 294 295 haze formation: (1) changes in sources, and (2) chemical processing.

Here we firstly investigated the Fe_T contributions from various primary emissions, 296 which were calculated by Fe_T concentrations dividing particulate matter 297 concentrations in different sources. Secondly, %Fes was investigated from various 298 299 primary emissions, and calculated by Fe_s concentrations dividing Fe_T concentrations in different sources. Fer contribution ranged approximately from 3.7%-11.9% for 300 coal combustion (Desboeufs et al., 2005; Fu et al., 2012), 0.4%-3.3% for biomass 301 burning (Yamasoe etal., 2000; Lee et al., 2005; Fuzzi et al., 2007; Fu et al., 2012), 302 303 0.86%-9.3% for oil combustion (Desboeufs et al., 2005; Schroth et al., 2009; Fu et al., 2012), and 3.1%-8.5% for mineral dust (Schroth et al., 2009; Fu et al., 2012; Shi 304 et al., 2011, 2012). % Fes ranged from 0.06%-0.2% for coal combustion (Desboeufs 305 et al., 2005; Oakes et al., 2012), 2%-46% for biomass burning (Guieu et al., 2005; 306 Bowie et al. 2009; Oakes et al., 2012), 35.7%-77% for oil combustion (Desboefus et 307 al. 2005; Schroth et al., 2009; Oakes et al., 2012), and 0.04%-0.54% for mineral dust 308 (Schroth et al., 2009; Oakes et al., 2012; Shi et al., 2012). In addition, our previous 309 study showed that %Fes in smelter particles (e.g., Fe oxides) from industrial 310 emissions was extremely low (Li et al., 2017). For this type of particles, we used 0.1% 311 as a conservative value. 312

313

The contributions of coal combustion, biomass burning, oil combustion, mineral

dust, and industrial emission to PM_{2.5} were 16%-57%, 7%-11.2%, 2%-17.1%,
10%-23.1%, and 12%-20% in Beijing (Yu et al., 2013; Zhang et al., 2013; Ma et al.,
2017), 22.3%-25.9%, 6.3%-10.6%, 10.2%-12.8%, 9.1%-10.9%, and 16.2%-24.2% in
Handan (Wei et al., 2014; Meng et al., 2016; Wang et al., 2015), 14%-23%,
12%-13%, 7%-23%, 8%-26%, and 4%-26% in Zhengzhou (Geng et al., 2013; Wang
et al., 2017; Jiang et al., 2018), and 12.8%-16.7%, 4%-14%, 10.2%-22%, 2%-8.2%,
and 2%-9% in Hangzhou (Zhen et al., 2010; Liu et al., 2015).

321 Based on above data, we can know that coal combustion, industry and mineral dust have extremely low %Fes (< 1%). Although biomass burning and oil 322 combustion having high %Fe₅, their Fe_T contribution to fine particles are low. Using 323 the equation of Fe_T content $\times PM_{2.5}$ source apportionment data $\times 100\%$, we find that 324 the Fe_T contributions of biomass burning and oil combustion are less than 3.0% in 325 $PM_{2.5}$. In a word, even though the solubility in the two sources is high, their 326 contribution to the Fes is low due to their small contribution to FeT. Therefore, 327 variations in primary emissions alone are not able to explain the enhanced %Fes 328 329 during the haze days, which suggests that chemical processing is the key reason leading to enhanced % Fes during haze events. 330

TEM observations further support this argument. We clearly identified abundant 331 fine Fe-containing particles (including Fe-rich and S-Fe particles) in the samples 332 with a size range of 25 nm to 4 µm (Fig. 4). The number contribution of 333 Fe-containing particles to the total analyzed particles was 9.2% in Beijing, 7.7% in 334 Handan, 6.6% in Zhengzhou, and 5.2% in Hangzhou (Table S1). In particular, we 335 found that S-Fe particles (internally mixed with secondary inorganic aerosols) were 336 337 dominant in Fe-containing particles. S-Fe particles accounted for 77%, 74%, 68% and 85% in all the Fe-containing particles in Beijing, Handan, Zhengzhou, and 338 Hangzhou, respectively (Table S1). TEM/EDS showed that secondary inorganic parts 339 in S-Fe particles more or less contained elemental Fe (Fig. 3). The phenomenon is 340 consistent with the findings of Li et al. (2017). 341

Size distributions of individual particles in Figure 4 showed that the peaks of Fe-rich particles were at 325 nm, 225 nm, 175 nm, and 175 nm in Beijing, Handan, 12

Zhengzhou and Hangzhou, while the corresponding internally mixed S-Fe particles
had peaks at 625 nm, 575 nm, 625 nm and 625 nm, respectively. Thus, secondary
sulfate/nitrate uptake led to an increase in particle size by 48%-72%.

