





## 22 **Abstract**

23       The global source-receptor relationships of sulfate concentration, direct and  
24 indirect radiative forcing (DRF and IRF) from sixteen regions/sectors for years  
25 2010-2014 are examined in this study through utilizing a sulfur source-tagging  
26 capability implemented in the Community Earth System Model (CESM) with winds  
27 nudged to reanalysis data. Sulfate concentrations are mostly contributed by local  
28 emissions in regions with high emissions, while over regions with relatively low SO<sub>2</sub>  
29 emissions, the near-surface sulfate concentrations are primarily attributed to non-local  
30 sources from long-range transport. The export of SO<sub>2</sub> and sulfate from Europe  
31 contributes 16–20% of near-surface sulfate concentrations over North Africa,  
32 Russia/Belarus/Ukraine (RBU) region and Central Asia. Sources from the Middle  
33 East account for 15–24% of sulfate over North Africa, Southern Africa and Central  
34 Asia in winter and autumn, and 19% over South Asia in spring. Sources in RBU  
35 account for 21–42% of sulfate concentrations over Central Asia. East Asia accounts  
36 for about 50% of sulfate over Southeast Asia in winter and autumn, 15% over RBU in  
37 summer, and 11% over North America in spring. South Asia contributes to 11–24% of  
38 sulfate over Southeast Asia in winter and spring. Regional source efficiencies of  
39 sulfate concentrations are higher over regions with dry atmospheric conditions and  
40 less export, suggesting that lifetime of aerosols, together with regional export, is  
41 important in determining regional air quality. The simulated global total sulfate DRF is  
42  $-0.42 \text{ W m}^{-2}$ , with  $-0.31 \text{ W m}^{-2}$  contributed by anthropogenic sulfate and  $-0.11 \text{ W m}^{-2}$   
43 contributed by natural sulfate, relative to a state with no sulfur emissions. In the



44 Southern Hemisphere tropics, dimethyl sulfide (DMS) contributes 17–84% to the total  
45 DRF. East Asia has the largest contribution of 20–30% over the Northern Hemisphere  
46 mid- and high-latitudes. A 20% perturbation of sulfate and its precursor emissions  
47 gives a sulfate incremental IRF of  $-0.44 \text{ W m}^{-2}$ . DMS has the largest contribution,  
48 explaining  $-0.23 \text{ W m}^{-2}$  of the global sulfate incremental IRF. Incremental IRF over  
49 regions in the Southern Hemisphere with low background aerosols is more sensitive to  
50 emission perturbation than those over the polluted Northern Hemisphere.



## 51 **1. Introduction**

52 Sulfate is an important aerosol that poses health risks (Fajersztajn et al., 2013;  
53 Xu et al., 2013; Peplow, 2014) and sulfur deposition is a major driver of ecosystem  
54 acidification (Driscoll et al., 2010). Due to long-range transport, local sulfate pollution  
55 could result from intercontinental influences, making domestic efforts of improving air  
56 quality inefficient (Part et al., 2004; Bergin et al., 2005; Liu and Mauzerall, 2007). In  
57 addition, sulfate aerosol substantially perturbs the radiation budget of the Earth  
58 directly through scattering incoming solar radiation and indirectly through modifying  
59 cloud microphysical properties (Lohmann and Feichter, 2005; Stevens and Feingold,  
60 2009; Myhre et al., 2013). On a global average basis, anthropogenic sulfate aerosol  
61 contributes a negative direct radiative forcing (DRF) of  $-0.4 \pm 0.2 \text{ W m}^{-2}$  (Boucher et  
62 al., 2013). The negative radiative forcing from sulfate partly offsets the positive  
63 radiative forcing from greenhouse gases. Therefore, accurate understanding of  
64 source attribution of sulfate and its radiative forcing is important for both regional air  
65 quality and global climate mitigation (Shindell et al., 2012), which are of great interest  
66 to not only science community but also the general public and policymakers.

67 Sulfate aerosol is produced through oxidation of sulfur dioxide ( $\text{SO}_2$ ) by the  
68 hydroxyl radical (OH) in gas phase and aqueous phase oxidation mainly by hydrogen  
69 peroxide ( $\text{H}_2\text{O}_2$ ) (Martin and Damschen, 1981). The  $\text{SO}_2$  precursor is mainly emitted  
70 from fossil-fuel combustion (Lu et al., 2010). In recent decades,  $\text{SO}_2$  emissions from  
71 many developing countries in East Asia and South Asia have increased substantially  
72 as a result of accelerated urbanization and rapid economic growth (Streets et al.,



73 2000; Pham et al., 2005). In contrast, due to air pollution regulations, SO<sub>2</sub> emissions  
74 in North America and Europe have decreased significantly since 1980–1990 (Smith  
75 et al., 2011; Prechtel et al., 2001). As a consequence, source attribution of sulfate has  
76 changed with time over recent decades.

77 Previous studies have reported that regional aerosols, including sulfate, are  
78 produced not only by domestic emissions, but also by distant sources through  
79 long-range transport (Jacob et al., 2003; Jaffe et al., 2003; Park et al., 2004; Heald et  
80 al., 2006; Liu et al., 2008; Liu et al., 2009; Yu et al., 2012). For example, the strong  
81 anthropogenic emissions over East Asia have led to an increasing interest in  
82 quantifying the impact of aerosols exported from East Asia. Recent studies indicate  
83 that the transpacific transport of sulfate from East Asia contributes to 30–50% of the  
84 background (sulfate produced from non-local emissions) surface concentrations in  
85 the Western U.S. and 10–30% in the Eastern U.S. (Park et al., 2004; Hadley et al.,  
86 2007; Liu et al., 2008), which are larger than contributions from all other foreign  
87 sources (Liu et al., 2009). In addition, among the major emitting regions assessed for  
88 2001 conditions, European sources were shown to account for 1–5 µg m<sup>-3</sup> of surface  
89 sulfate concentration over northern Africa and western Asia, and their contribution to  
90 East Asia (0.2–0.5 µg m<sup>-3</sup>) was twice as much as the contribution (0.1–0.2 µg m<sup>-3</sup>) of  
91 Asian sources to North America (Chin et al., 2007).

