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Representation of iron aerosol size distributions of anthropogenic emissions is critical in evaluating atmospheric soluble iron input to the ocean

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Abstract. Atmospheric aerosol deposition acts as a major source of soluble (bioavailable) iron in open ocean regions where it limits phytoplankton growth and primary production. The aerosol size distribution of emitted iron particles, along with particle growth from mixing with other atmospheric components, is an important modulator of its long-range transport potential. There currently exists a large uncertainty in the particle size distribution of iron aerosol, and the extent to which such uncertainty shapes global soluble iron deposition remains unclear. Here, we couple a sophisticated microphysical, size-resolved aerosol model with an iron-speciated and iron-processing module to disentangle the impact of iron emission size distributions on soluble iron input to the ocean, with a focus on anthropogenic combustion and metal smelting sources. We evaluate our model results against a global-scale flight measurement dataset for anthropogenic iron concentration and show that the different representations of iron size distribution upon emission, as adopted in previous studies, introduce a variability in modeled iron concentrations over remote oceans of a factor of 10. Shifting the iron aerosol size distribution toward finer particle sizes ($< 1 \, \mu m$) enables a longer atmospheric lifetime (a doubling), promoting atmospheric processing, which enhances the soluble iron deposition to ocean basins by up to 50 % on an annual basis. The monthly enhancements reach 110 % and 80 % over the Southern Ocean and North Pacific Ocean, respectively. Uniquely, our results highlight that compared with emission flux variability, iron emission size distribution plays an equally important role in regulating soluble iron deposition, especially to the remote oceans. Our new findings can help to interpret inter-model differences in iron deposition estimation and to better quantify the effects of atmospheric nutrient input on marine biogeochemistry, including but not limited to iron and phosphorus.

1 Introduction

Iron is a critical micronutrient supporting marine primary production, which is closely associated with marine carbonnitrogen cycles in the Earth system (Mahowald et al., 2018; Moore et al., 2001). The atmospheric deposition of soluble iron to many ocean basins has long been regarded as an important source of bioavailable iron for ocean biota uptake in iron-limited areas (Jickells et al., 2005; Jickells and Moore, 2015; Tagliabue et al., 2017). Understanding the amount and past-to-future evolution of atmospheric iron deposition to the ocean is critical in assessing ocean carbon sequestration under a changing climate (Bergas-Massó et al., 2023; Myriokefalitakis et al., 2020; Hamilton et al., 2020b).

The quantification of soluble (bioavailable) iron input to the ocean is linked to differences in iron emission source properties, the degree to which iron aerosol undergoes acidic or organic chemistry, and atmospheric transport (Hamilton et al., 2022). Atmospheric iron comes from three major emission sources, i.e., wind-blowing dust, wildfire and biomass burning, and anthropogenic activities (such as fossil fuel combustion and iron smelting). Dust storms, which frequently occur in arid or semi-arid regions of the world, such as North Africa and East Asia, provide an abundant iron source to the ocean and support primary production (Mahowald et al., 2009; Westberry et al., 2023). In addition, a growing body of evidence is showing that pyrogenic iron, with higher fractional solubility than dust (Ito et al., 2019), is a large source of atmospheric soluble iron deposition to many ocean basins, including the Southern Ocean, North Pacific Ocean, and North Atlantic Ocean (Conway et al., 2019; Liu et al., 2022; Matsui et al., 2018; Seo and Kim, 2023). Because the strength of each source could be affected by future climate change and/or human activities, their contributions to bioavailable iron input to the ocean may vary regionally and temporally by the end of the century (Bergas-Massó et al., 2023).

Atmospheric transport provides an essential pathway through which iron aerosol emitted from the land is supplied to the remote ocean. Atmospheric circulation patterns dictate the main transport pathways for aerosol to follow and thereby which source regions are important to consider in terms of their supply to ocean basins. Additionally, atmospheric transport enables internal mixing of iron-bearing aerosols with other aerosol and gas components, like sulfates and organics - a process commonly known as aging - which facilitates the dissolution of iron from an insoluble state to a soluble state (Shi et al., 2020; Shi et al., 2012; Solmon et al., 2009; Li et al., 2017). The atmospheric aging processes can significantly increase iron solubility and subsequent soluble iron deposition, evidenced by both in-field and laboratory observations and global model simulations (Ito, 2015; Li et al., 2017; Longo et al., 2016). Uncovering the underlying mechanisms of the aging processes and associated enhancement of iron solubility during transport is an ongoing topic of investigation (Meskhidze et al., 2019; Shi et al., 2020).

