Air pollution–aerosol interactions produce more bioavailable iron for ocean ecosystems

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It has long been hypothesized that acids formed from anthropogenic pollutants and natural emissions dissolve iron (Fe) in airborne particles, enhancing the supply of bioavailable Fe to the oceans. However, field observations have yet to provide indisputable evidence to confirm this hypothesis. Single-particle chemical analysis for hundreds of individual atmospheric particles collected over the East China Sea shows that Fe-rich particles from coal combustion and steel industries were coated with thick layers of sulfate after 1 to 2 days of atmospheric residence. The Fe in aged particles was present as a “hotspot” of (insoluble) iron oxides and throughout the acidic sulfate coating in the form of (soluble) Fe sulfate, which increases with degree of aging (thickness of coating). This provides the “smoking gun” for acid iron dissolution, because iron sulfate was not detected in the freshly emitted particles and there is no other source or mechanism of iron sulfate formation in the atmosphere.

INTRODUCTION
Iron (Fe) is a micronutrient that limits primary productivity in large areas of the surface ocean, particularly in high-nutrient, low-chlorophyll areas (1). Fe may also limit nitrogen (N) fixation in low-latitude, N-limited oceans (2, 3). Soluble Fe from atmospheric deposition can stimulate primary production and/or nitrogen fixation in the surface ocean (1, 3–5). Changes in the soluble Fe input to the oceans could have an important impact on oceanic carbon uptake and storage and indirectly affect the climate (6).

Recent modeling studies have suggested that anthropogenic activities may have led to a doubling or even tripling of atmospheric soluble Fe deposition to the oceans since the Industrial Revolution (6–11). If confirmed, this increase in soluble Fe could have a major impact on ocean productivity, carbon uptake, ocean oxygen depletion and associated biogeochemical feedback, and climate (6, 11). A key component of these models is the hypothesized “Fe acid dissolution” process: Acids formed from anthropogenic gaseous pollutants such as sulfur dioxide dissolve iron in aerosol particles (12–14), making them bioavailable and increasing the bioavailable iron input to the oceans. Because of the potential importance of this process in the Fe cycle and ocean biogeochemistry, a number of field and laboratory studies have been carried out to test this hypothesis in the last decade. Laboratory studies found a positive relationship between Fe solubility (soluble Fe–to–total Fe ratio) and aerosol acidity (8, 14–16), providing indirect support to the hypothesis. However, field observations have been less conclusive (17–19). A key limitation is that previous studies have been based on bulk aerosol analysis and do not provide information on the distribution of soluble Fe in individual aerosol particles and how it relates to acidic compounds on a per-particle basis (20). Oakes et al. (21) showed, using bulk aerosol analysis upon ambient particle samples, that soluble Fe was correlated with sulfate in aerosol, a relationship consistent with low-pH environments. Longo et al. (22) recently suggested, through a combination of bulk measurements and some Fe mineral speciation with x-ray absorption near-edge structure, that strong acidity likely contributes to higher aerosol Fe solubility. Recently, Rindelaub et al. (23) demonstrated the potential of Raman microspectroscopy in measuring the pH in individual particles, but challenges in its application to atmospheric particles remain. The limitation of bulk analysis and the difficulty of measuring Fe species in individual aerosol particles (23, 24) make it highly challenging to test the Fe acid dissolution hypothesis.

RESULTS AND DISCUSSION
We used novel individual particle analysis techniques including nanoscale secondary ion mass spectrometry (NanoSIMS) and scanning transmission electron microscopy (STEM) to provide indisputable evidence of the Fe dissolution process from acids deposited on atmospheric particles. We collected a number of aerosol samples during a research cruise over the Yellow Sea in June 2013 (fig. S1A). Back trajectory analyses (fig. S2) indicated that air masses reaching the sampling sites were chiefly from mainland China. We investigated the composition and sources of Fe-bearing particles in the collected aerosol samples. The sizes of the particles were measured on the basis of projected area on microscopic photographs and then corrected to volume-based diameters (fig. S3). The chemical composition of 5511 particles with a size range of 20 to 5000 nm was analyzed using a transmission electron microscope (TEM) with an energy-dispersive x-ray spectrometer (EDS). Fe was detected in 14% (ranging from 5 to 29%) of all analyzed particles.