92 Due to the important role of sulfate aerosol in the climate system, knowing the  
93 relative significance of sulfate radiative forcing from different source regions is useful  
94 for climate mitigation. Some previous studies examined the impact of emission



95 reductions on global and regional DRF and the influence of long-range transport (Yu  
96 et al., 2013; Bellouin et al., 2016; Stjern et al., 2016). Yu et al. (2013) examined  
97 changes in aerosol DRF resulting from a 20% reduction in anthropogenic emissions  
98 from four major polluted regions (namely North America, Europe, East Asia, and  
99 South Asia) in Northern Hemisphere, using simulations by nine models from the first  
100 phase of the Hemispheric Transport of Air Pollution (HTAP1). They found that 31% of  
101 South Asia sulfate aerosol optical depth over South Asia was contributed by non-local  
102 sources. Based on the HTAP2, Stjern et al. (2016), using results from ten models,  
103 further assessed global and regional DRF from a 20% reduction in emissions over  
104 seven regions including North America, Europe, South Asia, East Asia, Russia, the  
105 Middle East, and the Arctic. They found that the 20% reduction in emissions in South  
106 Asia and East Asia largely perturbed the radiative balance for other regions. However,  
107 these studies focused on only the limited number of source regions over the Northern  
108 Hemisphere. Continents and subcontinents over the tropics and Southern  
109 Hemisphere are also important source and receptor regions for the sulfate radiative  
110 forcing, especially for indirect forcing due to stronger aerosol-cloud interactions in  
111 clean environments (Koren et al., 2014). Bellouin et al. (2016) quantified the radiative  
112 forcing efficiency based on simulations of a 20% reduction in emissions from four  
113 source regions/sectors in year 2008, and reported that, with aerosol-cloud  
114 interactions included, models simulated higher radiative forcing efficiency of sulfate  
115 compared to previous studies (Myhre et al., 2013, Shindell et al., 2013; Yu et al.,  
116 2013). Few studies have quantified systematically the global source-receptor



117 relationships of sulfate indirect radiative forcing that can be attributed to  
118 local/non-local source regions and anthropogenic/natural source sectors.

119 In this study, we introduce an explicit sulfur tagging technique into the  
120 Community Earth System Model (CESM), in which sulfate aerosol and its precursor  
121 emissions from fourteen major source regions and two natural source sectors are  
122 tagged and explicitly tracked. We quantify source region/sector contributions to  
123 regional and global sulfate mass concentrations, and direct and indirect radiative  
124 forcing (DRF and IRF) of sulfate.

125 Model description, emissions datasets, and model experiments are shown in  
126 Sect. 2. Section 3 gives the comparison of modeled concentrations of sulfate and  
127 SO<sub>2</sub> with a variety of observations. Section 4 shows model results for source  
128 attributions of near-surface sulfate and SO<sub>2</sub> concentrations over various receptor  
129 regions. Source attributions of DRF and IRF of sulfate are discussed in Section 5.  
130 Section 6 summarizes all the results and main conclusions.

131

## 132 **2. Methods**

133 We use the version 5 of the Community Atmosphere Model (CAM5), which is the  
134 atmospheric component of CESM (Hurrell et al., 2013), to simulate the sulfate aerosol  
135 and calculate its DRF and IRF. The modal aerosol treatment in CAM5 (Liu et al., 2012)  
136 predicts number mixing ratios and mass mixing ratios of aerosols, distributed in three  
137 lognormal modes. A set of modifications to CAM5 that improves wet scavenging of  
138 aerosols and convective transport reported by Wang et al. (2013) has also been



139 implemented in the model used in this study. Parameterizations of aerosol optical  
140 properties, cloud droplet nucleation, and aerosol-cloud interactions are described in  
141 Neale et al. (2012). In addition to the standard radiative fluxes calculated with all  
142 aerosols included, the CESM model has the capability of diagnosing radiative fluxes  
143 for a subset of aerosol species. The difference between the standard and the  
144 diagnosed shortwave radiative fluxes represents the DRF of the excluded aerosol  
145 components in the diagnostic calculation (Ghan, 2013). To investigate IRF of sulfate  
146 from different sources, we define in this study an incremental IRF, calculated as the  
147 difference of cloud radiative forcing by perturbing 20% of sulfate and its precursor  
148 emissions. Note that, the model only considers aerosol effects on stratiform cloud  
149 (Morrison and Gettelman, 2008), and no microphysical impact on convective clouds  
150 is included in the present version.

151 To quantify the regional source attributions of sulfate, for the first time, we  
152 implemented in CESM/CAM5 a sulfur source-tagging capability, similar to the black  
153 carbon tagging method used in H. Wang et al. (2014) and Yang et al. (2017), through  
154 which sulfur gases and sulfate aerosols produced by emissions from independent  
155 sources are tagged. The tool can be used to quantify the source attributions of SO<sub>2</sub>  
156 and sulfate without perturbing source emissions. The black carbon tagging only  
157 required tagging interstitial and cloud-borne black carbon in the accumulation mode.  
158 In contrast, the sulfur tagging requires tagging of interstitial and cloud-borne sulfate in  
159 each of the three modes as well as SO<sub>2</sub>, H<sub>2</sub>SO<sub>4</sub> and dimethyl sulfide (DMS) gases. In  
160 this study, sulfur species produced by emissions from fourteen geographical source



161 regions and two natural source sectors including volcanic eruptions and DMS from  
162 oceans are tagged. The tagged and untagged models have been verified of producing  
163 the same SO<sub>2</sub>/sulfate properties and meteorology. While emissions of organic carbon,  
164 black carbon, sulfate and its precursor gases are all included in the simulations, the  
165 source tagging is used for sulfate and its precursor gases emissions alone.