To elucidate the atmospheric flux of iron-containing aerosols to the ocean, global-scale aerosol models have been developed to include a range of iron emission sources that currently show a large intermodal difference in flux estimates (Myriokefalitakis et al., 2018). Part of the problem is that it is difficult to realistically reproduce the distribution of soluble iron concentrations across all the different regions of the world, especially over the remote polar oceans, which are often characterized by low iron concentrations with a high fractional solubility (Ito et al., 2019). Among those global aerosol simulations, the size distribution of iron, which is an important consideration when determining aerosol lifetimes and thus its long-range transport potential, is key in shaping atmospheric iron distributions (Hamilton et al., 2020a; Myriokefalitakis et al., 2018). Compared to coarse-sized particles (e.g., larger than 1 µm), smaller particles generally feature lower loss rates with respect to dry deposition and wet removal, resulting in longer atmospheric lifetimes; being transported over a longer distance increases the potential for atmospheric processing (i.e., a longer period of aerosol aging) and thus more soluble iron deposition. Evidence from recent works (Zhang et al., 2023, 2022) has shown that iron sources and physicochemical properties largely differ between coarse and fine particles, leading to different iron solubility. Representation of iron size distribution in models could therefore be important.

Iron aerosol characteristics depend in part on differences between source types. The iron mass size distribution associated with natural dust sources commonly pertains to mineral dust aerosols, with the coarse-sized (diameter greater than 1 µm) fraction being dominant (Mahowald et al., 2014; Albani et al., 2014). Similarly, fire iron emissions are dominated (> 80%) by coarse-mode particles (Hamilton et al., 2019), suggested to be due to the entrainment of local dust iron-bearing aerosol in the pyro-convective updrafts generated by a fire (Hamilton et al., 2022). For iron aerosol with an anthropogenic source, however, the relative fractions between the fine and coarse particle size distribution upon emission are more divergent among previous investigations. Recent observational constraints have revealed large mass concentrations of anthropogenic iron oxide in the fine mode (Moteki et al., 2017), while a subset of modeling studies have treated most of this iron in the coarse mode (Wang et al., 2015). As opposed to these natural sources, anthropogenic iron size distributions may be highly variable with respect to diverse fuels, combustion temperatures, and industrial processes, as well as the abatement technologies applied to control air pollution (Rathod et al., 2020; Hamilton et al., 2020a).

The extent to which iron aerosol size distributions shape the pattern of atmospheric soluble iron inputs to different ocean regions is currently unknown. Herein, by leveraging a size-resolved global aerosol model configured with iron processes, we focus on the representation of anthropogenic iron size distributions upon emission, primarily involving their roles in altering global long-range transport and deposition fluxes of iron. We further place the effect of iron size distribution in the context of iron emission uncertainty to shed light on its relative importance in controlling global-scale iron deposition.

2 Methods and materials

2.1 Global aerosol model

We conducted global aerosol simulations using the Community Atmosphere Model version 5 (CAM5.3) with the Aerosol Two-dimensional bin module for foRmation and Aging Simulation (CAM-ATRAS) version 2 (Matsui, 2017; Matsui and Mahowald, 2017). The model treats a series of aerosol chemical and microphysical processes in a size-resolved manner with 12 aerosol size bins from 1 to 10 000 nm in diameter. Our recent study suggests that this size-revolved method can represent the growth of small particles to larger ones and the evolution of particle size distributions during atmospheric transport well (Liu and Matsui, 2022). Both dry deposition (Zhang et al., 2001) and wet deposition (Liu et al., 2012) of aerosols are treated in our model. The improvement of the aerosol in-cloud wet scavenging process was included to improve the modeling of aerosol long-range transport efficiency (Liu and Matsui, 2021). The CAM-ATRAS model has been adequately validated for aerosol mass and number concentrations at a global scale using comprehensive measurements from the ground to the upper troposphere (Matsui and Liu, 2021; Gliß et al., 2021; Kawai et al., 2021; Matsui et al., 2022; Matsui and Moteki, 2020).