Figure 5 provides further evidence to support the potential role of acidic species in 347 the enhancement of %Fes in PM_{2.5} during haze days. Hsu et al. (2014) have used the 348 molar ratio of acidic components to Fe_T to indicate the influence of aerosol 349 acidification on the %Fes. In this study, we followed the method to investigate the 350 impact of aerosol acidification on %Fes. NO3⁻/Fe_T and SO4²⁻/Fe_T molar ratios 351 showed high correlations with %Fes at each sampling site (r > 0.7). This suggested a 352 potential role of secondary species, such as sulfuric acid in the dissolution of 353 insoluble Fe in fine particles (Fig. 3 and 5). 354

RH is influential in the formation and phases of SO_4^{2-} and NO_3^{-} in the polluted air 355 of East China (Sun et al., 2018; Wu et al., 2018; Zhu et al., 2020). As a result, RH 356 should be an impact factor on %Fes by influencing secondary sulfate and nitrate 357 formation. Sun et al. (2018) showed that solid phases particle started to convert to 358 359 solid-liquid mixed phase particles when RH was > 50% in polluted urban air. Therefore, we assumed RH at 50% as the threshold of wet aerosols. Here, we 360 investigated the combined effects of RH and aerosol acidic species on %Fes (Fig. 6). 361 Hsu et al. (2010) reported that the dissolution of aerosol Fe was enhanced by the 362 presence of acidic constituents. SO_4^{2-} and NO_3^{-} , as the two major acidic constituents 363 in PM_{2.5}, were examined in this study. We used the molar ratio of $[2SO_4^{2-} +$ 364 NO_3 /Fe_T to represent the acidification degree. Figure 6 shows that %Fe_S ranged 365 from 0.7% to 3.8% in four cities at RH < 50%, even samples with a high degree of 366 367 acidification. However, %Fe₈ ranged from 1.3% to 11.4% at RH > 50% in four cities. Therefore, our results suggested that the sulfates/nitrates had a weaker effect 368 on %Fe_s at RH < 50% than at RH > 50% at all the sampling sites. Indeed, RH 369 showed high correlations with %Fe_s in Beijing, Handan, Zhengzhou and Hangzhou, 370 their correlation coefficients ranged from 0.51 to 0.92 (Fig. S5). In a word, RH 371 appears to be an indirect factor influencing the %Fes in fine particles. 372

4. Conclusions and atmospheric implications

374 The study suggests that acidic species contribute to the dissolution of Fe in the internally mixed particles collected in the four polluted urban sites. Our individual 375 particle analysis suggests that most of Fe-containing inorganic particles with size less 376 than 1 μ m in urban air have undergone acidic processes. A recent study shows that 377 aerosol acidity increases with decreasing particle size generated from the 378 (NH₄)₂SO₄-H₂SO₄ solution (Crag et al, 2018). Since most of the Fe-containing 379 particles are small with the peak size of Fe-rich particles of 175 to 325 nm and S-Fe 380 381 particles of 575 to 625 nm (Fig. 4), these particles may tend to be more acidic, even though the bulk aerosol pH may be higher (Shi et al., 2017; Liu et al., 2017; Song et 382 al., 2018). 383

The presence of large amount of Fe_s may catalyze the reactions for secondary 384 sulfate formation in polluted air of China. How the Fe_s, as the dominant soluble metal 385 in fine particles, changes the heterogeneous uptake of hydroxyl peroxy radicals (HO_2) 386 on aerosol should be paid more attention in polluted air in East China (Zou et al., 387 2019). Moreover, large amounts of tiny Fe particles and their associated Fes can be 388 389 inhaled into the respiratory tract, even into lung tissues, and can cause adverse health effects in urban cities through the generation of oxygen free radicals (Gonet and 390 Maher, 2019). 391

Under prevailing westerly winds in winter, these Fe-containing particles in Beijing, 392 Handan, Zhengzhou and Hangzhou urban areas can be transported into the East China 393 Sea and possibly influence the oceanic ecosystem. Li et al. (2017) collected 394 atmospheric particles during a research cruise over the East China Sea, and found that 395 14% of all analyzed particles were Fe-containing particles, and among them, 75% 396 397 were internal mixtures of sulfate coating and Fe inclusions. Takahashi et al. (2013) observed that anthropogenic Fe emitted from megacities in Eastern Asia was the most 398 important contributor to Fes in the North Pacific Ocean. Our study shows that these 399 anthropogenic Fe particles have already been partially dissolved into Fes in aerosols 400 401 before leaving the continental air.