166 The CEDS (Community Emissions Data System) anthropogenic emissions  
167 (Hoesly et al., 2017) and open biomass burning emissions from Van Marle et al. (2017)  
168 that were produced for the CMIP6 model experiments are used in our simulations. In  
169 CAM5, 97.5% of SO<sub>2</sub> is emitted directly into the atmosphere and 2.5% is emitted as  
170 sulfate aerosol. Natural emissions of volcanic SO<sub>2</sub> and DMS are the same as those  
171 used in AeroCom following Neale et al. (2012), which are kept constant throughout the  
172 selected years in this study. Figure 1a shows the fourteen geographical source  
173 regions tagged in this study, which are consistent with source-receptor regions  
174 defined in HTAP2, including North America (NAM), Central America (CAM), South  
175 America (SAM), Europe (EUR), North Africa (NAF), Southern Africa (SAF), the  
176 Middle East (MDE), Southeast Asia (SEA), Central Asia (CAS), South Asia (SAS),  
177 East Asia (EAS), Russia/Belarus/Ukraine (RBU), Pacific/Australia/New Zealand  
178 (PAN), and rest of the world (ROW, including oceans and polar continents). Table 1  
179 summarizes emissions of combustion SO<sub>2</sub> (anthropogenic + open biomass burning),  
180 volcanic SO<sub>2</sub> emissions (VOL), and DMS emissions over the sixteen tagged source  
181 regions/sectors averaged for the most recent five years (2010–2014) and Figure 1b  
182 presents relative contributions from individual source regions to the global



183 combustion SO<sub>2</sub> emissions. The global combustion SO<sub>2</sub> emissions rate is 57.6 Tg S  
184 yr<sup>-1</sup>, of which more than 98% come from anthropogenic sources. The combustion SO<sub>2</sub>  
185 and sulfate are referred to anthropogenic SO<sub>2</sub> and sulfate hereafter. Detailed  
186 information on the anthropogenic emissions of SO<sub>2</sub> can be found in Hoesly et al.  
187 (2017). East Asia, with regional emission of 17.8 Tg S yr<sup>-1</sup> (31% of global  
188 anthropogenic SO<sub>2</sub>), has the largest total SO<sub>2</sub> emissions, compared to the other  
189 tagged regions. South Asia also emits a large amount of SO<sub>2</sub>, 6.4 Tg S yr<sup>-1</sup> (11%),  
190 followed by 3.4 Tg S yr<sup>-1</sup> (6%) from the Middle East, 3.3 Tg S yr<sup>-1</sup> (6%) from Europe,  
191 3.1 Tg S yr<sup>-1</sup> (5%) from North America, and 2.7 Tg S yr<sup>-1</sup> (5%) from Southern Africa.  
192 The other individual tagged regions have weaker emissions, with a combined  
193 contribution of less than 5%. However, emissions from ROW contribute 11.2 Tg S yr<sup>-1</sup>  
194 (19%) of SO<sub>2</sub> that are mainly from shipping emissions near the continents. In addition,  
195 natural emissions of sulfur are also accounted for, including 12.6 Tg S yr<sup>-1</sup> of SO<sub>2</sub> from  
196 volcanic eruptions, in the range of 10–13 Tg S yr<sup>-1</sup> derived from the Ozone Monitoring  
197 Instrument (OMI) measurement (McLinden et al., 2016), and 18.2 Tg S yr<sup>-1</sup> of DMS.  
198 Figure 2 shows the spatial distribution of SO<sub>2</sub> emissions from each tagged  
199 region/sector as well as DMS emissions. Emissions are spatially heterogeneous even  
200 within the individual tagged regions. For instance, SO<sub>2</sub> emissions in North America are  
201 mainly located in Eastern U.S., and Eastern China accounts for the majority of SO<sub>2</sub>  
202 emissions from East Asia. In addition, seasonal variations in emissions are quite  
203 different among the source regions (Table 1). East Asia, RBU and Europe have  
204 seasonal peak emissions in boreal winter, and Southern Africa shows larger



205 emissions in boreal summer, while emissions from North America are comparable in  
206 winter and summer. Although volcanic SO<sub>2</sub> emissions are scattered near continents, a  
207 large amount of them are injected into the free troposphere. DMS is emitted over  
208 oceans with a boreal winter peak. These heterogeneous spatial and temporal  
209 distributions of emissions could lead to different influences on air quality and radiative  
210 forcing over continents and subcontinents near the source regions.

211 The CAM5 simulation is conducted using a meteorological nudging method (Ma  
212 et al., 2013; Zhang et al., 2014), with winds nudged to the MERRA reanalysis  
213 (Rienecker et al., 2011) every 6 hours. The simulation is integrated for years 2009–  
214 2014, with 2009 for spin-up and 2010–2014 for analysis. A sensitivity simulation with  
215 the same model configuration but having a uniform 20% reduction in sulfur (SO<sub>2</sub>,  
216 sulfate, DMS) emissions globally is performed to quantify source attributions of  
217 incremental IRF of sulfate. Two additional sensitivity simulations with the same  
218 standard model configuration but having a 20% reduction in global DMS emissions  
219 and regional sulfur emissions over North America, respectively, are performed to  
220 validate the decomposition of global incremental IRF into contributions from source  
221 regions/sectors using the tagging method. And one additional sensitivity simulation  
222 with anthropogenic SO<sub>2</sub> emissions fixed at 1850 level globally is performed to  
223 compare incremental IRF and anthropogenic IRF of sulfate. All simulations are  
224 performed at 1.9° latitude by 2.5° longitude horizontal grids and 30 vertical layers.

