Mixing of Asian mineral dust with anthropogenic pollutants over East Asia: a model case study of a super-duststorm in March 2010

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Abstract. Mixing of Asian mineral dust with anthropogenic pollutants allows pollutants (e.g. sulfate and nitrate) to be transported over longer distances (e.g. to the northern Pacific, even to North America) along with dust particles. This mixing therefore affects the atmospheric and oceanic environment at local, regional and even continental scales. In this study, we used a three-dimensional regional chemical transport model (Nested Air Quality Predicting Modeling System, NAQPMS) to examine the degree of mixing between Asian mineral dust and anthropogenic pollutants in a super-duststorm event during 19–22 March 2010. Influences of the mixing processes on regional atmospheric environmental and oceanic biogeochemical cycles were also investigated. A comparison with measurements showed that the model reproduced well the trajectory of long-range dust transport, the vertical dust profile, and the chemical evolution of dust particles. We found that along-path mixing processes during the long-range transport of Asian dust led to increasingly polluted particles. As a result, ∼60 % of the sulfate and 70–95 % of the nitrate in the downwind regions was derived from active mixing processes of minerals with pollutants sourced from the North China Plain and enhanced by transport over South China. This mixing had a significant impact on the regional-scale atmospheric composition and oceanic biogeochemical cycle. Surface HNO₃, SO₂ and O₃ were decreased by up to 90 %, 40 % and 30 %, respectively, due to the heterogeneous reactions on dust particles. Fe solubility rose from ∼0.5 % in the Gobi region to ∼3–5 % in the northwestern Pacific, resulting from oxidization of SO₂ on dust particles. Total Fe(II) deposition in the ocean region of East Asia reached 327 tons during the 4-day dust event, and created a calculated primary productivity of ∼520 mgC m⁻² d⁻¹ in the Kuril Islands, which can support almost 100 % of the observed mean marine primary productivity in spring in this region (526 mgC m⁻² d⁻¹).

1 Introduction

Over the last few decades, China has experienced urbanization and industrialization processes on an unprecedented scale. More than 120 cities have populations of more than one million. This rapid growth in such a short period of time has not only led to a remarkable increase in material wealth and a higher standard of living, but has also caused serious so-called “complicated air pollution”, which means various types of pollution (e.g. acid rain, coal smog, photochemical smog, duststorm and haze) are present together in China (He et al., 2002). The chemical transformation and transport processes in this “complicated air pollution” make emissions controls more difficult. In particular, the interaction between gas and aerosols creates major challenges for policy makers. Therefore, approaches to reduce “complicated air pollution” and its impact on other regions have already become the research focus for Chinese environmental scientists.

The mixing of Asian mineral dust with anthropogenic pollutants is one of the major research issues. This is because extremely high levels of sulfur dioxide (SO₂) and nitrogen dioxide (NO₂) in China allow dust particles to take up high levels of both these pollutants, leading to the formation of
water-soluble sulfate and nitrate (Jordan et al., 2003). Compared with pure secondary anthropogenic sulfate and nitrate, sulfate and nitrate on dust can be transported over much longer distances (e.g. to the northern Pacific, even to North America) (Duce, 1980; Wilkening et al., 2000) along with dust particles. These pollutants alter the photochemistry, acid deposition and oceanic primary productivity along the pathway of long-range transport, through chemical transformation and dry/wet deposition (Jordan et al., 2003; Zhuang et al., 2003; Tang et al., 2004). Consequently, studying the mixing of Asian mineral dust with anthropogenic pollutants is crucial for a detailed understanding of the impact of China’s anthropogenic emissions on the atmospheric and oceanic environments at local, regional and even continental scales.

The mineralogy of Asian dust particles consists primarily of carbonates (CaCO$_3$), quartz (SiO$_2$), feldspars, micas, clays and hematite (Fe$_2$O$_3$). Ca in Asian dust accounts for 39 % of seven crustal elements (Si, Al, Mg, Ca, Na and K), in contrast with Saharan dust particles where Ca only takes up 17 % (Kreuger et al., 2004). The greater Ca concentration allows Asian dust particles to easily react with anthropogenic sulfuric and nitric acid, which consequently chemically alter the dust particles (Jordan et al., 2003). By collecting and analyzing aerosol samples over long time scales at North Pacific island stations, Zhuang et al. (1992) proposed a hypothesis for the coupling and feedback between natural iron (Fe) and anthropogenic S in the atmosphere and ocean. In their hypothesis, the heterogeneous oxidation of SO$_2$ to sulfate drives the acidification of dust particles. In highly acidic ambient aerosol solutions, Fe (III) in atmospheric dust particles is reduced to Fe (II) with OH radical production. More Fe(II) could result in the marine production of more DMS which in turn provides more SO$_2$ into the atmosphere. The more SO$_2$ result in more acidic SO$_4^{2-}$ aerosol on mineral dust and again more Fe(II). Charlson et al. (1987) suggested that the heterogeneous oxidation of SO$_2$ to sulfate is a key step in this feedback between iron and sulfate. Furthermore, Mesekhidze et al. (2003) proposed that the SO$_2$ from urban and industrial centers plays a key role in Fe mobilization in Asian mineral dust. Zhuang et al. (1992) believed that this hypothesis can explain high Fe(II) and non-sea salt sulfate (nss-sulfate) observed in aerosol samples at a remote pacific station (Midway, Hawaii). Zhuang et al. (1992) believed that this hypothesis can explain high Fe(II) and non-sea salt sulfate (nss-sulfate) observed in aerosol samples at a remote pacific station (Midway, Hawaii).

Partial evidence for Zhuang’s hypothesis has been found in observations of Asian dust plume outflows in the Yellow Sea (East of China) and northwestern Pacific (downwind region of China) since the late 1990s. Choi et al. (2001) reported that SO$_4^{2-}$, NO$_3^-$, Ca$^{2+}$ and Mg$^{2+}$ were the dominant water soluble ions in bulk dust aerosols in Seoul, and SO$_4^{2-}$ and NO$_3^-$ were primarily found in the coarse fraction associated with Ca$^{2+}$ (Kim and Park, 2001). This is in contrast with non-dust observations suggesting that anthropogenic SO$_4^{2-}$ and NO$_3^-$ are associated with NH$_4^+$ in fine particles. Lidar observations showed that the aerosols were possibly an internal mixture of dust and sulfate during dusty days in the northeast Pacific (Sugimoto et al., 2002). Jordan et al. (2003) estimated that 43 % of sulfate came from Asian dust particles during an intensive aircraft-based aerosol sampling program as part of TRACE-P. Aerosol samples on the NASA DC-8 aircraft also revealed that the capture of SO$_4^{2-}$ and NO$_3^-$ by Asian mineral particles was accompanied by acid-mobilization of Fe (Mesekhidze et al., 2003). This acid-mobilization can increase Fe solubility from 0.5 % to 1–2 % within 3–5 days (Mesekhidze et al., 2003). Excellent correlation between the marine primary productivity observed at two remote sites in the northwestern Pacific and dust storm frequency in China seems to support their conclusion (Deng et al., 2008).

Numerical modeling is a very useful tool for estimating the degree of mixing of minerals with anthropogenic pollutants (Tie et al., 2005). Various models have been used to estimate the contribution of China’s anthropogenic emissions to this mixing process by comparing model results and observations in regions downwind of China’s emissions (e.g. Korea, Japan and the northwestern Pacific). Lasserre et al. (2008) developed a model for assessing this mixing. Using a regional model, Song and Carmichael (2001) estimated that chemical conversion of SO$_2$ on the surface of dust particles accounted for 10–40 % of SO$_4^{2-}$ production in Asian dust plumes. Tang et al. (2004) found heterogeneous reactions on dust particles can produce > 20 % of SO$_4^{2-}$ and > 70 % of NO$_3^-$ under heavy dust loadings over the northwestern Pacific. Luo et al. (2005) and Fan et al. (2006) also suggested that increasing SO$_2$ emissions alone could have caused significant Fe mobilization in the Northern Hemisphere oceans. However, the lack of model evaluation in China (the major region where minerals mix with pollutants) leads to large uncertainties in their estimations. A recent dust model intercomparison (DMIP) study over Asia found that simulated dust emissions by 8 models ranged from 27 to 336 Tg in a Asian dust event, although they agreed with each other in downwind regions, and suggested that modeling of dust transport and removal processes between China and Japan was one of the most important issues in improving dust modeling (Uno et al., 2006). In addition, greatly varied uptake coefficients are another origin of uncertainties (e.g. uptakes of nitric acid (HNO$_3$) varied greatly in the studies of Tang et al. (2004) and Tie et al. (2005), with values of 0.01 and 0.1, respectively). Therefore, a simulation evaluated by observations in China is urgently needed for assessing the extent to which mineral dust mixes with China’s anthropogenic emissions.