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We observed three main types of Fe-bearing particles: Fe-rich (Fig. 1A and fig. S4A), fly ash (Fig. 1B), and mineral dust. Fe-rich particles are distinct from coal fly ash particles: Fe in the former is the major element in their EDS spectra (for example, top EDS spectrum in Fig. 1), while that in the latter is a minor element (for example, bottom EDS spectrum in Fig. 1). Fly ash and Fe-rich particles were darker (more electron-dense) than secondary sulfate or organic matter (OM) under the TEM (Fig. 1, A and B). Fe-rich particles and fly ash usually displayed a spherical shape (Fig. 1, A and B), with the former mainly containing Fe and the latter containing Si, Al, and Fe (Fig. 1B). Mineral dust particles usually displayed an irregular shape and contained Si and Al with a small amount of Fe.

Fe-bearing particles were classified as externally mixed particles (that is, primary Fe-rich, fly ash, or mineral dust) and internally mixed particles (that is, primary particles mixed with sulfate and OM) (fig. S5). The contribution of externally mixed fly ash and Fe-rich particles to total Fe-bearing particles decreased from 64% for particles with a diameter of 20 to 200 nm, to 20% for particles with a diameter of 200 to 400 nm, to less than 5% for those with a diameter of >400 nm (Fig. 2). Overall, 75% of the Fe-bearing particles were internal mixtures of Fe-rich/fly ash and sulfate (Fig. 2).

STEM elemental maps show that Fe was distributed throughout the entire particle, in addition to the Fe hotspots (for example, see Fig. 3, A and B). The spherical morphology of Fe-rich or fly ash particles (Figs. 1 and 3A) suggests that they were formed via high-temperature processes followed by a fast cooling process (25). They were primarily emitted from anthropogenic emissions, such as coal combustion or industrial processes.

No sulfur was detected in Fe-rich particles collected from a steelworks or in coal fly ash particles collected from a power plant using TEM-EDS (fig. S6, A and B). However, most of the fly ash and Fe-rich particles collected over the Yellow Sea contained a significant amount of sulfate. More than 80% of the Fe-bearing particles larger than 200 nm were coated with sulfate (marked as “Sulfate with Fe inclusions” in Fig. 2), suggesting that the sulfate was formed in the atmosphere via gas-to-particle conversion. Almost none of the analyzed individual particles showed evidence of sodium, suggesting that the S is not associated with sea salt. Because most SO$_2$ in East Asia is emitted from power plants and industries (26, 27), and most of the sulfate aerosol is formed from anthropogenic SO$_2$ in the Northern Hemisphere (28), we argue that the sulfate aerosol found in the individual aerosol particles is primarily formed from anthropogenic SO$_2$.

The overall size of Fe-bearing particles ranges from 20 nm to 5 μm, with a peak at 700 nm; the size of Fe-rich/fly ash grains within these particles are much smaller, ranging from 20 nm to 1 μm, with a peak at 160 nm (fig. S7). On average, the size of the primary particles (that is, Fe-rich particles or fly ash) increased fourfold after 1 to 2 days of transport. This suggests that secondary aerosol formation on the Fe-rich/fly ash particles is very efficient, possibly involving trace metal or acid-catalyzed reactions in aqueous aerosol (29, 30).

STEM elemental maps showed an overlap of elemental Fe and S in most of the analyzed Fe-bearing particles, in addition to that in the Fe hotspots (for example, see Fig. 3, A and B). This suggests that the Fe dispersed in the secondary coating is released from the Fe-rich or fly ash inclusions, because there is no vapor-phase Fe in the atmosphere. In addition, if this Fe was present as insoluble Fe, such as Fe oxides or aluminosilicate, it would have only existed as individual grains, that is, nanoparticles. However, this is not the case, suggesting that the Fe in the secondary coating is likely to be present as soluble Fe, such as Fe sulfate.