### 225 **3. Model evaluation**



226 To evaluate the model's performance in simulating sulfate with the latest  
227 emissions from CEDS inventory, the simulated sulfur concentrations are compared  
228 with measurements from regional observation networks. These datasets include the  
229 Interagency Monitoring of Protected Visual Environments (IMPROVE), the European  
230 Monitoring and Evaluation Programme (EMEP), the East Asian Monitoring Network  
231 (EANET), and the China Meteorological Administration Atmosphere Watch Network  
232 (CAWNET, Zhang et al., 2012). Sulfate concentrations observed from IMPROVE,  
233 EMEP and EANET being used here are from 2010 to 2014, covering the same time  
234 period as the simulation, while CAWNET only collected data over 2006–2007. In  
235 order to use the CAWNET data to evaluate 2010-2014 simulation results, we decide  
236 to scale the observed sulfate mass concentrations using the ratio of CEDS  
237 2010-2014 SO<sub>2</sub> emissions to 2006-2007 emissions over China (which is 0.92) for  
238 comparison, thus assuming a linear relationship between SO<sub>2</sub> emissions and sulfate  
239 concentrations.

240 Figure 3 shows the comparison of modeled annual mean near-surface sulfate  
241 concentrations with those from the observational networks. The model successfully  
242 reproduces the global spatial distribution of sulfate with high concentrations over East  
243 Asia and low concentrations over North America and Europe, as well as the spatial  
244 patterns within major continents, for instance, high (low) values over Eastern  
245 (Western) U.S. and high (low) sulfate concentrations over Eastern (Western) China.  
246 The spatial correlation coefficient between simulated and observed sulfate  
247 concentrations globally is +0.86 and is statistically significant at the 95th percentile.



248 Compared to the measurements at the IMPROVE sites over North America, at the  
249 EMEP sites over Europe, and at the EANET sites over part of East Asia (only one site  
250 in China) and Southeast Asia, the model reproduces sulfate concentrations with  
251 biases within  $\pm 20\%$ . However, the model largely underestimates the simulated sulfate  
252 concentrations in China, with normalized mean biases (NMB) of  $-54\%$ , compared to  
253 the CAWNET observations.

254 A few factors could be responsible for the bias between the observed and  
255 modeled sulfate concentrations. Underestimation of local  $\text{SO}_2$  emissions could result  
256 in the simulated low sulfate concentrations (Liu et al., 2012; Wang et al., 2013). Too  
257 frequent liquid clouds and too strong wet scavenging at the mid- and high latitudes in  
258 CESM model can lead to shorter aerosol lifetime and lower concentrations in the  
259 simulation (Wang et al., 2011; Liu et al., 2012; Wang et al., 2013). In addition, the  
260 underestimation of emissions from upwind regions or strong wet scavenging of  
261 aerosols during transport could be another reason for the simulated low bias (Yang et  
262 al., 2017). A too low rate of transformation from  $\text{SO}_2$  gas to sulfate particles in the  
263 model could also contribute to the low bias in sulfate concentrations (Wang et al.,  
264 2016; Li et al., 2017). The bias can also result from the fact that the site  
265 measurements are point observations, while the model results are grid-cell average  
266 that does not consider subgrid aerosol variations (Qian et al., 2010; R. Wang et al.,  
267 2014). Considering the longer lifetime of  $\text{SO}_2$ /sulfate than black carbon, this effect  
268 would be expected to be less significant for  $\text{SO}_2$ /sulfate. In addition, different models  
269 show large discrepancies in simulating sulfate over China (Kasoar et al., 2016). The



270 underestimation of sulfate in China can lead to an underestimation of source  
271 contribution from East Asia of sulfate concentrations, direct and indirect radiative  
272 forcing of sulfate, and forcing efficiencies of sulfate.

273 To evaluate the model results more broadly, we compare the simulated total  
274 column burden of SO<sub>2</sub> with that derived from the OMI measurements (Li et al., 2013),  
275 as shown in Fig. S1. Both the model results and the OMI satellite data are averaged  
276 over 2010–2014. Compared to the OMI SO<sub>2</sub>, the spatial distribution of column burden  
277 of SO<sub>2</sub> is reproduced in CAM5, with a statistically significant spatial correlation  
278 coefficient of +0.57. However, the model largely overestimates the magnitude of SO<sub>2</sub>,  
279 especially over China where the simulated values are about 8 times larger than OMI  
280 data. The large difference between SO<sub>2</sub> burden and OMI retrievals over China must  
281 be due to either an underestimation of SO<sub>2</sub> in OMI products and/or an overestimation  
282 of SO<sub>2</sub> burden in the model results. He et al. (2012) compared in situ measurements  
283 with OMI SO<sub>2</sub> burden over central China and reported a negative bias of 50% in OMI  
284 data, which probably came from cloud contamination, reduced satellite sensitivity to  
285 SO<sub>2</sub> due to aerosols, and spatial sampling bias in the satellite data. It is also worth  
286 mentioning that satellite column-SO<sub>2</sub> retrievals depend on the vertical distribution of  
287 SO<sub>2</sub> assumed in the retrieval algorithm, which could be different from either the  
288 modeled SO<sub>2</sub> profile in this study or the actual profile, which would introduce a bias.

289 The simulated SO<sub>2</sub> near-surface concentrations, however, are also  
290 underestimated by 25% compared to observations over thirteen sites in China (Gong  
291 et al., 2014) shown in Fig. S2a, also suggesting a large bias in satellite retrievals or too



292 much SO<sub>2</sub> simulated in higher altitude. The modeled SO<sub>2</sub> concentrations over  
293 downwind regions of China are underestimated by 45% compared to observations  
294 from EANET sites (Fig. S2b), indicating that the transport of SO<sub>2</sub> from China is  
295 probably underestimated in the model.

296 A less efficient of transformation from SO<sub>2</sub> to sulfate could also lead to  
297 underestimation of sulfate. A recent study by Wang et al. (2016) focusing on the  
298 sulfate pollution over China and London found that aqueous oxidation of SO<sub>2</sub> by NO<sub>2</sub>  
299 was key to an efficient sulfate formation, which has typically been neglected in  
300 atmospheric models and is not considered in the CAM5. Another study by Li et al.  
301 (2017) found that including an aerosol water (HRSO<sub>2</sub>) parameterization in SO<sub>2</sub>  
302 oxidation in a box model could reproduce the observed rapid sulfate formation in Xi'an  
303 over China. More rapid oxidation of SO<sub>2</sub> would reduce SO<sub>2</sub> loss by dry and wet  
304 removal and increase sulfate production, which can partly explain the low bias in the  
305 simulated sulfate concentrations and high bias in SO<sub>2</sub>. In CAM5, 36% of total sulfur  
306 converts into column-integrated sulfate over China, similar to 33% in the Community  
307 Multiscale Air Quality (CMAQ) model (He et al., 2012). However, it changes to 21% in  
308 the bottom model layer (about 992 hPa), indicating that the oxidation of SO<sub>2</sub> may be  
309 underestimated near the surface, which most directly affects the comparison to  
310 near-surface observations. This appears to be a plausible explanation for the  
311 underestimated sulfate concentrations over China and points to a potentially important  
312 direction for future model development.