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Table 1

 $PM_{2.5}$, Fe_T , and Fe_S concentrations as well as % Fe_S at the four urban sites from NCP to YRD (the numbers in parentheses were minimum and maximum).

	Beijing	Handan	Zhengzhou	Hangzhou
PM _{2.5} (µg m ⁻³)	155 ± 60 (74-270)	237 ± 71 (117-378)	179 ± 90 (51-357)	93 ± 18 (71-156)
$\operatorname{Fe}_{\mathrm{T}}(\mathrm{ng}\ \mathrm{m}^{-3})$	1490 ± 428 (971-2601)	1310 ± 271 (977-1996)	1132 ± 467 (342-1945)	869 ± 215 (433-1258)
$\operatorname{Fe}_{S}(\operatorname{ng m}^{-3})$	68 ± 46 (15-148)	59 ± 33 (16-119)	32 ± 20 (2.4-72)	24 ± 8.5 (13-53)
%Fe _S	$5.0\pm 3.8\;(0.911)$	$4.5\pm 2.6\ (0.7\text{-}9.6)$	$2.7 \pm 1.5 \ (0.7-5.6)$	3.0 ± 1.1 (1.2-5.5)

Table 2

Total Fe (Fe_T) concentration and Fe solubility (%Fe_S) reported in this study and literature data from ocean sites in the world.

Location	Туре	Sampling period	Size	Fe _T , ng m ⁻³	%Fes	References
Beijing	Urban	21-31 December 2017	PM _{2.5}	1490	5.0	This study
Handan	Urban	21-31 December 2017	PM _{2.5}	1310	4.5	This study
Zhengzhou	Urban	21-31 December 2017	PM _{2.5}	1132	2.7	This study
Hangzhou	Urban	21-31 December 2017	PM _{2.5}	869	3.0	This study
North Atlantic Ocean	Ocean	20 June-7 August 2003	TSP	218	9.0	Buck et al., 2010
Pacific Ocean	Ocean	2004-2006	TSP	28.4	8.1	Buck et al., 2013
Bay of Bengal	Ocean	March-April 2006	TSP	590	6.0	Srinivas et al., 2012
East China Sea	Ocean	Spring 2005- Spring 2007	TSP	761	7.7	Hsu et al., 2010

Figure Captions

Fig. 1. Map showing the locations of Beijing, Handan, Zhengzhou and Hangzhou sampling sites. The map is color-coded by surface elevation heights, which were obtained from SRTM (Shuttle Radar Topography Missionv) data (http://srtm.csi.cgiar.org/srtmdata/).

Fig. 2. Correlations of %Fe_s and Fe_T (ng m⁻³) (red), PM_{2.5} (μ g m⁻³) (blue) and Fe_s (ng m⁻³) (black) at the Beijing (a), Handan (b), Zhengzhou (c) and Hangzhou (d).

Fig. 3. TEM images and EDS of Fe-containing particles in this study: (a) TEM image of Fe-containing particle, (b) EDS of Fe-rich particle, (c) EDS of S-Fe particle.

Fig. 4. Size distributions of Fe-rich particle (blue) and internally mixed S-Fe particle (green) at the four urban sites. The distribution pattern is normalized.

Fig. 5. Correlations between %Fe_s and NO₃⁻/Fe_T (red) and SO₄²⁻/Fe_T (blue) molar ratio at Beijing (a), Handan (b), Zhengzhou (c) and Hangzhou (d).

Fig. 6. Relationships of %Fe_s with RH and acidification degree molar ratio $(2SO_4^{2-} + NO_3^{-})/Fe_T$ in Beijing (red), Handan (blue), Zhengzhou (green) and Hangzhou (purple).

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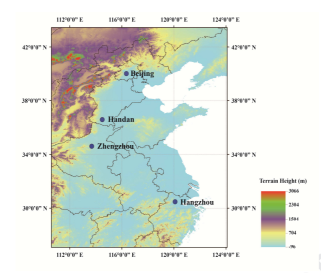


Fig. 1. Map showing the locations of Beijing, Handan, Zhengzhou and Hangzhou sampling sites. The map is color-coded by surface elevation heights, which were obtained from SRTM (Shuttle Radar Topography Missionv) data (http://srtm.csi.cgiar.org/srtmdata/).