We confirmed the presence of Fe sulfate in the sulfate coatings by NanoSIMS analysis of individual particles in the same samples. Figure 3C shows an example of ion intensity maps of CN$^-$, FeO$^-$, S$^-$, and FeS$^- in an individual particle: The particle has two FeO$^-$ hotspots, with FeS$^-$ distributed in the S$^-$ matrix. The FeO$^-$ hotspots are likely from Fe oxides or fly ash inclusions, similar to those shown in standard samples (fig. S8A), whereas the FeS$^-$ signal is from Fe sulfate (fig. S8B). Furthermore, the acid dissolution process reproduced in the laboratory showed a distribution pattern for FeS$^-$, FeO$^-$, and S$^-$ signals in NanoSIMS ion.
maps as well as for Fe and S elemental maps (figs. S8C and S9) that is similar to that of the real atmospheric particles (Fig. 3 and fig. S10). These results demonstrated that at least part of the Fe in the sulfate coating of ambient particles is present as Fe sulfate, which can only be formed from primary Fe particles (for example, fly ash or Fe-rich particles).

To further demonstrate the environmental relevance of the above-mentioned results, we carried out NanoSIMS measurements of ion intensity maps in a number of individual particles collected during the cruise. We then plotted the $\text{FeS}^-/($Fe$^3+ + \text{FeO}^-$) ratio against the $\text{S}^-/($Fe$^3+ + \text{FeO}^-$) ratio in Fig. 4. The former is an indicator of solubility, while the latter reflects the thickness of sulfate coating relative to total Fe. Figure 4 shows that Fe$^3+/(\text{FeO}^- + \text{FeS}^-)$ ratio increased with the $\text{S}^-/(\text{FeS}^- + \text{FeO}^-)$ ratio. Moreover, selected-area electron diffraction (SAED) confirmed that the sulfate coating in the randomly selected individual particles is more likely to be (acidic) ammonium bisulfate (fig. S4). This finding provides further support for higher solubility in individual particles associated with sulfate availability (in proportion to the total Fe present in the particles), an indicator of acidity in an ammonia-poor atmosphere (31), particularly at higher altitudes. Particles generated from a filtered water leach of an urban PM$_{2.5}$ (particulate matter with an aerodynamic diameter of less than 2.5 μm) sample collected during a haze event in the North China Plain produced a medium Fe$^3+/(\text{FeO}^- + \text{FeS}^-)$ ratio that is close to the regression line (Fig. 4). Because haze particles in China are predominantly secondary in origin (32), the acidic aqueous phase contained within the haze provides the necessary medium for acid dissolution.

A potential contradiction to the inferred acid dissolution process is that some of the Fe in the sulfate coating may be directly released from the soluble Fe on the surface of the primary Fe-rich or fly ash particles. However, this possibility is unlikely because no Fe was detected in the sulfate coating around coal fly ash generated from the suspension of coal fly ash in nonacidic ammonium sulfate solution (figs. S6C and S11; see experiment 3). This is consistent with the results of a previous study in which the initial solubility of Fe measured in coal fly ash is very low (much less than 1%) (16). Fe-rich particles from the iron/steel industry are also unlikely to be soluble in water because they are mainly
composed of Fe oxides (for example, magnetite in fig S4). Although Fe particles from ship and vehicle emissions or biomass burning may be highly soluble (for example, >50%) (21, 33), their contribution to all Fe-bearing particles is very small. In summary, the detection of Fe sulfate, mixed within the sulfate coatings, provides the “smoking gun” for acid dissolution—because there is no other atmospheric source of Fe sulfate or process that leads to its formation.