313 **4. Source attribution of sulfate mass concentrations**



314 Figure 4 shows spatial distributions of modeled fractional contributions to annual  
315 near-surface sulfate concentrations. (The absolute concentrations of sulfate are  
316 shown in Fig. S3). East Asia, ROW, South Asia and the Middle East contribute 16%,  
317 14%, 10% and 7%, respectively, to global annual mean near-surface sulfate  
318 concentration, whereas contributions from the other individual source regions are all  
319 less than 5%. Natural emissions of volcanic SO<sub>2</sub> and ocean DMS account for 11% and  
320 16% of global mean sulfate concentrations. Sulfate concentrations are mostly  
321 contributed by local sources in regions with high emissions, such as Eastern U.S.,  
322 Southern Africa, South Asia, and Eastern China, where local source contributions are  
323 larger than 80%. Over regions with relatively low SO<sub>2</sub> emissions, the near-surface  
324 sulfate concentrations are primarily attributed to non-local sources from long-range  
325 transport. Natural DMS emissions are the source of 80% of near-surface sulfate  
326 concentrations over Southern Hemisphere oceans and 20–60% for Northern  
327 Hemisphere oceans. Over downwind ocean regions of East Asia, emissions from  
328 DMS only account for 20–40% of near-surface sulfate concentrations, showing a  
329 stronger influence of regional transport. Sources from volcanic eruption strongly  
330 influence sulfate concentrations over eruption regions. They are responsible for 10–  
331 40% of near-surface concentrations over Central America and South America, 40–80%  
332 over North Africa and Southeast Asia, but only account for about less than 5% over  
333 East Asia and South Asia where anthropogenic emissions dominate.

334 The spatial distribution of sulfate column burden and relative contributions are  
335 shown in Figs. S4 and S5, respectively. The global average source attribution of



336 column burden does not differ significantly from that of near-surface concentration.  
337 The exception is an increase from 11% to 15% of the relative contribution from VOL to  
338 column burden as compared to near-surface concentration due to injection mostly into  
339 the free troposphere. The DMS contribution decreases from 16% to 11% to  
340 compensate the increase of VOL contribution over oceans. In general, the relative  
341 contribution from local source to column burden within a source region is lower than  
342 that of near-surface concentration.

343 Figure 5 presents relative contributions of major sources to near-surface sulfate  
344 concentrations in neighboring receptor regions along with seasonal mean wind fields  
345 at 850 hPa. (Table S1 summarizes a complete list of numbers characterizing the  
346 source-receptor relationships.) Transport of sulfate shows different patterns in  
347 different seasons, due to the seasonal variability in local precursor emissions, lifetime  
348 of sulfate, and meteorology, such as wind fields and precipitation. Sulfate originating  
349 from North America, Central America and South America do not show significant  
350 contributions (relative contribution less than 10%) to sulfate over other tagged regions  
351 in all seasons because of the relatively low sulfate concentrations over these regions  
352 and the long intercontinental transport pathways.

353 The export of sulfate from Europe contributes to about 16–20% of near-surface  
354 sulfate concentrations over North Africa, RBU and Central Asia in all seasons due to  
355 the westerly jet over the eastern European boundary and northerly winds over  
356 southern boundary. Sulfate concentrations in North Africa and Southern Africa are  
357 relatively low and there is no significant export to other regions.



358       The Middle East has relatively high concentrations of sulfate (Fig. S3). Sulfate  
359       from this region can be effectively transported to the surrounding receptor regions:  
360       North Africa, Southern Africa, Central Asia, and South Asia. This export accounts for  
361       15–24% of sulfate concentrations over North Africa, Southern Africa and Central Asia  
362       in DJF and SON, and 19% over South Asia in MAM. Sources in the RBU explain  
363       about 21–42% of sulfate concentrations over Central Asia, especially in JJA, with  
364       northerly winds over north boundary of Central Asia driving transport from this region.  
365       Central Asia does not have a discernable export of sulfate due to low emissions in  
366       most seasons, except that it accounts for 13% of sulfate over the RBU region in DJF  
367       when source emissions are the largest.

368       Northerly winds over East Asia in DJF and SON associated with the East Asian  
369       winter monsoon transport sulfate from highly polluted Eastern China to Southeast  
370       Asia, which accounts for about 50% of near-surface sulfate concentrations over  
371       Southeast Asia in these months. The oxidation of SO<sub>2</sub> is expected to peak in JJA  
372       because of the high temperature and humidity, and more sunlight. With the help of  
373       southerly winds of East Asian summer monsoon, East Asia contributes to 15% of  
374       sulfate concentrations over RBU in JJA. Due to the strong westerly jet in MAM,  
375       sulfate originating from East Asia has a long-range transport across the North Pacific  
376       and accounts for 11% of near-surface sulfate concentrations and 25% of total  
377       imported sulfate (without local contributions) over North America. The transport of  
378       sulfate from South Asia contributes 11–24% of sulfate in Southeast Asia in DJF and



379 MAM. These results, however, have additional uncertainties due to the SO<sub>2</sub>/sulfate  
380 bias in the model for East Asia discussed previously.