To represent the iron cycle from emission to deposition, we explicitly treated iron constituents within the aerosol model, similar to our previous study (Liu et al., 2022), with updates to iron processing in the current work. All ironbearing components were assumed to be internally mixed with other aerosols and underwent emission, physical transport, chemical aging (e.g., solubilization), and deposition in the atmosphere. The internal mixing assumption is reasonable for anthropogenic iron aerosols, which are often mixed together with other aerosol compounds in polluted environments, e.g., East Asia, and enables the growth of iron aerosols via condensation and coagulation. Our model simulated iron solubility and atmospheric processing of ironbearing aerosols through online coupling with the Mechanism of Intermediate complexity for Modelling Iron (Hamilton et al., 2019). For anthropogenic iron, we consider five different minerals, namely magnetite, hematite, illite, kaolinite, and sulfate iron, following the global emission inventory by Rathod et al. (2020), which was developed by a bottomup approach at a 1° spatial resolution and 1-month temporal resolution. The reference year of the inventory was 2010. We account for a wide range of anthropogenic sources, including iron smelting and fossil fuel combustion sources.

Dust iron emission was calculated by assuming a constant iron content of 3.5% in dust aerosol emission (Shi et al., 2012). The model estimated total dust emission fluxes using the scheme of Zender et al. (2003), with modifications by Albani et al. (2014) and the size distribution from Kok (2011). Dust mineralogy was not involved in our simulations. The global interannual mean iron (insoluble + soluble) emissions from dust, biomass burning, and anthropogenic sources were 87, 1.1, and 2.2 Tg Fe yr⁻¹, respectively.

We validated our modeled anthropogenic iron oxide concentrations against a global-scale aircraft measurement in the troposphere consisting of eight campaigns for the periods of 2009–2011 and 2016–2018 (Lamb et al., 2021). These observations provide mass concentrations of anthropogenic iron oxide, i.e., magnetite, with volume equivalent diameters between 180–1290 nm. The model results were extracted along the flight tracks in time and space and further averaged in several latitudinal bands across the Pacific and Atlantic oceans (see Fig. S1 in the Supplement). More details can be found in Liu et al. (2022).

The model was compiled with a horizontal resolution of $1.9^{\circ} \times 2.5^{\circ}$ (latitude × longitude) and 30 vertical layers from the surface to 40 km. We ran the model for the two periods, 2008–2011 and 2015–2018, with the first year in each period functioning as spin-up. The meteorological fields were nudged by the Modern-Era Retrospective analysis for Research and Applications version 2. In addition, to provide implications for ocean biogeochemistry, we estimated the changes in marine net primary production induced by iron inputs following the methods used by Rathod et al. (2022) and Okin et al. (2011), in which a cutoff (4 µmol L⁻¹) of nitrate concentrations at the surface water was chosen to define the geographical areas of iron-limited ocean basins.

2.2 Representation of iron size distribution of anthropogenic emissions

The number and mass size distribution of aerosol upon emission is an essential parameter in aerosol modeling. However, due to the limited knowledge about iron emission characteristics, the representation of iron size distributions varies greatly for anthropogenic sources. We tested four different size distributions of anthropogenic iron upon emission (Fig. 1), briefly described as follows.