Recent studies have extended observations into China. Sun et al. (2005) found SO$_4^{2-}$/Al and NO$_3^-$/Al ratios in dust particles in Beijing (Fig. 1) were 60 and 8 times higher than those in surface soils of the Gobi Desert and Loess Plateau, respectively. They concluded that these high ratios in Beijing were
attributed to the capture of \( \text{SO}_2 \) or \( \text{HNO}_3 \) by mineral dust during long-range transport (Sun et al., 2005). Meanwhile, Fe (II) in total Fe greatly increased, from 0.4 % to 1.3–5.3 %, during the long-range dust transport in Beijing (Zhang et al., 2005). By applying positive matrix factorization (PMF) combined with the enrichment factors method, Yuan et al. (2008) suggested that dust particles brought Beijing a lot of sulfate, but little nitrate, which is opposite to other studies (Jordan et al., 2003). However, high concentrations of both sulfate and nitrate were observed in Xiamen, China (Fig. 1) during dusty days (Zhao et al., 2011). More evidence of the mixing of minerals with pollutants in China can be found in the literature (Zhang et al., 2003; Zhuang et al., 2003, 2010; Wang et al., 2011a). The above observations provide excellent opportunities for model simulations to estimate the extent to which dust mixes with China’s anthropogenic emissions.

On 19–23 March 2010, a super-duststorm event occurred in China. In contrast to previous events, which have all occurred at mid-latitudes, this event covered the whole of eastern China from the Gobi desert (45° N) to South China Sea. Even at Xiamen city (24.5° N) in South China and on Dongsha Island (20° N) in the South China Sea, aerosol concentrations reached 990 \( \mu \text{g m}^{-3} \) and 300 \( \mu \text{g m}^{-3} \), respectively, which are the highest levels in history (Wang et al., 2011b; Zhao et al., 2011). Remote sensing showed that this event lasted in eastern China for 4 days (Li et al., 2011b), which allowed dust particles to mix thoroughly with China’s anthropogenic emissions.

The goal of this study is to estimate the extent to which Asian aerosols mixed with China’s pollutants in the dust episode of 19–23 March 2010. We utilized a regional chemical transport model (NAQPMS) and evaluated its performance by comparison with in-situ and remote sensing observations in China. We also assessed changes of gas concentrations by comparison with in-situ and remote sensing observations in China. We also assessed changes of gas concentrations by comparison with in-situ and remote sensing observations in China. We also assessed changes of gas concentrations by comparison with in-situ and remote sensing observations in China.
study, \( b_{\text{ext550}} \) (expressed as inverse megameters, \( 10^{-6} \text{ m}^{-1} \), at 550 nm), was calculated by Eq. (1):

\[
b_{\text{ext550}} = 3.0 \times f(\text{RH}) \{ [(\text{NH}_4)_2\text{SO}_4] + [\text{NH}_4\text{NO}_3] \} + 4.0 \times [\text{OMC}] + 10.0 \times [\text{LAC}] + 1.0 \times [\text{SOIL}] + 0.6 \times [\text{CM}] \tag{1}
\]

where \( f(\text{RH}) \) represents the hygroscopic growth factor or extinction enhancement factor resulting from water uptake on aerosols. OMC, LAC, SOIL and CM represent concentrations (\( \mu g \text{ m}^{-3} \)) of organic matter, light-absorbing carbon, soil aerosol and coarse mass, respectively.

### 2.2 Description of a unique dust deflation module

The generation of desert dust is critical in this study. Here, we applied a modified size-segregated dust deflation module designed by Wang et al. (2000) to simulate the long-range transport of Asian dust. The model was developed after detailed analysis of the meteorological conditions, landforms and climate available in daily weather reports from about 300 local weather stations in northern China (Wang et al., 2000). It has been successfully applied in the simulations of dust long-range transport, impact of dust on air quality and dust-climate interactions (Wang et al., 2002; Yue et al., 2010).

Recently, Luo and Wang (2006) modified the mineral dust emission intensity (\( F \)), considering soil categories, vegetation fraction percentages and snow/ice coverage, by introducing the following equation:

\[
F = C_1 \times \frac{\rho_d}{g} \times E \times \left(1 + \frac{u_0^3}{u^3}\right) \times \left(1 - \frac{u_0^2}{u^2}\right) \times \left(1 - \frac{\text{RH}}{\text{RH}_0}\right) \tag{2}
\]

where \( F \) is the dust flux (\( \text{kg m}^{-2} \text{ s}^{-1} \)). The constant \( C_1 \) is set to \( 1.0 \times 10^{-5} \). \( \rho_d \) (\( \text{kg m}^{-3} \)) and \( g \) (\( \text{m} \text{s}^{-2} \)) are the air density and acceleration due to gravity. Weighting factor (\( E \)) represents the uplifting capability of land surface, and reflects the impact of landuse categories, vegetation fractions and snow/ice coverage on dust fluxes. In the desert without vegetable and snow cover, dust particles are easily uplifted to the boundary layer, and \( E \) is set to 1.0. On the contrary, \( E \) is set to 0.0 in evergreen forest. In this study, soil categories, vegetation fractions and snow/ice coverage are derived from Moderate Resolution Imaging Spectrometer (MODIS) data. Figure 1 shows the distribution of calculated \( E \) in this study. \( u^3 \) and \( u_0^3 \) are the fraction and threshold friction velocities. \( u_0^3 \) is related to soil type, mineral particle size distribution, surface roughness and soil moisture. Here, \( u_0^3 \) is set to 0.45, 0.35 and 0.6 m s\(^{-1} \) in the Gobi, China Loess and Hunshandak deserts, respectively. These values were obtained from observations by Zhu and Zhang (2010). \( u_0^3 \) in other dust source regions is set as 0.4 m s\(^{-1} \) as suggested by Yue et al. (2010). RH and \( \text{RH}_0 \) represent relative humidity and its threshold value. In this study, \( \text{RH}_0 \) is set to 40%. The dust particle size is divided into 4 size bins covering the range 0.43–10 \( \mu \text{m} \) in diameter (0.43–1 µm; 1–2.5 µm; 2.5–5 µm and 5–10 µm).

### 2.3 Uptakes of anthropogenic gases on aerosol

In this study, 28 heterogeneous reactions are considered when simulating the mixing of aerosols with pollutant gases. Table 1 lists the all first-order rate constant (\( k_{\text{het}} \)) is calculated by uptake coefficients (\( \gamma \)) and surface area density of particles (\( A \)) using the following equation suggested by Jacob (2000):

\[
k_{\text{het}} = \left( \frac{r}{D_g} + \frac{4}{cr} \right)^{-1} \times A \tag{3}
\]

where \( r \) is the black carbon and sulfate mean radius, \( D_g \) is the gas phase diffusion coefficient, \( c = \left( \frac{8 k T}{m} \right)^{0.5} \) is the mean molecular speed of the gas, \( \gamma \) is the uptake coefficient and \( A \) is the surface area density of the particles.