381 Source-receptor relationships for sulfate column burden are summarized in Table  
382 S2. Compared to the near-surface concentrations, the sulfate column burden  
383 contributed by local sources is much lower in all the receptor regions due to the more  
384 efficient long-range transport of aerosols in the free atmosphere. Annually, the local  
385 contribution over North America decreases from 67% for near-surface concentration  
386 to 33% for column burden. The contributions of non-local sources from East Asia and  
387 South Asia increase from 7% and 1% for near-surface concentration to 24% and 10%  
388 for column burden, respectively, to the sulfate over North America. In addition, South  
389 Asia contribution to sulfate in East Asia, and East Asia contribution to sulfate in RBU  
390 and Europe also significantly increase for column burden compared to near-surface  
391 concentrations.

392 Figure 6 shows local contributions (i.e., from sources within the tagged regions)  
393 to near-surface sulfate concentrations. Averaged over individual tagged regions,  
394 contributions from local sources dominate (i.e., local contributions > 50%) over North  
395 America, South America, Europe, Southern Africa, the Middle East, South Asia, and  
396 East Asia. Imports dominate near-surface sulfate concentrations (i.e., local  
397 contributions < 50%) over the rest of tagged land regions. Within each tagged region,  
398 whether local source or import dominates depends on specific locations. For instance,  
399 over Eastern China, because of high anthropogenic emissions, local contribution to  
400 sulfate concentration is larger than 80%, whereas import from other source regions



401 dominates sulfate over the less economically developed Western China. The same  
402 difference can be found between Eastern and Western U.S. of the tagged North  
403 America. Over oceans in the Southern Hemisphere, natural sources of DMS  
404 contribute the largest to local sulfate concentrations (Fig. 4), whereas long-range  
405 transport dominates over the North Pacific in DJF and MAM.

406 Figure 7 presents the aggregate, seasonal relative source contributions to area  
407 weighted average near-surface sulfate concentrations over land/ocean in the  
408 Northern/Southern Hemisphere. Over land in the Northern Hemisphere, sulfate  
409 concentration is mainly attributed to sources from East Asia, South Asia, the Middle  
410 East, ROW and volcanic eruption, with relative contributions of 22–29%, 9–16%, 8–  
411 14%, 9–11%, and 6–13%, respectively. Over ocean in the Northern Hemisphere,  
412 although contribution from ROW, volcanic SO<sub>2</sub> and DMS increase dramatically  
413 compared to land, contributions from East Asia and South Asia do not have a large  
414 decrease, especially in DJF, MAM and SON when aerosol outflow from Asia is strong  
415 (Yu et al., 2012; Yang et al., 2015). Over land in the Southern Hemisphere, mean  
416 sulfate concentration is dominated by sources in Southern Africa, having a  
417 contribution of 33–43%, followed by 13–25% from South America. Emissions from  
418 DMS drive sulfate over ocean in the Southern Hemisphere in all seasons contributing  
419 27–63% of sulfate, although Southern Africa contributes 20% of sulfate in JJA.

420 Figure 8 shows seasonal and annual mean regional concentration efficiencies of  
421 sulfate from the tagged source regions/sectors, defined as the local contribution to  
422 near-surface sulfate concentration divided by the corresponding sulfur emissions



423 from that region. (Table S3 provides the numeric values.) The regional concentration  
424 efficiency represents the relationship between local contribution to sulfate  
425 concentration and local emission, which is influenced by many factors, such as local  
426 production of sulfate from the emitted SO<sub>2</sub>, aerosol removal and export. Note that, the  
427 receptor region of ROW is used to calculate efficiencies of the VOL and DMS source  
428 sectors, which leads to low biases in efficiencies. The efficiencies over the Middle  
429 East show high values in almost all seasons due to dry atmospheric conditions  
430 favoring long aerosol lifetime (e.g., Wang et al., 2014; Stjern et al., 2016). The  
431 efficiencies are also high over South Asia in DJF and SON, but low in MAM and JJA  
432 due to strong wet removal during the South Asian summer monsoon season. North  
433 Africa and Central Asia also show high efficiencies resulted from less precipitation.  
434 Although East Asia does not have much precipitation in DJF, the efficiency is low  
435 because a large amount of sulfate is transported outside East Asia. It suggests that  
436 the lifetime of aerosols, mainly driven by wet deposition, together with regional export,  
437 is important in determining the local contribution to near-surface concentrations or  
438 regional air quality.

439

#### 440 **5. Source attribution of direct and indirect radiative forcing of sulfate**

441 The modeled global annual mean sulfate total DRF here is  $-0.42 \text{ W m}^{-2}$ , with –  
442  $0.31 \text{ W m}^{-2}$  contributed by anthropogenic sulfate and  $-0.11 \text{ W m}^{-2}$  contributed by  
443 natural sulfate (e.g., relative to a state with no natural emissions). The DRF of  
444 anthropogenic sulfate is  $-0.4 \pm 0.2 \text{ W m}^{-2}$  provided in the Fifth Assessment Report of



445 the Intergovernmental Panel on Climate Change (IPCC, 2013). Note that, the DRF of  
446 anthropogenic sulfate calculated here is total anthropogenic sulfate, whereas values  
447 from IPCC represent changes in anthropogenic sulfate between 1750 and  
448 present-day conditions, although this difference is small since 1750 SO<sub>2</sub> emissions  
449 are less than 1% of 2010 emissions. Spatial distributions of sulfate DRF, originating  
450 from the individual sixteen sources are shown in Fig. S6. The spatial distributions and  
451 global contributions of sulfate DRF are similar to those of sulfate column burden (Fig.  
452 S4), except that contribution of DMS to global sulfate DRF (18%) is much larger  
453 relative to its global column burden (11%). It is because DMS-produced sulfate  
454 burden is mostly located between 30°S–30°N (Fig. S4), where insolation is much  
455 stronger than at mid- and high latitudes, leading to stronger DRF over these regions.  
456 East Asia is the second largest contributor to global sulfate DRF, contributing 16% of  
457 global sulfate DRF, followed by 13% from ROW and 11% from South Asia.