First, we adopted the same size distribution as our previous studies (Matsui et al., 2018; Liu et al., 2022) based on Moteki et al. (2017), abbreviated as Moteki2017 hereafter, which was derived from a power law function to fit the observed anthropogenic iron oxide concentrations within the boundary layer in the outflow of East Asian sources. Note that the observed size-revolved number concentrations were confined within 170 to 2100 nm in diameter given the detection limit. By extrapolating the observation results, we obtained a mass

Figure 1. Mass size distribution functions for anthropogenic iron emission adopted in four previous studies. The Moteki2017 curve is provided by fitting in situ measurements for iron particles within the range of 170 to 2100 nm in diameter over East Asia; Rathod2020 is based on the latest combustion iron emission inventory, with updates on iron estimates in fine-mode sizes ($< 1 \mu m$); and the Hamilton2019 and Wang2015 cases are modeling studies that assume most anthropogenic combustion in the coarse-mode bin.

size distribution between 1 and 10 000 nm with negative values excluded. In this case, more than 90% of iron mass was allocated to the size range of 100 to 2500 nm. In addition, we varied the emissions by a factor of 2 ($\times 0.5$ and $\times 2.0$, respectively) in two other parallel experiments to account for the potential uncertainties in iron emission estimates.

The second case was derived from Rathod et al. (2020) (abbreviated as Rathod2020 hereafter). They developed a new mineralogy-based iron emission inventory by introducing more details in anthropogenic sources, especially the inclusion of a metal smelting source. These improvements increase the fine-aerosol (less than 1 μ m in diameter) fractions by a factor of 10 compared to most previous inventories. Consequently, this inventory was characterized by almost-equal fractions between fine- and coarse-sized emissions, while previous inventories always applied a much larger fraction for the coarse mode. Consistent with Rathod et al. (2020), we allocated 10 % and 90 % of fine iron mass to the Aitken mode and the accumulation modes of aerosols, respectively.

The third case was derived from Hamilton et al. (2019) (abbreviated as Hamilton2019 hereafter). They revised the anthropogenic iron emission inventory based on Luo et al. (2008) (no metal smelting) and showed that the ratio of fine-sized iron mass to coarse-sized iron mass was 1 : 5.6, which resulted in the coarse mode dominating. A similar ratio for anthropogenic iron emissions was applied by Ito (2013). Moreover, 10% of fine-sized emissions were allocated to the Aitken mode and the remaining 90% to the accumulation mode.

The fourth case was derived from Wang et al. (2015) (abbreviated as Wang2015 hereafter). In their combustion iron inventory, the ratio of fine-mode mass to coarse-mode mass was as low as 1:24 because only 0.1 %–0.3 % of iron mass

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from coal fly ash was emitted in the fine mode. Thus, we allocate 96 % of iron in the coarse mode and the remaining 4 % in the fine mode.

To enable intercomparison among these cases, we used the global-scale anthropogenic iron emission mass inventory from Rathod et al. (2020) but with different allocations between fine and coarse sizes in each case. The size distribution of anthropogenic iron emission in each case was treated uniformly on a global scale. Therefore, the simulated variability in atmospheric iron input to the ocean between cases should be attributable to iron size distributions rather than emission amount. For the last three cases, we adopted three constant lognormal modes to distribute iron emissions, namely Aitken, accumulation, and coarse modes, with their determining parameters including geometric standard deviations, number mode diameter, and density reported by Hamilton et al. (2019). We then separated these three modes into 12 size bins from 1 to 10000 nm, adapted for our size-resolved aerosol modeling. The size distributions of iron from biomass burning and dust sources were consistent with Liu et al. (2022) in all cases. In the following analysis, we grouped Moteki2017 and Rathod2020 into the fine-sized group, while Hamilton2019 and Wang2015 were grouped into the coarse-sized group.

3 Results and discussion

3.1 Atmospheric iron aerosol concentrations

We first examined the effect of changes to iron particle size distributions for anthropogenic sources (unless otherwise stated) on the atmospheric iron aerosol burden and its global distributions. Figure 2 illustrates that the global-mean anthropogenic iron lifetimes differ by a factor of approximately 2 among the four examined cases. Both the Moteki2017 and the Rathod2020 cases simulated a lifetime around 3.0 d. In contrast, the Hamilon2019 and Wang2015 cases simulated a lifetime around half as long (between 1.4 and 1.7 d). As only the size distribution is changed in these simulations, the change in lifetime is directly linked to the apportionment of mass aerosol between fine- and coarse-particle size modes. This result is in line with previous reports (Hamilton et al., 2020a) and demonstrates that shifting emitted iron toward fine sized diminishes atmospheric loss rates of iron aerosols via dry (sedimentation) and wet (precipitation) removal pathways and extends their lifetime. Consequently, given the same emission amount, the atmospheric iron burdens are enhanced accordingly by a factor of approximately 2 from $\sim 9.0 \,\text{Gg}$ in the coarse-sized cases (Hamilton2019 and Wang2015) to \sim 18.0 Gg in the fine-sized cases (Moteki2017 and Rathod2020). In a similar manner, the lifetimes and mass burdens of anthropogenic iron in the soluble form are almost doubled in the fine-sized cases (Fig. 2b). The extended lifetimes also enhance the globally averaged iron solubility