Depending on types of dust sample, experimental and analytical methods, measured uptake coefficient (\( \gamma \)) on certain reaction usually varies greatly, in some cases several orders of magnitude. This could bring uncertainties to assess the mixing extents of dust and urban aerosols. Considering impact of dust mineralogy, we tried our best to select those values on China losses and Gobi desert which are potential sources to reduce the uncertainties due to \( \gamma \). For example, the reaction \( \text{HNO}_3 \) on dust particles in this study is experimental \( \gamma \) on China Losses at 0%, 40% and 80% RH (Wei, 2010).

In this study, we compute uptakes of gases on mineral dust separately for each dust bin (4 bins from 0.43–10 \( \mu \text{m} \)) which is expected to reduce errors caused by coarse size distributions in previous studies, and maintain dust sulfate and dust nitrate as separate size-segregated components in the model. These dust-nitrate and dust-sulfate coat mineral dust particles by a shell on the surface of dust particles, and experience the same transport and deposition processes as the corresponding dust particles.

### 2.4 Acid dissolution of iron containing mineral dust

Mixing of urban acidic gases with mineral dust might result in an enhancement of iron mobilization in mineral dust plumes (Mesekhidze et al., 2003). To the extent that this mobilized iron is then acting as a limiting marine primary productivity. This would imply that iron mobilization is one of important indicators reflecting impacts of mixing processes on regional environment. Air pollution controls in China which reduced \( \text{SO}_2 \) and \( \text{NO}_x \) emissions might actually depress C uptake in pacific waters, and thus affect the regional climate.

A large number of studies have demonstrated that dissolved iron (Fe(II)) in surface seawater is the limiting nutrient factor for primary productivity in high-nitrate low-chlorophyll (HNLC) regions of ocean (Zhuang et al., 2003). However, Fe in surface soils of the Gobi deserts is found...
Table 1. Heterogeneous reactions and reactive uptake coefficients.

<table>
<thead>
<tr>
<th>No.</th>
<th>Heterogeneous reactions</th>
<th>( \gamma^a )</th>
<th>Ref(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HR1</td>
<td>( \text{N}_2\text{O}_5 + \text{sulfate} \rightarrow 2\text{HNO}_3 )</td>
<td>( \gamma = \alpha \times 10^\beta )</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>( \alpha = 2.79 \times 10^{-4} + 1.3 \times 10^{-4} \times \text{RH} )</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(-3.43 \times 10^{-6} \times \text{RH}^2 + 7.52 \times 10^{-8} \times \text{RH}^3 )</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>( \beta = 4 \times 10^{-2} \times (T - 294)(T \geq 282 \text{K}) )</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>( \beta = -0.48(T &lt; 282 \text{K}) )</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HR2</td>
<td>( \text{NO}_3 + \text{sulfate} \rightarrow \text{HNO}_3 )</td>
<td>( 3 \times 10^{-3} )</td>
<td>2</td>
</tr>
<tr>
<td>HR3</td>
<td>( \text{HO}_2 + \text{sulfate} \rightarrow 0.5\text{H}_2\text{O}_2 )</td>
<td>( 2.5 \times 10^{-1} )</td>
<td>3</td>
</tr>
<tr>
<td>HR4</td>
<td>( \text{OH} + \text{sulfate} \rightarrow \text{products} )</td>
<td>( 2 \times 10^{-1} )</td>
<td>4</td>
</tr>
<tr>
<td>HR5</td>
<td>( \text{HCHO} + \text{sulfate} \rightarrow \text{products} )</td>
<td>( 2.2 \times 10^{-2} )</td>
<td>5</td>
</tr>
<tr>
<td>HR6</td>
<td>( \text{O}_3 + \text{soot} \rightarrow \text{products} )</td>
<td>( 1.8 \times 10^{-4} \times e^{-\frac{T}{1000}} )</td>
<td>5</td>
</tr>
<tr>
<td>HR7</td>
<td>( \text{NO}_2 + \text{soot} \rightarrow \text{ONO} )</td>
<td>( 3.3 \times 10^{-4} )</td>
<td>6</td>
</tr>
<tr>
<td>HR8</td>
<td>( \text{HNO}_3 + \text{soot} \rightarrow \text{NO}_2 )</td>
<td>( 2.1 \times 10^{-2} )</td>
<td>6</td>
</tr>
<tr>
<td>HR9</td>
<td>( \text{NO}_2 + \text{soot} \rightarrow 0.5\text{HONO} + 0.5\text{HNO}_3 )</td>
<td>( 3.0 \times 10^{-3} )</td>
<td>7</td>
</tr>
<tr>
<td>HR10</td>
<td>( \text{N}_2\text{O}_5 + \text{soot} \rightarrow 2\text{HNO}_3 )</td>
<td>( 5.0 \times 10^{-3} )</td>
<td>1</td>
</tr>
<tr>
<td>HR11</td>
<td>( \text{O}_3 + \text{dust} \rightarrow \text{products} )</td>
<td>( 2.7 \times 10^{-5} )</td>
<td>8</td>
</tr>
<tr>
<td>HR12</td>
<td>( \text{HNO}_3 + \text{dust} \rightarrow \text{NO}_3^- )</td>
<td>( \gamma = c \times \text{RH} \times \frac{1}{(1-\text{RH}) \times (1-\text{RH}^{0.018}(c = 8))} )</td>
<td>9,10c</td>
</tr>
<tr>
<td>HR13</td>
<td>( \text{NO}_2 + \text{dust} \rightarrow 0.5\text{HONO} + 0.5\text{HNO}_3 )</td>
<td>( 2.1 \times 10^{-6} )</td>
<td>8</td>
</tr>
<tr>
<td>HR14</td>
<td>( \text{NO}_3 + \text{dust} \rightarrow \text{HNO}_3 )</td>
<td>( 1.0 \times 10^{-3} )</td>
<td>11</td>
</tr>
<tr>
<td>HR15</td>
<td>( \text{N}_2\text{O}_5 + \text{dust} \rightarrow 2\text{HNO}_3 )</td>
<td>( 3.0 \times 10^{-2} )</td>
<td>8</td>
</tr>
<tr>
<td>HR16</td>
<td>( \text{OH} + \text{dust} \rightarrow \text{products} )</td>
<td>( 1.0 \times 10^{-1} )</td>
<td>8</td>
</tr>
<tr>
<td>HR17</td>
<td>( \text{H}_2\text{O}_2 + \text{dust} \rightarrow 0.5\text{H}_2\text{O}_2 )</td>
<td>( 2.0 \times 10^{-1} )</td>
<td>8</td>
</tr>
<tr>
<td>HR18</td>
<td>( \text{H}_2\text{O}_3 + \text{dust} \rightarrow \text{products} )</td>
<td>( \gamma = 12 \times \text{RH}^2 - 5.95 \times \text{RH} + 4.08 )</td>
<td>12</td>
</tr>
<tr>
<td>HR19</td>
<td>( \text{SO}_2 + \text{dust} \rightarrow \text{SO}_4^{2-} )</td>
<td>( 1.0 \times 10^{-4} )</td>
<td>13</td>
</tr>
<tr>
<td>HR20</td>
<td>( \text{CH}_3\text{COOH} + \text{dust} \rightarrow \text{products} )</td>
<td>( 1.0 \times 10^{-3} )</td>
<td>8</td>
</tr>
<tr>
<td>HR21</td>
<td>( \text{CH}_3\text{OH} + \text{dust} \rightarrow \text{products} )</td>
<td>( 1.0 \times 10^{-5} )</td>
<td>8</td>
</tr>
<tr>
<td>HR22</td>
<td>( \text{HCHO} + \text{dust} \rightarrow \text{products} )</td>
<td>( 1.0 \times 10^{-5} )</td>
<td>8</td>
</tr>
<tr>
<td>HR23</td>
<td>( \text{N}_2\text{O}_5 + \text{SSA} \rightarrow 2\text{HNO}_3 )</td>
<td>( 5.0 \times 10^{-3}(\text{RH} &lt; 62 %) )</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>( 3.0 \times 10^{-2}(\text{RH} &lt; 62 %) )</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HR24</td>
<td>( \text{NO}_3 + \text{SSA} \rightarrow \text{HNO}_3 )</td>
<td>( 1.0 \times 10^{-3} )</td>
<td>7</td>
</tr>
<tr>
<td>HR25</td>
<td>( \text{HO}_2 + \text{SSA} \rightarrow 0.5\text{H}_2\text{O}_2 )</td>
<td>( 2.0 \times 10^{-1} )</td>
<td>7</td>
</tr>
<tr>
<td>HR26</td>
<td>( \text{SO}_2 + \text{SSA} \rightarrow \text{SO}_4^{2-} )</td>
<td>( 5.0 \times 10^{-3}(\text{RH} &lt; 50 %) )</td>
<td>14</td>
</tr>
<tr>
<td></td>
<td>( 5.0 \times 10^{-2}(\text{RH} &lt; 50 %) )</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HR27</td>
<td>( \text{NO}_3 + \text{SSA} \rightarrow \text{NO}_3^- )</td>
<td>( 1.7 \times 10^{-2} )</td>
<td>15</td>
</tr>
<tr>
<td>HR28</td>
<td>( \text{HNO}_3 + \text{SSA} \rightarrow \text{NO}_3^- )</td>
<td>( 5.0 \times 10^{-1} )</td>
<td>16</td>
</tr>
</tbody>
</table>