458 Figure 9 shows seasonal and zonal mean DRF of sulfate originating from the  
459 tagged regions/sectors and the global total. The meridional distribution of DRF is  
460 jointly determined by many factors, e.g. sulfate loading, the insolation, cloud cover,  
461 and surface albedo. The total sulfate DRF shows a seasonal pattern that has the  
462 maximum DRF over 0°–10°N in DJF and over 30°–40°N in JJA, with values between  
463 –0.9 and –1.3 W m<sup>-2</sup>. Emissions originating from East Asia have the largest zonal  
464 mean sulfate DRF in almost all seasons, with a maximum around –0.45 W m<sup>-2</sup> in JJA  
465 because of the higher sulfate loading and the more abundant sunlight in JJA. South  
466 Asia also strongly contributes to sulfate DRF, followed by sources from ROW, VOL,



467 RBU, and the Middle East. DMS has the largest contribution over 60°S–0° in DJF due  
468 to the stronger insolation over the Southern Hemisphere in winter and more ocean  
469 DMS emission over these regions, and its DRF contribution is more widespread.  
470 Other tagged source regions have a relatively small contribution to the global total  
471 DRF, with a seasonal peak DRF less than  $-0.10 \text{ W m}^{-2}$ . The global and annual  
472 average sulfate DRF has a contribution of  $-0.074 \text{ W m}^{-2}$  from DMS,  $-0.068 \text{ W m}^{-2}$   
473 from East Asia,  $-0.054 \text{ W m}^{-2}$  from ROW,  $-0.047 \text{ W m}^{-2}$  from South Asia,  $-0.035 \text{ W}$   
474  $\text{m}^{-2}$  from VOL,  $-0.031 \text{ W m}^{-2}$  from the Middle East,  $-0.023 \text{ W m}^{-2}$  from Southern Africa,  
475  $-0.018 \text{ W m}^{-2}$  from Europe,  $-0.016 \text{ W m}^{-2}$  from North America, and a total of  $-0.057$   
476  $\text{W m}^{-2}$  from all other regions (Table S4).

477 Figure 10 shows seasonal fractional contributions to sulfate DRF in different  
478 latitudinal bands. Over the Southern Hemisphere tropics (30°S–Equator), mid-  
479 (60°S–30°S) and high (90°S–60°S) latitudes, DMS has the largest contribution to  
480 sulfate DRF in all seasons, with contribution about 17–84%. Sources from Southern  
481 Africa contribute about 11–20% of sulfate DRF over the Southern Hemisphere tropic  
482 and mid-latitudes, followed by about 10% from South America and ROW. Sources  
483 from East Asia account for 6–19% of sulfate DRF over the Southern Hemisphere high  
484 latitudes. In the Northern Hemisphere, influence from DMS becomes much weaker,  
485 but still substantial. Over the Northern Hemisphere tropics, East Asia, South Asia,  
486 ROW, and DMS exert equal contributions of 10–20%. East Asia has the largest  
487 contribution of 20–30% over the Northern Hemisphere mid- and high-latitudes,  
488 followed by South Asia and ROW.



489 Sulfate incremental IRF is estimated by using an additional simulation in which  
490 sulfur emissions are reduced by 20% for all regions and sectors. The difference in  
491 cloud radiative forcing between the control simulation and this second simulation gives  
492 the sulfate incremental IRF of the last 20% of sulfur emissions. Regional incremental  
493 IRF contributions are calculated by scaling the total incremental IRF in a grid column  
494 by regional source contributions to sulfate mass concentration reduction averaged  
495 from the surface layer to 850 hPa, which is the approximate altitude of cloud base.

496 Figure 11 shows regional contributions to sulfate incremental IRF from the tagged  
497 source regions/sectors. The sulfate incremental IRF is  $-0.44 \text{ W m}^{-2}$ . The spatial  
498 pattern is consistent with that of stratiform clouds since the model only considers  
499 aerosol effects on stratiform cloud. The strong negative forcing is mainly over oceans.  
500 All source contributions to sulfate incremental IRF from the fourteen tagged source  
501 regions are less than  $-0.04 \text{ W m}^{-2}$ , probably due to the polluted conditions over or near  
502 land. Particles originating from North America, South America, Southern Africa, and  
503 East Asia are also transported to ocean regions, leading to a strong negative forcing  
504 there. DMS has the largest contribution, explaining  $-0.23 \text{ W m}^{-2}$  of the global sulfate  
505 incremental IRF, because complex cloud adjustments are likely to respond sensitively  
506 to small changes in aerosol under clean conditions (Rosenfeld et al., 2014), followed  
507 by  $-0.06 \text{ W m}^{-2}$  from volcanic emissions. Note that the regional contribution to  
508 incremental IRF is simply calculated by decomposing the total incremental IRF with  
509 mass concentrations based on two simulations without and with the reduction in



510 emissions. This assumption could introduce biases considering non-linear relationship  
511 between mass concentration and IRF of sulfate.

512 To evaluate this new method for decomposing incremental IRF into different  
513 source regions/sector contributions, the IRF for one region (North America) and one  
514 sector (DMS) were calculated in a traditional manner using two additional simulations  
515 in which SO<sub>2</sub> emissions from North America and DMS emissions were reduced by  
516 20%, respectively. The incremental IRF calculated with the two methods are  
517 compared in Fig. S7. Although the incremental IRF outside the source regions  
518 obtained from the emission perturbation method is noisy, these two methods show  
519 similar negative incremental IRF within and near source regions. The 20% DMS leads  
520 to strong negative IRF over oceans and sources from North America result in negative  
521 IRF over Eastern U.S. and downwind ocean regions for both of these two methods.  
522 Globally, DMS and North America contribute to  $-0.231 (\pm 0.012)$  and  $-0.014 (\pm 0.002)$   
523  $W m^{-2}$ , respectively, of sulfate incremental IRF from the method with sulfur tagging  
524 technique, similar to  $-0.248 (\pm 0.020)$  and  $-0.018 (\pm 0.019) W m^{-2}$  from the individual  
525 emission-perturbation simulations.