Figure 2. Global anthropogenic iron concentration burdens and lifetimes varied by the emission size distributions. The scatter plots are shown for (**a**) anthropogenic iron (labeled AN iron) burden vs. lifetime and (**b**) soluble anthropogenic iron vs. lifetime. Four representative cases are examined in this work: Moteki2017, Rathod2020, Hamilton2019, and Wang2015. The first two cases are grouped into fine sized, and the other two are grouped into coarse sized.

(Fig. S2) by allowing more iron to be subject to aerosol aging and solubilization processes.

To evaluate each iron simulation, we compare simulated aerosol characteristics against global-scale aircraft measurements of anthropogenic magnetite within nine regions of the troposphere (Fig. 3). We extracted the modeled mass concentrations of iron aerosols with a size range similar to that of the measurements. Note that magnetite, comprising about 70 % of anthropogenic iron emissions, can be used as an indicator of anthropogenic iron abundance in the atmosphere (Rathod et al., 2020; Matsui et al., 2018). Despite the fact that the same emission fluxes are considered in all cases, their simulated magnetite aerosol concentrations can differ by up to a factor of 10. Specifically, the Moteki2017 and Rathod2020 cases show a much better performance in reproducing the observed profiles over all ocean basins compared to the Hamilon2019 and Wang2015 cases. In particular, the cases with a more uniformly distributed particle size distribution across modes capture the high concentrations (> 1 ng m⁻³) in the North Pacific, which can be linked to atmospheric plumes transported from East Asia with intensive emission rates (Seo and Kim, 2023; Moteki et al., 2017). Some underestimations still exist near the surface or at high altitudes. Doubling the emission fluxes based on the Moteki2017 case can appreciably narrow the differences from the observations (dashed red lines in Fig. 3). In contrast, the coarse-sized-dominated simulations (i.e., Hamilton2019 and Wang2015) underrepresent the magnetite concentrations over global remote oceans, particularly by up to 1 order of magnitude over the Pacific Ocean. The shorter lifetimes in this group limit the long-range transport of iron aerosols from continental sources to the remote atmosphere.

These results imply that agreement between observations and model simulations can be improved by reducing uncer-



Figure 3. Comparison of modeled anthropogenic iron mineral (magnetite) vertical concentration profiles in four cases with aircraft measurements across global oceans. The flight routes and model–observation sampling methods have been documented in Lamb et al. (2021) and Liu et al. (2022). The geographical location of each oceanic area is marked in Fig. S1. We also up- and downscaled emission fluxes by a factor of 2 from the Moteki2017 case to account for potential uncertainties in emission estimates (dashed red lines in the panels).

tainties in the emission inventory and in the long-range transport efficiency associated with the representation of iron size distributions. Moreover, as illustrated in Fig. 3, the variabilities (the ratio between the maximum and minimum) of the simulated magnetite vertical profiles due to iron size distribution changes are wider than those due to the emission uncertainties, for which emission fluxes were perturbed by a factor of 2 ($\times 2.0$ and $\times 0.5$, respectively) to test the sensitivity of the simulated iron concentrations. We therefore highlight that in order to observationally constrain iron emissions more realistically in global aerosol simulations, it is a prerequisite to use a realistic empirical representation of anthropogenic iron aerosol size distributions.