\( ^a \gamma \) is the reaction probability; \( T \) is temperature (K), \( \text{RH} \) is relative humidity (%).


\( ^c \) Vlasenko et al. in reference 98 suggested the BET isotherm has a similar shape with the experimental \( \gamma \) as a function of \( \text{RH} \). Wei et al. in reference 108 reported \( \gamma \) on China Losses at 0 %, 40 % and 80 % \( \text{RH} \). So \( \gamma \) in this study is calculated by the BET isotherm forced by \( \gamma \) values reported by Wei (2010).

mainly in the form of undissolved hematite (\( \alpha \)-Fe2O3). Acid mobilization which reduce undissolved hematite to dissolved Fe(II) is thought as one of possible mechanisms (Mesekhidze et al., 2003). Fan et al. (2006) proposed a two-step mechanism for the acid-mobilization of iron in mineral dust particles. In this mechanism, formation of dissolved Fe (Fe(II)) experiences two processes: sulfate coating and dissolution of iron (hydr)oxide on the coated dust aerosols. The sulfate coating process can be achieved by the heterogeneous oxidation of SO2 and HNO3 (HR12 and HR19 in Table 1), and represents the acidification of mineral dust surface. Dissolution of iron (hydr)oxide represents the mass transfer from insoluble Fe to dissolved Fe in acid highly acidic ambient aerosol solutions. Three types iron/dust are included: fresh, coated and dissolved (for Fe). In this study, we used this two-step mechanism to reproduce the impact of mixing processes on Fe mobilization.
In the first step, gas uptakes convert dust particles from fresh to coated by the heterogeneous reactions HR12 and HR19. The rate coefficient represents how many fresh dust particles are converted into coated particles and is calculated from $k_x \times [SO_4^{2-}] + k_n \times [HNO_3]$, where $k_x$ and $k_n$ represent the first order rates of HR19 and HR12. In the second step, Fe(III) in the coated dust is transferred to dissolved Fe (Fe(II)). The production rate of Fe(II) is calculated from $R_{Fe} = R_d \times n A M / w$, where $R_{Fe}$ is grams of Fe dissolved per gram of Fe(III) per second, $R_d$ is the dissolution rate per unit surface area, $A$ is the specific surface area of hematite ($m^2 g^{-1}$), $n = 2$ (moles Fe/moles hematite), $M$ is the gram-molecular weight of Fe, and $w$ is the mass fraction of Fe in hematite (i.e. 0.7). Here, $R_d$ and $A$ are set as $1 \times 10^{-10} \text{mol m}^{-2} \text{s}^{-1}$ and $100 \text{m}^2 \text{g}^{-1}$ as suggested by Fan et al. (2006) and Duckworth and Martin (2001). These values correspond to an e-folding time of 6–10 days for dissolution. Besides the heterogeneous chemistry, the effects of radiation process and cloud process on Fe-mobilization are also considered based on the scheme devised by Luo et al. (2005).

Here the Fe solubility is assumed to have a lower limit of 0.5% (Fan et al., 2006).

Overall, five categories of aerosols are explicitly treated in our model: (1) anthropogenic aerosols (sodium-chloride-sulfate-nitrate-ammonium, black carbon and organic carbon) (2): dust particles; (3) sea salt, (4) dust-sulfate and dust-nitrate, (5) sea salt-nitrate and sulfate. The aerosol and gas gases are coupled through (1) gas chemistry module, (2) aqueous chemistry module, (3) heterogeneous chemistry module, (4) gas-aerosol partitioning of inorganic compounds. In the category of anthropogenic aerosols, sulfate, nitrate and ammonium are in the form of NaCl, Na$_2$SO$_4$, NaNO$_3$, NH$_4$NO$_3$, (NH$_4$)$_2$SO$_4$, NH$_4$Cl, NH$_4$HSO$_4$, NaHSO$_4$, and (NH$_4$)$_3$H(SO$_4$)$_2$ and internally mixed each other. They are in fine mode with a lognormal size distribution. Anthropogenic aerosols are externally mixed with the other categories of aerosols (dust and sea salt). Previous studies shows that the assumption of external mixture of anthropogenic aerosols with dust or sea salt is reasonable (Maxwell-Meier et al., 2004; Tang et al., 2004; Song et al., 2005).

### 2.5 Model setup

Figure 1 shows the NAQPMS model domain used in this study, which is composed of two nested domains. The coarser domain is $7760 \times 6160$ km on a Lambert conformal map projection with 80-km grid resolution. The nested domain is divided into $148 \times 160$ horizontal grids with 20-km resolution. Vertically, the model uses 20 terrain-following layers from the surface to 20 km a.s.l., with the lowest 10 layers below 3 km.

The first meteorological guess fields and boundary conditions for the coarse domain for every 6 h are obtained from NCAR/NCEP FNL reanalysis data ($1^\circ \times 1^\circ$). Nudging, or four dimensional data assimilation (FDDA) (OTTE, 2008), consisting of 6-h 3-D analyses of temperature, water vapor mixing ratio and horizontal wind components for both domains, and 6-h surface analyses of horizontal wind components for the coarse domain, were used with nudging coefficients of $3.0 \times 10^{-4}$ for the wind, temperatures and humidity fields of the coarse and nested domains. In this study, The Yonsei University (YSU) Boundary Layer (BL) scheme is used for calculating BL height. The surface layer and land surface schemes are the MM5 similarity and Noah Land Surface schemes, respectively.

The anthropogenic emissions of air pollutants (fossil fuels, biofuels and industrial emissions) in Asia are derived from the year 2006 bottom-up Regional Emission inventory in Asia (REAS) data at $(0.5^\circ \times 0.5^\circ)$ resolution (Ohara et al., 2007). The initial and boundary conditions are taken from a global chemical transport model MOZART-V2.4 with $2.8^\circ$ resolution. The mass fraction of Fe(III) in dust particles is set as 2.8% in this study, following Zheng et al. (1994).

The simulation started at 00:00 UTC on 10 March 2010 and ran through to the 24 March 2010. The first 5 days are regarded as the spin-up period, used to reduce the influence of initial conditions.

### 3 Model validation

Before validating concentrations of simulated dust particles and pollutants, we compare the predicted and observed hourly averaged meteorological variables (temperature, relative humidity and wind) during 19–23 March 2010 (figure not shown). The results show the model successfully reproduced the synoptic features of the meteorological variables, notably the change of wind direction from northward to southward during the dust episode.

#### 3.1 Observational data

In this study, a combination of aspaceborne instruments, backscatter lidars, and mass aerosol instruments is used to evaluate the model’s ability in simulating horizontal and vertical distributions of aerosols over East Asia during the dust event.