Our work also indirectly confirms the presence of the aerosol acidification process in the atmosphere. Although thermodynamic models, ion balance, or phase partitioning often predicts high acidity in wet aerosols (12, 13, 31, 34–36), aerosol acidity in individual particles has never been measured because of the technological challenges in measuring pH in the miniscule amounts of water associated with aerosol particles (23, 24). Our results confirm the existence of an acid dissolution process, providing indirect evidence that the aerosols become acidic after transport to the Yellow Sea. This is further supported by the SAED measurement, which showed that S is present as acidic ammonium bisulfate in the analyzed individual particles (fig S4). Recently, Wang et al. (37) argued that acids in aerosols in two Chinese megacities are completely neutralized by ammonium but we showed that aerosols in the Asian outflow is acidic. Future work needs to determine the spatial and temporal variability of aerosol acidity across size.

Current Fe emission inventories indicate that coal fly ash and dust globally contributed 0.45 and 7.8 Tg of total Fe per year deposited to the oceans (38). Ito and Shi (8) simulated the atmospheric acid Fe dissolution processes and calculated that the Fe solubility at deposition is ca. 8% for coal fly ash in the present day and ca. 0.27% for dust in the preindustrial era. Using these estimates, we calculated that soluble Fe deposition to the ocean from coal fly ash alone may be 50% more than that from natural dust. Acid dissolution of Fe in particles from the steel industry (38) and deserts (7–12, 35, 36) due to anthropogenic acidic gas emissions will certainly add more soluble Fe to the oceans. Thus, the Fe acid dissolution process associated with anthropogenic gas emissions may have significantly enhanced the soluble Fe deposition to the ocean. Mahowald et al. (6) estimated that even a doubling in soluble Fe deposition to the oceans could have resulted in a 6% increase in ocean productivity and an additional 8 Pg of carbon drawdown. Furthermore, Ito et al. (11) showed that the impact of increased soluble Fe supply to the oceans on productivity and carbon export may have contributed to the observed expansion of the oxygen depletion in many regions of the world with important impacts on ecosystems and biogeochemical cycles. It is also important to note that soluble Fe from anthropogenic aerosols (for example, coal fly ash) may have a disproportionately larger impact on marine ecosystems than dust—because they tend to deposit to the Fe-limited ecosystems (1, 2, 39) and their emissions are continuous throughout the year, in contrast to those of the dust (9, 31, 38–40). Looking into the future, changes in acidic gas emissions from natural or anthropogenic processes and atmospheric acidity will affect this Fe dissolution process and thus the soluble Fe to the oceans. Therefore, it is essential that Earth system models capture this process to better understand the interaction of air pollution with the atmospheric Fe cycle and ocean biogeochemistry (6, 11).

MATERIALS AND METHODS
Sample collection
Eighteen atmospheric particle samples were collected during a research cruise (Dongfanghong 2) over the Yellow Sea (fig. S1A) from 23 to 30 June 2013. The Yellow Sea is located between mainland China and the Korean Peninsula and is the northern part of the East China Sea, a marginal sea of the Pacific Ocean. All aerosol samples were collected onto copper TEM grids coated with carbon film (carbon type-B, 300-mesh copper; Tianld Co.) using a single-stage cascade impactor. Sampling durations varied from 10 to 15 min depending on particle loading to avoid particle aggregation during sample collection. Each sample was placed in a sealed dry plastic tube and stored in a desiccator at 25°C and 20 ± 3% relative humidity to minimize exposure to ambient air until further analysis. The red dots on the cruise track indicated where the samples were collected (fig. S1).

Aerosol optical depths (AODs) at 550 nm from the MODIS Terra sensor (data product: MODIS-Terra ver. 5.1) were obtained through the online Giovanni interface (41). The regional AOD distribution from 20 to 30 June 2013 suggests a gradient of high to low AOD from mainland China to the Yellow Sea (fig. S1B). Air mass back trajectory analysis indicates that most of the air masses reaching the cruise passed over the polluted coastal cities of East China (fig. S2).