Our simulations further demonstrate that the representation of iron size distribution shapes the iron aerosol concentrations at a global scale (Fig. 4). Anthropogenic iron in the fine-sized group shows column concentrations more than twice as high as those in the coarse-sized group over



Figure 4. Global map of variability in yearly averaged iron column concentrations between the fine-sized group and the coarse-sized group. The variability ratios (fine sized/coarse sized) are calculated for (**a**) anthropogenic iron, (**b**) total iron, and (**c**) soluble iron. Herein, total iron is a combination of iron from dust, biomass burning, and anthropogenic sources, the soluble form of which denotes soluble iron. The ratios represent the maximum differences between the fine-sized group (the Moteki2017 and Rathod2020 cases) and the coarse-sized group (the Hamilton2019 and Wang2015 cases) and indicate the spread of iron simulations.

the oceans. The differences (ratios) are larger in these remote oceans compared to source regions like East Asia, southern Africa, and South America because iron-bearing aerosols of smaller sizes can be transported over a longer distance. The variability between the two groups is relatively minor for total iron (Fig. 4b), which includes contributions from dust and biomass burning sources. By contrast, the differences for soluble iron are more pronounced than for total iron over much of the global ocean (Fig. 4c) because of the higher solubility of anthropogenic iron compared to dust iron. Over East Asia and its outflow areas, the rapid aging processes in polluted environments are capable of enhancing iron solubility, particularly that of anthropogenic origin, and thus amplifying the differences in soluble iron concentrations between the fine-sized and coarse-sized groups (Li et al., 2017; Zhu et al., 2022).

3.2 Atmospheric soluble iron input to the ocean

Next, we examine the extent to which iron size distributions upon emission can alter soluble iron input to the global ocean basins, which is vital for net primary productivity, especially in high-nutrient, low-chlorophyll (HNLC) areas (Hamilton et al., 2022; Moore et al., 2013). As illustrated in Fig. 5, although the global emission and resulting annual iron (insoluble + soluble) deposition amounts are the same between the cases, their geographical distributions vary substantially. Using the ratio of annual soluble iron deposition in the finesized group to that in the coarse-sized group as a proxy of the variability, we find the fine-sized distributions lead to much more soluble iron input to remote ocean basins, including the North Pacific and Southern Ocean, by up to a factor of 4 for the anthropogenic sources and 1.5 for the total of all sources (i.e., anthropogenic + fire + dust). Similar spatial patterns emerge regarding the total iron deposition, suggesting the importance of long-range transport efficiency in regulating iron distributions (Fig. S3). The variability is negligible in the equatorial and subtropical Atlantic, where dust iron dominates soluble iron input to this area. Ratios of less than 1, indicating reduced deposition fluxes in the fine-sized group, are found near the continental sources, including the western US, Australia, and southern Africa, because of the slower deposition speed. However, in East Asia, which has the most intensive anthropogenic iron emissions, the shift toward finer sizes also increases soluble iron deposition near the sources (e.g., eastern China). This is attributable to the rapid aging processes of the fine-sized iron in such a polluted environment, which convert more insoluble iron to its soluble form (Baldo et al., 2022; Zhang et al., 2022).

The source appointment of soluble iron deposition across ocean basins also varies with iron size distributions (Fig. 5cd). The anthropogenic iron source becomes more dominant in the North Pacific, North Atlantic, and parts of the Southern Ocean, with its fractional contribution reaching more than 50 % in the fine-sized group compared to that of 30 %-40 % in the coarse-sized group. Such variability is attributable to the enhancement of anthropogenic soluble iron fluxes to these remote oceans due to the shift in anthropogenic iron emissions toward finer-sized bins. Globally, the soluble iron deposition from anthropogenic sources is $55.0 \,\mathrm{Gg} \,\mathrm{yr}^{-1}$ for the fine-sized group, which is larger than that of $35.3 \,\mathrm{Gg} \,\mathrm{yr}^{-1}$ for the coarse-sized group. Hence, even though the same emission is applied in these simulations, the diversity of iron size distributions upon emission yields considerable variability of soluble iron deposition on a global basis. As discussed earlier, the extended iron lifetime by a factor of approximately 2 in the fine-sized group allows for more iron to be transported to a remote region and simultaneously increases the amount of atmospheric iron processing and dissolution to a soluble form. Of the total iron deposition, the soluble fraction is thus notably elevated. We also find that the chemical aging process, as the major source of soluble iron, controls the differences in soluble iron deposition over remote oceans between the fine-sized and coarse-sized groups (Fig. S4).