The Moderate Resolution Imaging Spectroradiometer (MODIS) field of AOD at 550 nm is obtained from the MODIS/AQUA Deep Blue Collection 005. MODIS is one of the most widely used passive satellite instruments in aerosol remote sensing. In this study, the daily level 3 AOD product at 550 nm (Ver5.1) was retrieved as an average value for an area defined by a $1^\circ \times 1^\circ$ square cell. The expected accuracy level over land is $\pm 0.05 \pm 0.15 \times$ AOD (Remer et al., 2005). Previous validations suggest that MODIS is capable of deriving AOD over land and ocean within expected uncertainty (Song et al., 2001; Cheng et al., 2012).
Table 2. Statistical summary of comparisons of the model results with observations.a.

<table>
<thead>
<tr>
<th></th>
<th>AOD550</th>
<th>Modis</th>
<th>5701</th>
<th>0.60</th>
<th>0.65</th>
<th>0.64</th>
<th>−0.05</th>
<th>0.52</th>
<th>56.0</th>
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<tr>
<td>SO₂</td>
<td>Shanghai</td>
<td>211</td>
<td>8.3</td>
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<td></td>
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<tr>
<td></td>
<td>Beijing</td>
<td>231</td>
<td>15.3</td>
<td>0.58</td>
<td>4.2</td>
<td>9.36</td>
<td>57.3</td>
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<tr>
<td>NO₂</td>
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<td>0.49</td>
<td>−22.0</td>
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<td>62.0</td>
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<tr>
<td></td>
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<td>21.0</td>
<td>0.67</td>
<td>−0.2</td>
<td>9.93</td>
<td>42.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PM₁₀</td>
<td>Beijing</td>
<td>9</td>
<td>217.5</td>
<td>0.91</td>
<td>−19.1</td>
<td>174.9</td>
<td>43.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jinan</td>
<td>9</td>
<td>213.2</td>
<td>0.82</td>
<td>52.6</td>
<td>203.0</td>
<td>47.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Shanghai</td>
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<td>0.94</td>
<td>−44.5</td>
<td>89.4</td>
<td>42.6</td>
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<tr>
<td></td>
<td>Taipei</td>
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<td>64.8</td>
<td>0.80</td>
<td>−5.6</td>
<td>38.5</td>
<td>52.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Xiamen</td>
<td>9</td>
<td>116.9</td>
<td>0.91</td>
<td>−37.8</td>
<td>110.1</td>
<td>67.6</td>
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<tr>
<td></td>
<td>Nanchang</td>
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<td>0.86</td>
<td>−34.1</td>
<td>129.7</td>
<td>73.5</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a N is the number of observed samples, 〈Cₘ, Cₒ, r, MB, RMSE and NME represent the mean modeled and observed values, correlation coefficient, mean bias, root mean square error and normalized mean error. Units of SO₂, NO₂ and PM₁₀ are ppbv, ppbv and µg m⁻³, respectively. AOD550 is unitless.

b Observed PM₁₀ is daily mean values, except Taipei which is an hourly mean value.

The aerosol backscatter data at Nagasaki, Chiba and Hedo shown in Fig. 1 is taken from the National Institute of Environmental Studies (NIES) lidar network (http://www-lidar.nies.go.jp/). In this ground-based lidar network over Asia, Mie-lidar extinction coefficients of non-spherical Asian dust are measured by the two-wavelength Mie-scattering lidar which has three channels and measures the backscattering at 1064 and 532 nm and the depolarization at 532 nm. Vertical profiles averaged for 5 min are recorded every 15 min. The height resolution was 6 m and the profiles are recorded up to 6 km.

Surface mass aerosol concentrations are determined by the Chinese air pollution index (API) which is officially reported for the principal pollutant among SO₂, NO₂ and PM₁₀ (particles on the order of ~10 µm or less) (http://www.sepa.gov.cn/english/air_s.php3). API is calculated from daily averaged concentration (from 12:00 the day before to 12:00 on this day). During the study period, PM₁₀ were always the prevailing pollutant in observation stations shown in Fig. 1, which allowed us to validate simulated PM₁₀ mass concentrations. In this study, we use a polynomial regression to derive the PM₁₀ mass loading from Chinese API, which has been already validated in the study of Lasserre et al. (2008). Note that API has an upper limit of 500, which means this value of 600 µg m⁻³ is a “saturation” maximum value.

\[
PM₁₀(\mu g m⁻³) = 8 \times 10⁻⁶(API)^³ - 87 \times 10⁻³(API)^² + 36706(API) - 12127.
\]

3.2 Comparison with satellite retrievals

Figure 2 presents the aerosol optical depth (AOD550) at 550 nm retrieved by MODIS (available at http://gdata1.sci.gsfc.nasa.gov/daac-bin/G3/gui.cgi?instance_id=MODIS_DAILY_L3) and the AOD550 simu-

![Fig. 2. MODIS-derived (left panel) and simulated AOD550 (right panel) during 19–23 March 2010. The white color in left panel indicates cloud contaminations.](http://example.com/image.jpg)
lated by NAQPMS during the 19–23 March 2010. Table 2 shows a series of statistical measures of agreements or disagreements between the simulation and observation. In general, the simulation reproduced the long-range transport of dust particles and anthropogenic aerosols over East Asia reasonably well over the study period. The correlation coefficients between model and satellite reach 0.64. On 19 March, a duststorm episode started over the Gobi desert, as shown in Fig. 2e. This was associated with the formation of a strong Siberian-Mongolian high pressure system and a much stronger westerly wind in the Gobi, which entrained desert sand into the atmosphere (figure not shown). Unfortunately, MODIS data are missing over the Gobi due to either the lack of interorbital coverage or cloud contamination (Fig. 2a). However, the aerosol index measured by OMI (Ozone Monitoring Instrument) captured this outbreak of dust deflation over the Gobi (Li et al., 2011b). On 20 March, both the model and MODIS revealed that dust particles moved southeasterly to the North China Plain, the Gulf of Bohai and the Korean peninsula (Fig. 2b, f), where AOD550 was generally greater than 2. On 21 March, the dust plume and cold front advanced further, reaching southern China (including Xia’men City) and Taiwan. Over the ocean, a narrow plume extended from Taiwan to the southwest of Japan. Due to relatively calm winds associated with the southeastward-moving high, dust particles still accumulated in central China and the Yangtze River delta (Fig. 2c, g). On 23 March, MODIS showed a relatively high AOD550 belt in the mid-latitudes, which we attribute to the combination of weakened dust and anthropogenic particle (e.g. BC, sulfate and nitrate) concentrations. This pattern is well reproduced by the simulation, which indicates good model performance with respect to both dust and anthropogenic particles.

3.3 Comparison with ground-based observations

Simulated concentrations of PM\textsubscript{10} (particles on the order of \(\sim 10\mu m\) or less) at six stations covering most of east China (20–40\(^\circ\)N) are compared with observations in Fig. 3. Different from other five sites, daily PM\textsubscript{10} at Taipei is directly calculated using the 24-hourly averaged values instead of calculated values from Chinese API. As shown in Fig. 3, the model yielded good agreement with observations. The correlation coefficients and Normalized Mean Errors range from 0.8–0.95 and 40–70\% respectively (Table 2). As we expected, dust particles dominated the peak PM\textsubscript{10} concentrations at these six sites in the dust event. Concentrations of PM\textsubscript{10} abruptly increased on 20, 21 and 22–23 March in north China (Beijing and Jinan), the Yangtze River Delta (Shanghai and Nanchang) and south China (Xiamen and Taipei), respectively. The model correctly captured this pattern and showed a similar peak magnitude (e.g. in Taipei, Shanghai, and Nanchang). In Beijing and Jinan, the model slightly over-estimated the peak. This is attributed to the imposed upper limit of API (600\(\mu g m^{-3}\)). However, other observations by Zhang (H. Zhang, Peking University, China) revealed a peak of 3000\(\mu g m^{-3}\) hourly concentrations (unpublished data), which are consistent with this simulation (3300\(\mu g m^{-3}\)). The simulation under predicted observed PM\textsubscript{10} in Xiamen city, probably due to the stronger simulated deposition processes.