Laboratory standard sample preparation
We have prepared a series of standards to verify our field results. Coal fly ash and Fe-rich particle samples were collected from the hoppers of electrostatic precipitators of a coal-fired power plant and a steelworks, respectively, in Linyi City, Shandong. Hematite, as an Fe oxide standard, was purchased from Sinopharm Chemical Reagent Co. Ltd. Four types of additional “standard” samples were prepared:

1. Hematite and sulfuric acid: 2.5 mg of hematite powder was added to diluted sulfuric acid to a final pH of 2.0. This experiment was carried out to simulate the acidic processing of Fe oxides, similar to previous studies (16–19, 34–36).
2. Hematite and sulfuric + oxalic acid: 15.7 mg of hematite and 22.5 mg of H2C2O4 powders (Sigma-Aldrich) were added to dilute sulfuric H2SO4 to a final pH value of 1.8. This experiment aimed to simulate the acid and oxalate processing of Fe oxides (8, 31).
3. Coal fly ash in a near-neutral ammonium sulfate solution: 5.0 mg of coal fly ash was added to (NH4)2SO4 solution (0.1 M) (pH 5.6). This experiment was used to test whether there is soluble Fe in untreated coal fly ash.
4. Haze particle leach: A PM2.5 sample was collected onto a Teflon filter during a haze event in Jinan (in the North China Plain); the sample was leached with 10 ml of Milli-Q water and then filtered using a membrane filter with a pore size of 0.2 μm.

Suspensions from Exps. 1 to 3 were shaken for 24 hours at room temperature before further processing. The water leach of the PM2.5 sample was prepared immediately before aerosol atomization. Aerosol particles were then generated from the particle suspensions (Exps. 1 to 3) and the filtrate (experiment 4) using an atomizer (Badger Airbrushes 150-7) and collected onto TEM grids.

Offline microscopic and spectroscopic analysis
Aerosol and standard particles collected on TEM grids were analyzed with a JEOL JEM-2100 TEM operated at 200 kV. To ensure that the analyzed particles were representative of the ambient sample, five areas were chosen from the center and periphery of the sampling spot on each grid. Every particle in the selected area was analyzed using EDS. EDS spectra were collected for 30 s to minimize radiation exposure and potential beam damage (52). Chemical composition of a total of 5511 individual particles ranging from 20 to 5000 nm was recorded. The area of an individual particle or the inclusion therein was
measured by using the iTEM system, on the basis of which the equivalent area diameter (EAD) was calculated (53). The SAED and fast Fourier diffraction patterns derived from crystal lattice spacings were used to confirm the phases of minerals in individual particles.

To correct the EAD to volume-based equivalent spherical diameters (ESDs), we obtained the two- and three-dimensional images of airborne particles using a Digital Nanoscope IIIa atomic force microscope (AFM) (54, 55). The instrument was operated in tapping mode with a cantilever and conical tip with a radius of 10 nm for imaging. Usually, particle images were taken at a full scan size of 10 µm, which included 10 to 15 particles. By comparing the ESD and EAD of the same particles, we observed a linear relationship between the two parameters (fig. S3), which was used to convert EAD measured under iTEM into ESD (fig. S7).

On the basis of the TEM analysis, we chose four typical aerosol samples for STEM analysis using a JEOL 2100F field-emission TEM equipped with an Oxford EDS system. STEM provides high-resolution elemental mapping (53, 56). These four samples were further analyzed by NanoSIMS 50L (Institute of Geology and Geophysics, Chinese Academy of Sciences). NanoSIMS can provide high-resolution chemical mapping (54, 57). The instrument was set to simultaneous secondary ion collection mode with pulse counting on electron multipliers, enabling us to simultaneously collect seven secondary ions originating from the same sputtered volume of the sample. A microerosion source was used to generate Cs+ primary ions, with an impact energy of 16 kV for sample interrogation. Fe2(SO4)3 powder was used to calibrate peaks in standard samples.

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