Figure 5. Comparison of yearly accumulated soluble iron input to the ocean between the fine-sized and coarse-sized cases. Shown here are (**a**) the ratio of anthropogenic soluble iron simulated in the fine-sized case to the result of the coarse-sized case, (**b**) the ratio of total soluble iron, (**c**) the fractional contribution (in percentage) of anthropogenic emissions to total soluble iron input in the fine-sized case, and (**d**) the fractional contribution of anthropogenic emissions in the coarse-sized case.

In this study, the explicit treatment of anthropogenic iron mineralogy enables us to identify iron-mineral-dependent variability. We find that for minerals coming primarily from fossil fuel combustion and iron smelting on land, namely magnetite, hematite, and clays, the shift in iron emissions toward finer-sized bins promotes their long-range transport and enhances corresponding soluble iron deposition to the North Pacific, equatorial Pacific, and Southern Ocean by more than a factor of 4 (Fig. 6a-c). However, the size distribution effects on iron sulfates are pronounced only in the polar areas, which are subject to plumes of middle-latitude shipping emissions. In line with a previous study (Rathod et al., 2022), iron sulfates constitute an important contribution to soluble iron deposition away from major continental sources, predominately associated with shipping emissions over local oceans rather than long-range transport from land (Fig. S5). Hence, the variability induced by iron size distributions is less remarkable for iron sulfates than for other anthropogenic minerals. These results also suggest that the relative importance of iron sulfates in total soluble iron deposition to remote oceans is altered by the size distributions of all other iron minerals that originate from continental sources. Because the size distributions of anthropogenic iron minerals may depend on different combustion processes, source- and region-specific size distribution representation is desirable in future work.

It is critical to examine monthly soluble iron availability altered by emission size distributions because ocean primary production can respond to iron inputs of the order of days to months (Guieu et al., 2014). From our results in Fig. 7, the importance of anthropogenic emission size distributions in shifting soluble iron deposition varies by month over potentially iron-limited ocean basins, i.e., HNLC regions, due to the episodic nature of natural iron sources (dust and wildfire) and their deposition. For the Southern Ocean, the monthly ratios of the fine-sized case to those of the coarse-sized case span 1.1 in September to 2.1 in December (Fig. 7a). The September peak in fire iron (shown in Fig. S6), possibly linked to low precipitation in the southern winter, masks the variability in anthropogenic iron contributions due to emission size distributions. Conversely, the largest difference in December is associated with the lowest contribution of nat-



Figure 6. Ratios of yearly accumulated soluble iron deposition for four anthropogenic iron-containing minerals, comparing fine-sized and coarse-sized cases.



Figure 7. Differences in monthly total soluble iron deposition between the fine-sized and coarse-sized groups over specific ocean basins. Only anthropogenic iron emission sizes have been examined here. Histograms describe the ratio of monthly results for the fine-sized case to those for the coarse-sized case for the (a) Southern Ocean, (b) North Pacific, and (c) equatorial Pacific. For comparison, the black dots represent the ratio of the fine-sized results to those with global anthropogenic emission amounts downscaled by a factor of 2. The ocean basins of interest are indicated in the top-left corner of each panel. The dashed red lines indicate the ratio of 1.0.

ural sources (Fig. S6). By contrast, the monthly differences are less variable in the North Pacific, ranging between 1.3 and 1.8 (Fig. 7b). Anthropogenic emission dominates soluble iron throughout the year, except in March–May, during which dust storms originating from East Asia occur frequently and regulate soluble iron inputs to the North Pa-

cific. The equatorial Pacific has the lowest ratio amongst the three regions because anthropogenic aerosol-bearing plumes rarely arrive in this region, and large amounts of rain here can efficiently remove aerosols. For the three ocean basins, such differences related to the emission size treatments are even larger than those caused by adjusting the iron emission



Figure 8. Difference in marine net primary production sustained by atmospheric soluble iron between the two iron size distribution groups. The panels display (**a**) absolute differences and (**b**) percentage differences in net primary production of the fine-sized group relative to the coarse-sized group. Here, following Rathod et al. (2022), we focus only on the iron-limited ocean basins, which are defined using a cutoff of nitrate concentrations at surface oceans.

amount by a factor of 2 considering emission uncertainty (black dots in Fig. 7). We therefore suggest that compared with iron emission fluxes, the representation of size distributions for anthropogenic iron is equally or even more important for the estimation of total soluble iron deposition to remote oceans.