Lidar observations provide a good opportunity to evaluate the model’s ability to predict vertical aerosol profiles. Figure 4 shows the observed and simulated vertical profiles of dust extinction coefficients during 15–24 March 2010. The two-wavelength Mie-scattering lidar observations (1064 nm and 532 nm) are taken from the National Institute for Environmental Studies (NIES), Japan (http://www-lidar.nies.go.jp/). Clearly, the simulation agrees well with the overall evolution of the Lidar extinctions pattern. During the 19–22 March dust episode, the dust plume at three stations was constrained below 3 km and maintained a rather homogeneous profile. At Nagasaki and Hedo, another dust event on 19 March in the 1–3 km layer was accompanied by observations of a few dust particles on the ground. The model also captured this phenomenon.

Figure 5 compares observed and simulated SO\textsubscript{2} and NO\textsubscript{2} in Beijing and Shanghai during 15–24 March 2010. The model agrees with observations quite well, except for a significant underestimation of NO\textsubscript{2} in Shanghai. The correlation coefficients are in the range 0.49–0.67. In particular, the decrease of SO\textsubscript{2} and NO\textsubscript{2} concentrations in the dust episode is captured by the model. The underestimation of NO\textsubscript{2} in Shanghai is caused by the uncertainties of the REAS emission inventory (0.5\(^\circ\) \times 0.5\(^\circ\)) in the Yangtze River Delta.

The model’s ability to simulate sulfate and nitrate in non-dust and dust periods is critical in assessing the mixing extent in this study. Figure 6 presents a comparison of simulated and observed sulfate and nitrate in Shanghai and Xiamen during the pre-dust, dust and after-dust periods in base case. Both the model and observations show that SO\textsubscript{4}\(^{2-}\) and NO\textsubscript{3}\(^{-}\) significantly increased by factors of 2–3 times those in the pre-dust
Fig. 4. Lidar measurements (532 nm) (left panel) and simulations (550 nm) (right panel) of extinction coefficients during 15–24 March 2010 at Chiba, Nagasaki and Hedo.

Fig. 5. Observed (black) and simulated (red) concentrations (ppbv) of SO$_2$ (upper panel) and NO$_2$ (lower panel) at Beijing (left panel) and Shanghai (right panel) during 15–24 March 2010. N and r represent numbers of paired samples and correlation coefficients, respectively. $C_{\text{obs}}$ and $C_{\text{mod}}$ are average concentrations of observed and modeled species. The gray shades are the periods in the dust-storm episode at two cities.

and after-dust periods at both sites. The simulation clearly indicates that the mixing of minerals with anthropogenic pollutants during the long-range transport played the dominant role in this increase, while anthropogenic SO$_2$$^-$$^-$ and NO$_3$$^-$$^-$ particles decreased along with their precursors (SO$_2$ and NO$_2$) during the duststorm (Fig. 5). In the next section, we will discuss where the mixing of minerals with pollutants occurred (Figs. 7 and 8). Figure 6 also shows that the simulation underestimated the concentration of sulfate at Xiamen. This was caused by the underestimation of dust particle concentrations (as shown in Fig. 3). In fact, the simulated mass fraction of sulfate (5.2 %) in the total particles is consistent with observations (4.8 %).

Unfortunately, few observations of the size distribution of dust particles are available for this dust episode, although we
Fig. 7. Simulated sulfate (a, d) and nitrate (c, e) (µg m\(^{-3}\)) captured by dust particles by heterogeneous chemistry along the 3-day back-trajectories at Shanghai and Xiamen during the dusty periods. Also shown is the mass fraction of dissolved Fe (Fe(II)) in total Fe (e, f). The lines in colored lines represent the locations of air masses arrived at Shanghai and Xiamen during dusty days. The colors in colored lines represent the concentration of dust-sulfate and dust-nitrate (µg m\(^{-3}\)) and the mass fraction of dissolved Fe (e, f). The blue contours represent the SO\(_2\) (a, c, d, f) and NO\(_x\) (b, e) emission rate (µg m\(^{-2}\) s\(^{-1}\)).

were able to collect fine and coarse particles from Shanghai and Xiamen (Table 3). These results showed the deposited particles coarsened as the plume moved south. The mass fraction of fine particles in Xiamen was twice of that in Shanghai. The model reproduced this pattern reasonably well. The simulated fine particle fraction (47.9 %) in Xiamen was very close to the observation (43 %). However, the model overestimated the mass fraction of fine particles in Shanghai. This is likely caused by the local re-suspension of deposited dust particles. Shanghai is closer to the dust source regions than Xiamen, and hence depositions contained more coarse particles. The strong wind in the dust episode may have re-entrained more coarse deposited particles into the atmosphere in Shanghai. Yuan et al. (2008) found that this re-suspension contributed 10–30 % of TSP (Total suspended particles) in Beijing. Most dust models, including NAQPMS in this study, fail to reproduce the re-suspension process.

In general, the above model validation indicates that NAQPMS is capable of reproducing the temporal and spatial distributions of observed dust particles and anthropogenic pollutants. It can also reproduce the chemical evolution of pollutants on dust particles. The reasonable performance provides some confidence in the model-derived results.

4 Results

4.1 Level of mixing between mineral dust and pollutants

As shown in Fig. 6, SO\(_2\)\(^{-}\) resulting from the mixing of minerals with pollutants contributed more than 50 % to the total sulfate in both Shanghai and Xiamen. Figure 7 shows the chemical evolution of SO\(_2\)\(^{-}\), NO\(_3\)\(^{-}\) and dissolved Fe along the back-trajectories during dusty days. The back trajectories are calculated by a Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT v4) model based on WRF output with 80 km resolution (http://ready.arl.noaa.gov/HYSPLIT.php). For Shanghai, Fig. 7a–c shows dust particles stayed fresh and experienced no significant mixing with pollutants until they arrived on the North China Plain (30–40° N, 110–120° E). The concentration of SO\(_2\)\(^{-}\) taken up by dust particles significantly increased from 5 µg m\(^{-3}\) to 20 µg m\(^{-3}\) when the air mass reached the North China Plain. Both NO\(_3\)\(^{-}\) on dust particles and Fe solubility show a similar pattern to SO\(_2\)\(^{-}\). This indicates that high concentrations of anthropogenic pollutants on the North China Plain extensively coated these fresh dust particles.
particles transported from the Gobi desert, due to heterogeneous reactions (Table 1). The fraction of Fe(II) in total Fe in Shanghai was in the range 0.8–1.5 %, which was less than previously observed values in a duststorm event in Beijing in April 2002 (1.3–5.3 %) (Zhang et al., 2005). This is likely to be due to the underestimation of the dissolution rate of Fe on dust particles \( R_d \) in this study. The different transport pathway between two episodes is another likely cause. In our study, after the air mass passed Shanghai, mixed dust particles were transported to South China (e.g. Xiamen) along the coast line, where high relative humidity promoted the heterogeneous reactions. Consequently, anthropogenic gases \( \text{SO}_2 \) and \( \text{HNO}_3 \) were more rapidly converted to \( \text{SO}_4^{2-} \) and \( \text{NO}_3^- \) on the dust particles. The fraction of Fe(II) reached 1.6–5.3 % in Xiamen, which indicates that mixing in South China was more active than in North China.