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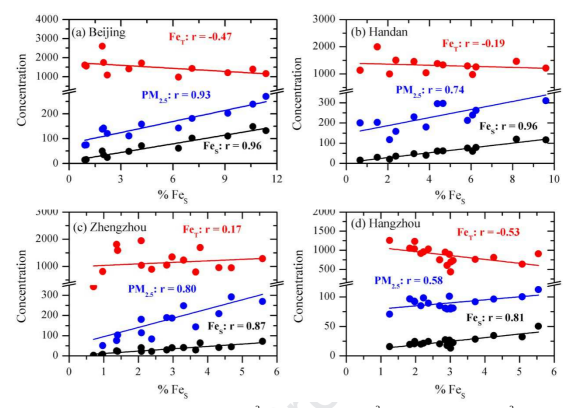


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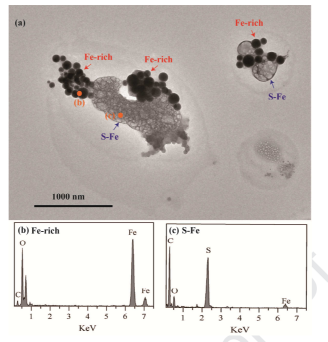


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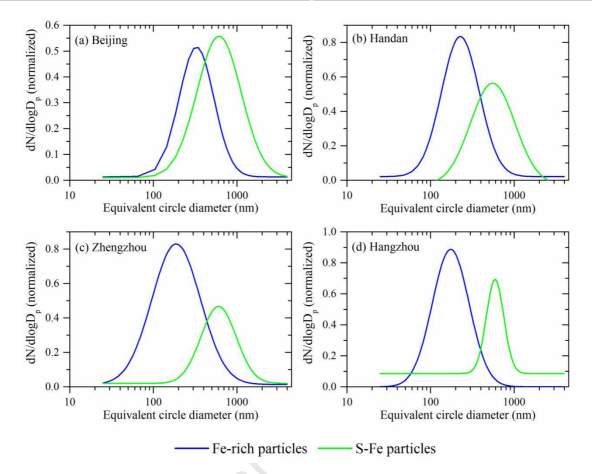


Fig. 4. Size distributions of Fe-rich particle (blue) and internally mixed S-Fe particle (green) at the four urban sites. The distribution pattern is normalized.

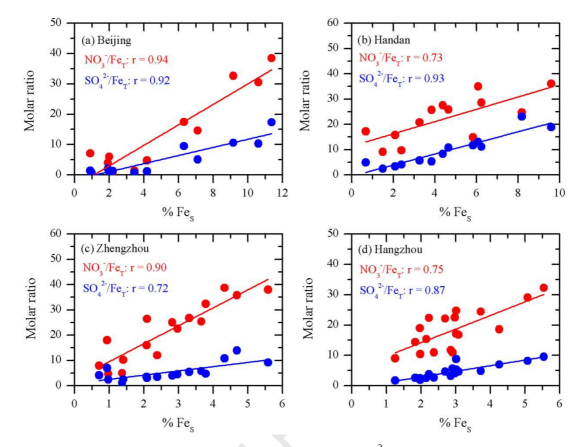


Fig. 5. Correlations between %Fe_s and NO₃⁻/Fe_T (red) and SO₄²⁻/Fe_T (blue) molar ratio at Beijing (a), Handan (b), Zhengzhou (c) and Hangzhou (d).

S

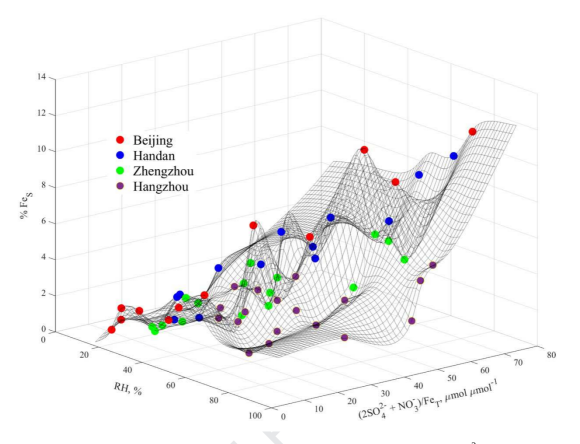


Fig. 6. Relationships of %Fe_s with RH and acidification degree molar ratio $(2SO_4^{2-} + NO_3^{-})/Fe_T$ in Beijing (red), Handan (blue), Zhengzhou (green) and Hangzhou (purple).

Highlights

1. Total iron and soluble iron concentrations as well as iron solubility in polluted air at four urban sites across East China were investigated.

2. A majority of nano-sized Fe-containing particles were internally mixed with sulfates and nitrates.

3. Chemical processing plays an important role in enhancing iron solubility in the polluted atmosphere.

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Declaration of interests

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