We also provide an estimate of the changes in ironsustained marine net primary production between the finersized and coarse-sized groups (Fig. 8). In line with the distributions of soluble iron deposition, the effects of finer-sized iron distributions can enhance primary production over remote oceans, including the North Pacific Ocean and Southern Ocean, by as much as 50%. Considering that anthropogenic iron aerosols may contribute to > 10% of the total marine productivity in the North Pacific Ocean (Rathod et al., 2022; Ito et al., 2020), the representation of their size distributions upon emissions, mostly from East Asia, is particularly important in Earth system modeling. The evolution of atmospheric iron aerosol size characteristics and their emission fluxes can be critical to ocean carbon sequestration from the past to the future. Hamilton et al. (2020a) found that historical air pollution control has cut down anthropogenic emission amounts of coarse-sized particles, in turn elevating the mass fraction of finer-sized iron particles and thus the overall lifetime of atmospheric iron. Hence, the complex interactions between iron and the Earth system are linked to the effects of human activity on soluble iron availability to ocean basins.

4 Conclusion

This study explores the extent to which iron size distribution upon emission, specifically from anthropogenic sources, alters estimates of soluble iron deposition to the open ocean. A global microphysical, size-resolved aerosol model is used to simulate the iron cycle, involving emissions, atmospheric processing, and deposition on a global scale. The model treats iron mineralogy, size evolution, and chemical aging processes during atmospheric transport, which enables the investigation of the relationship between iron size distributions and iron long-range transport and subsequent deposition. We test four representative size distribution schemes for anthropogenic iron sources employed in previous studies.

We find that allocating a more balanced fraction of iron aerosol upon emission to particle sizes less than 1 µm results in a longer atmospheric lifetime and larger mass burden of total iron aerosols by a factor of approximately 2 compared to a coarse-sized-dominated case, primarily associated with decreased loss rates via dry and wet removal processes. The evaluation of anthropogenic iron aerosols against the globalscale observation dataset reveals that despite the fact that the same emission fluxes are considered in all cases, their simulated magnetite aerosol concentrations differ by up to a factor of 10, while the higher fine-sized cases agree better with the observations. It is therefore necessary to accurately represent iron size distributions in order to constrain iron emission fluxes more realistically with aerosol simulations and observations (Liu et al., 2022). Our simulations show that the resulting annual soluble iron deposition differs by up to a factor of 1.5 over remote oceans, including the North Pacific Ocean and Southern Ocean, because the fine-sized group allows more iron to be transported over a long distance with enhanced atmospheric processing. More importantly, the monthly soluble iron deposition, which is relevant to ocean primary production responses over days to months, would be enhanced by 110 % and 80 % in the finesized case over the Southern Ocean and North Pacific Ocean, respectively. Such differences are similar to or even larger than those considering emission uncertainty, suggesting an equally important role of iron size distribution treatment.

This study unravels the critical role of iron size distributions in shaping atmospheric soluble iron inputs to global oceans, especially to remote regions. However, a realistic understanding of iron emission size distributions is still inadequate given limited observation data. Targeted on-site measurements of iron aerosol size along with its mass and solubility at source areas are highly desirable. Higher-resolution models with finer grids and detailed microphysics are useful for exploring iron aerosol deposition and chemical aging processes at regional scales. Furthermore, our finding may be extended to other key trace elements that are of importance to ocean biogeochemistry, like copper, manganese, and phosphorus.

Code availability. The CAM model is publicly available from NCAR (Neale et al., 2010). The source code for the atmospheric iron dissolution scheme is available at https://doi.org/10.5194/gmd-12-3835-2019 (Hamilton et al., 2019).

Data availability. The model data used to generate the figures can be made available upon request.

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