Similarly to the fraction of Fe(II) in total Fe, the fractions of \( \text{SO}_4^{2-} \) and \( \text{NO}_3^- \) captured by dust particles in total sulfate and nitrate are regarded as indices of the degree of mixing. Figure 8 shows the spatial distributions of the percentages (at the surface) of \( \text{SO}_4^{2-} \) and \( \text{NO}_3^- \) captured by dust particles in total sulfate and nitrate masses during the dusty period. Fe(II) in total Fe is also shown. Here the dusty days are defined as the periods with total dust levels greater than 100 µg m\(^{-3}\), as suggested by Tang et al. (2004). It was found that \( \text{SO}_4^{2-} \) mixing with minerals contributed over 60 % to total sulfate over eastern China and its downwind regions (northwestern Pacific), while its contribution was only 10–20 % in the Gobi. The contribution of mixed \( \text{NO}_3^- \) was 70–95 % over the Gobi, eastern China and northwestern Pacific, which indicates that the level of mixing of \( \text{NO}_3^- \) was higher than that of \( \text{SO}_4^{2-} \). This is partially due to the higher \( \text{HNO}_3 \) uptake (HR12 in Table 1) on dust particles than that of \( \text{SO}_2 \) (HR19 in Table 1), particularly in South China and the northwestern Pacific with high RH conditions (Jordan et al., 2003). The more complicated origin of \( \text{SO}_4^{2-} \) is likely to be another cause. In the semi-arid area of the Chinese Loess Plateau, sulfate usually covers 0.01–0.45 % of soils or sands. In plumes with high dust loadings (i.e. dust source regions), the emitted mineral dust significantly contributes to total sulfate mass, which decreases the fraction of mixed sulfate. Our results for \( \text{SO}_4^{2-} \) are higher than those in previous studies. For example, Tang et al. (2004) estimated that dust enhanced sulfate by up to 25 % during a duststorm event in 2001. This indicates the strong mixing intensity of the duststorm event during 19–22 March 2010.

### 4.2 Regional dust influences

In order to investigate the influences of dust on the regional atmospheric and oceanic environment, a sensitivity simulation without mixing processes was performed. The difference between the sensitivity and base simulation reveals the influence of dust.

#### 4.2.1 Influences on concentrations of atmospheric tracer gases

As well as aerosol compositions, the mixing of minerals with anthropogenic pollutants also alters the regional-scale concentrations of gaseous pollutants. Figure 9 shows the averaged influence of dust on surface \( \text{HNO}_3 \), \( \text{O}_3 \) and \( \text{SO}_2 \) during 19–22 March 2010. The average \( \text{HNO}_3 \) decrease due to heterogeneous reactions was 20–90 %, with the strongest decrease occurring in western China and in the Pacific to the south and east of Japan (up to 80–90 %) (Fig. 9a). The distribution of the \( \text{SO}_2 \) decrease was similar to that of \( \text{HNO}_3 \), but its impact was smaller (5–40 %) (Fig. 9c). Moreover, the impact area was much smaller. Surface \( \text{O}_3 \) are decreased 5–60 % due to the heterogeneous reactions (HR13). The band of large decrease is in the western Pacific which is consistent with \( \text{SO}_2 \). For \( \text{O}_3 \), the decrease was in the range 5–30 % (Fig. 9b). An interesting point is that the influences of dust in polluted regions (e.g. the North China Plain: 30–40° N and 110–120° E) were much smaller than those in remote regions. This is because high emission rates and complex photochemical systems on the North China Plain can offset part of their heterogeneous losses. Our estimations are consistent with the study of Tang et al. (2004), who estimated that surface \( \text{HNO}_3 \), \( \text{SO}_2 \) and \( \text{O}_3 \) decreased by 20 %, 20 % and 95 %, respectively, due to heterogeneous reactions on dust particles.

#### 4.2.2 Fe(II) deposition and its potential influences on marine primary productivity

A large number of studies have demonstrated that dissolved iron (Fe(II)) in surface seawater is the limiting nutrient factor for primary productivity in high-nitrate low-chlorophyll (HNLC) regions of ocean. Deposition of Asian mineral dust is thought to be the major source of Fe(II) in the Pacific. The Fe(II) deposited into the ocean may substantially affect ocean productivity, and by altering the uptake of carbon dioxide through the ocean, may even affect the global greenhouse effect (Zhuang et al., 2003).

The oxidation of \( \text{SO}_2 \) and \( \text{HNO}_3 \) on dust particles during the long-range transport largely enhances dissolved Fe deposition in Pacific. Observations show that this oxidation increased Fe(II) mass fraction in total Fe from 0.4±0.3 % in Gobi to 1.3–2.6 % in the offshore region (Zhuang et al., 1992, 1993a, 2002).

Table 3. Fractions of fine and coarse particles at Shanghai and Xiamen during the dust episode. Observations at Xiamen are from Zhao et al. (2011).

<table>
<thead>
<tr>
<th></th>
<th>Obs</th>
<th>Mod</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shanghai</td>
<td>Fine (&lt;2.5 µm)</td>
<td>18.9</td>
</tr>
<tr>
<td></td>
<td>Coarse (&gt;2.5 µm)</td>
<td>81.1</td>
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<tr>
<td>Xiamen</td>
<td>Fine (&lt;2.5 µm)</td>
<td>43.0</td>
</tr>
<tr>
<td></td>
<td>Coarse (&gt;2.5 µm)</td>
<td>57.0</td>
</tr>
</tbody>
</table>
2003). Over the remote mid-Pacific, Fe(II) on Asian dust particles continued rising, to 5–10%. Figure 8c shows the simulated mass fraction of Fe(II) in total Fe on dust particles in this study. The simulation captures this observed increase of Fe solubility with distance from dust source regions (i.e. the Gobi) well.

Figure 10 shows simulated dry, wet and total deposition of Fe(II) during this duststorm event. In order to isolate the impact of mixing processes on Fe(II) deposition, Fe(II) deposition calculated in a sensitivity simulation without mixing processes is summarized in Table 4.

Fe(II) deposition in the ocean off East Asia ranged from 20 to 100 µg m\(^{-2}\) (Fig. 10a). The largest depositions occurred in the East China Sea (40–60 µg m\(^{-2}\)), and in the Sea of Japan and the Kuril Islands (100 µg m\(^{-2}\)), which comprised dry deposition (Fig. 10b) and wet deposition (Fig. 10c), respectively. As shown in Fig. 10b and c, dry deposition dominated the total deposition of Fe(II) close to the Chinese mainland, while wet deposition played a more important role in the far downwind ocean. Table 4 lists atmospheric deposition fluxes of Fe(II) to the ocean in base and sensitivity simulations. Clearly, the mixing of minerals with anthropogenic pollutants significantly increased the deposition of Fe(II).

Assuming that all the Fe(II) transported and deposited into oceans are bioavailable to phytoplankton. The input can be converted to carbon uptake by using a cellular Fe : C ratio of 10 µmol mol\(^{-1}\) in the subarctic Pacific (e.g. the Kuril Islands: 150° E, 45° N) where observed concentration of dissolved Fe (e.g. the Kuril Islands: 150° E, 45° N) was about 0.2 nmol l\(^{-1}\) (Sunda and Huntsman, 1997). If we neglect removal by abiotic scavenging, our simulated 100 µg m\(^{-2}\) Fe(II) deposition (over 4 days) in the Kuril Islands can create a marine primary productivity of \(\sim 520\) mg C m\(^{-2}\) d\(^{-1}\), which can support almost 100% of the observed mean marine primary produc-

| Table 4. Atmospheric deposition Fluxes of Fe(II) to the ocean in the model domain during 19–22 March 2010 in the base (mixing processes) and sensitivity (no mixing processes) simulations. |
|---|---|---|
| Base(ton) | 128 | 199 | 327 |
| Sensitivity(ton) | 51 | 20 | 71 |

5 Discussion

5.1 Uncertainties in the modeling of mixing between mineral dust and pollutants

5.1.1 Uncertainty in uptake coefficients

Previous studies show that large uncertainties in the uptake coefficients (\(\gamma\)) exist in laboratory studies, which could bring uncertainties to evaluate the impact of mineral dust on the tropospheric chemistry. Zhu et al. (2010) suggested that differences of uptake coefficients of SO\(_2\) and HNO\(_3\) on mineral dust (HT12 and HR19) are several orders of magnitude (HR12: 5.0 \times 10^{-7} - 2.6 \times 10^{-4}; HR19: 1 \times 10^{-3} - 1.7 \times 10^{-1}). In this study, we make use of the sensitive analysis to examine the uncertainties caused by HR12 and HR19. In order to cover the factor of uncertainties reported by Zhu et al. (2010), simulations in which one heterogeneous reaction (HR12 or HR19) was calculated individually with either the “lower limit” or “upper limit” are performed. This uncertainty (\(U\)) was calculated by the following formulation:

\[
U = \left| \frac{y_{\gamma}(\text{upper}) - y_{\gamma}(\text{lower})}{y_{\gamma}(\text{base})} \right|
\]

where \(y_{\gamma}(\text{upper}), y_{\gamma}(\text{lower})\) and \(y_{\gamma}(\text{base})\) represents the impacts of mineral dust on pollutants under conditions with upper limit, lower limit and values in Table 1, respectively.

Figure 11 presents the observed and simulated sulfate and nitrate at Shanghai and Xiamen during the dusty period in simulations with base, upper limit and little limit of \(y_{\text{SO}_2}\) (HR19) and \(y_{\text{HNO}_3}\) (HR12). Clearly, the simulated nitrate and sulfate strongly depended on uptake coefficients. At Shanghai, the simulated nitrate and sulfate in base case where \(y\) values are taken from Table 1 were very close to observations. This indicates \(y_{\text{SO}_2}\) and \(y_{\text{HNO}_3}\) in this study is suitable...
Fig. 10. Simulated (a) total deposition (µg m$^{-2}$), (b) dry deposition (µg m$^{-2}$), (c) wet deposition (µg m$^{-2}$) of Fe(II) over East Asia during 19–22 March 2010.

Fig. 11. Observed and simulated sulfate and nitrate at Shanghai and Xiamen during the dusty period in simulations with base, upper limit and lower limit of uptake coefficients of HR19 and HR12.

5.1.2 Uncertainty in dust shapes

It is known that mineral dust particles have irregular shapes. Okada et al. (2001) found that most of the dust particles in China (0.2–4 µm) can be considered as ellipsoid characterized by an aspect ratio of $\sim 1.5$ (ratio between longer and short axis of an ellipse fitted to the particle outline). So the assumption of spherical particles in this study might cause errors because irregular particles could have different surface areas with sphere particles.

In this study, a method correcting surface area of irregular shapes was used to evaluate the impact of dust shapes on the heterogeneous reactions (Giroux, 2003). In this method, surface area of irregular particle can be calculated by a sphericity factor ($\phi_p$) in the formula:

$$S_p = S/\phi_p$$

where $S$ and $S_p$ represent the surface area of spherical and irregular particle, respectively. $V_p$ is the particle volume. When $\phi_p$ is equal to 1.0, the particle is sphere, and when it is equal to 0.1, the particle is elongated.

The sphericity factor ($\phi_p$) is expressed in terms of aspect ratio ($\lambda$) as the following formula:

$$\phi_p = \frac{2\lambda^2}{1 + \frac{\lambda^2}{\sqrt{\lambda^2-1}}} \arcsin \left( \sqrt{1 - \frac{1}{\lambda^2}} \right)$$

In China deserts (Taklamakan desert, Gobi), observed aspect ratio of dust is $\sim 1.5$ (Okada et al., 2001). So $\phi_p$ is calculated to be 0.97, which means surface areas errors caused by irregular shape are less than 5%.
5.2 Implications on regional atmosphere

From a photochemical perspective, heterogeneous chemistry on mineral dust surfaces that directly affects gaseous (NO$_2$, HNO$_3$ and O$_3$) is of great importance. By altering the complex set of photochemical reactions of O$_3$ and its precursors, the heterogeneous pathway perturbs photochemical oxidant (OH and HO$_2$) and thus atmospheric oxidation capacity (Bian et al., 2003). Furthermore, the heterogeneous pathway may impact the self-cleaning of many atmospheric primary pollutants (VOCs) and formation of secondary pollutants. In this study, mineral dust decreased HNO$_3$ and NO$_2$ concentrations by up to 80–90% and 40%, respectively. Indirect ozone due to HNO$_3$ uptake likely exceeds direct heterogeneous uptake of O$_3$. This indicates that heterogeneous chemistry on mineral dust surfaces may be of importance in the chemical balance of the troposphere over East Asia.

The chemical interactions between the gas and dust particles are able to result in increased amounts of sulfate and nitrate in the aerosol, and a change in size distributions of these anthropogenic pollutants. Under non-dust conditions, the sulfate formation is controlled by the nucleation of H$_2$SO$_4$ and H$_2$O and coagulation processes, and then concentrates in the fine particles. At a large dust loading, the sulfate formation due to surface uptake becomes dominant and its mass exhibits a maximum in coarse mode (Jordan et al., 2003). As shown in Fig. 8, sulfate coating dust particles by heterogeneous reactions covered 60–80% of total sulfate in this study. This would indicate that dust particles shifted sulfoates from fine to coarse mode over Eastern China and Pacific in this study. The coating processes also leads dust particles an increase in their hygroscopicity. The change of size distributions of sulfate and nitrate and hygroscopicity could alter aerosol optical properties, direct radiative forcing characteristics and has important implications for regional climate in East Asia (Tang et al., 2004; Liao et al., 2005).

6 Conclusions

The objective of this study was to examine the extent to which Asian mineral dust is mixed with anthropogenic pollutants in a super duststorm event during 19–22 March 2010. Meanwhile, influences of mixing processes on regional atmospheric environmental and oceanic biogeochemical cycles were also investigated.

A variety of observational data including in-situ ground-based concentrations of PM$_{10}$ and its chemical compositions, extinction coefficient data of dust particles from two-wavelength Mie-scattering lidar observations, and MODIS satellite AOD550 retrievals were used to validate the performance of the NAQPMS model. The model validation indicates that NAQPMS is capable of reproducing the long-range transport of dust particles from the Gobi desert to the South China Sea and northwestern Pacific. The vertical profile of dust plumes is also reproduced well. Even more importantly, NAQPMS captures the chemical evolution of dust particles during the long-range transport: for example, the increasing concentrations of sulfate, nitrate and Fe(II).

We found that mixing processes along the trajectory of the long-range transport of Asian dust strongly increased pollutant levels in the dust particles. In Shanghai, a megacity located in the Yangtze River Delta, SO$_4^{2-}$ and NO$_3^-$ levels captured by minerals reached $\sim$15 and 10 µg m$^{-3}$, respectively. Even in southern China (Xiamen City), the mixing processes contributed $\sim$60% of the SO$_4^{2-}$ and $\sim$90% of the NO$_3^-$, with the help of high RH. Analysis based on back trajectories indicates that the active mixing stared in the North China Plain, which is consistent with the distribution of anthropogenic emissions. In general, mixing SO$_4^{2-}$ with minerals made an overall contribution to total sulfate of $\sim$60% over eastern China and its downwind regions (northwestern Pacific), while its contribution was only 10–20% in the Gobi. The equivalent contribution of mixed NO$_3^-$ was 70–95% over the Gobi, eastern China and northwestern Pacific. The Fe solubility (the fraction of Fe(II) in total Fe) resulting from oxidization of SO$_2$ on dust particles showed an increase with distance from dust source regions, of the order of $\sim$0.5–5%.

The mixing of minerals with pollutants could have important impacts on the atmospheric composition and marine primary productivity over East Asia. In this study, dust particles significantly decreased surface HNO$_3$, SO$_2$ and O$_3$ by up to 90%, 40% and 30%, respectively, due to the heterogeneous chemistry. We estimated that the Fe(II) deposition in the ocean off East Asia totally reached 327 tons during the 4-day dust event, mostly due to the mixing of minerals with pollutants. The largest deposition occurred in the East China Sea (40–60 µg m$^{-2}$), and in the Sea of Japan and Kuril Islands (100 µg m$^{-2}$). A rough estimation showed that our simulated 100 µg m$^{-2}$ Fe deposition (over 4 days) in the subarctic Pacific(e.g. Kuril Islands) can create a marine primary productivity of $\sim$520 mgC m$^{-2}$ d$^{-1}$, which is consistent with the observed mean marine primary productivity in spring in this region (526 mgC m$^{-2}$ d$^{-1}$). Our results suggest that the Fe(II) deposition in dust particles has important impact on biological primary marine productivity in subarctic Pacific.

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