526 Table S5 summarizes the DRF and incremental IRF of sulfate over land/ocean in  
527 the Northern/Southern Hemisphere contributed by the tagged source regions/sectors.  
528 Over the fourteen tagged source regions, the total anthropogenic source region  
529 contribution to DRF is  $-0.54/-0.18 W m^{-2}$  over land in the Northern/Southern  
530 Hemisphere, larger than  $-0.48/-0.12 W m^{-2}$  over ocean due to the larger sulfate  
531 burden near sources. Anthropogenic source contributions to incremental IRF are



532 larger over ocean, with values of  $-0.23/-0.13 \text{ W m}^{-2}$  compared to  $-0.08/-0.10 \text{ W m}^{-2}$   
533 over land in the Northern/Southern Hemisphere, because clouds are more  
534 susceptible to aerosol changes in clean environment and there are more stratiform  
535 clouds over ocean. For natural source sectors, their contributions are larger over  
536 oceans for both DRF and incremental IRF. Over land in the Northern Hemisphere,  
537 DRF is mainly driven by emissions from East Asia, South Asia, and the Middle East,  
538 whereas incremental IRF is dominated by emissions from North America, RBU and  
539 East Asia. The difference in major contributing regions for DRF vs. incremental IRF  
540 may be due to changes in cloud susceptibility when background aerosol  
541 concentrations are different. North America and RBU have more relatively clean  
542 areas (Alaska, N. Canada, parts of Siberia) than South Asia and East Asia, and  
543 clouds in the cleaner areas are more susceptible to the 20% emissions reductions.  
544 The non-linearity in DRF is much weaker, so the high emissions from South Asia and  
545 East Asia dominate DRF. Over ocean in the Northern Hemisphere, East Asia also  
546 contributes the largest to DRF and it is the second largest contributor to incremental  
547 IRF of sulfate following DMS. Over land in the Southern Hemisphere, emissions from  
548 Southern Africa and South America control DRF, whereas incremental IRF are  
549 largely attributed to sources from South America, DMS, and PAN  
550 (Pacific/Australia/New Zealand). Over ocean in the Southern Hemisphere, both  
551 sulfate DRF and incremental IRF are dominated by DMS emissions.

552 Figure 12 shows the seasonal and annual global DRF and incremental IRF  
553 efficiencies of sulfate. (Table S6 gives values.) Global DRF efficiency of a source



554 region is defined as the global DRF of sulfate originating from the source  
555 region/sector divided by the total sulfur emissions from that region/sector. The global  
556 DRF efficiency treats the whole globe as a receptor region, as opposed to a specific  
557 region in the regional concentration efficiency definition, considering that aerosol  
558 climatic impacts are on a global scale whereas air quality impacts are more important  
559 on a regional scale. As the DRF is more closely related to sulfate burden, global  
560 sulfate burden efficiencies are also provided in Table S7. The global DRF efficiency  
561 for total sulfur emissions is  $-4.8 \text{ mW m}^{-2} (\text{Tg S yr}^{-1})^{-1}$ . The Middle East, North Africa,  
562 and Southern Africa present high DRF efficiencies, as a result of both long aerosol  
563 lifetime and strong tropical insolation. These source regions also have high global  
564 burden efficiencies.

565 Table S6 compares the annual DRF efficiencies simulated in this study with the  
566 average from a previous multi-model study (Stjern et al., 2016). In Stjern et al. (2016),  
567 efficiency was calculated as the response of global DRF to a 20% reduction in local  
568 emissions divided by the 20% of total  $\text{SO}_2$  emissions based on two separate  
569 simulations, whereas the efficiency in this present study is calculated as the global  
570 contribution of DRF divided by 100% of local emissions in a single simulation, taking  
571 advantage of the sulfur tagging capability and radiation diagnostic calculations. Both  
572 studies show high efficiencies over the Middle East and South Asia, and low  
573 efficiencies over North America, Europe, East Asia and RBU. In addition, the  
574 magnitude of efficiencies in this study are very similar to Stjern et al. (2016), with  
575 differences less than 15%.



576 The global IRF efficiency of a source region is calculated as the global  
577 contribution of sulfate incremental IRF divided by the changes (i.e., 20% reduction) in  
578 sulfur emissions in that region. Unlike the DRF efficiencies, IRF efficiencies are  
579 higher over or near ocean regions, with a global IRF efficiency of  $-5.0 \text{ mW m}^{-2} (\text{Tg S}$   
580  $\text{yr}^{-1})^{-1}$  for the global total 20% of sulfur emissions. PAN and DMS have the largest IRF  
581 efficiencies because PAN has a relatively clean environment compared to other  
582 regions and DMS is emitted over clean oceans. Cloud properties are more  
583 susceptible to aerosol perturbations in a more pristine environment. Although the  
584 background aerosols in South America are not so low, sulfate originating from this  
585 region has a large contribution to sulfate over oceans of the Southern Hemisphere,  
586 explaining a large IRF efficiency from that region.

587 In addition to the incremental IRF and efficiency, we also calculated the  
588 anthropogenic sulfate IRF and its efficiency between present-day and preindustrial  
589 conditions with an additional simulation, in which anthropogenic  $\text{SO}_2$  emissions are  
590 fixed at the 1850 level, and compared these values with those from the 20% sulfur  
591 emission reduction simulation in Table S8. The modeled annual and global mean  
592 anthropogenic sulfate IRF here is  $-0.74 \text{ W m}^{-2}$ , which is comparable to  $-0.45 \pm 0.5 \text{ W}$   
593  $\text{m}^{-2}$  of IRF for total anthropogenic aerosols from IPCC (2013). The anthropogenic IRF  
594 contributed from individual source regions is about 3–6 times larger than the  
595 incremental IRF, in agreement with about 5 times more reduction in  $\text{SO}_2$  emissions in  
596 the preindustrial simulation than in the 20% sulfur emission reduction simulation. The  
597 forcing efficiencies are roughly similar between the incremental and the



598 anthropogenic IRF, indicating a nearly linear relationship between SO<sub>2</sub> emission and  
599 sulfate IRF, except for the Middle East and South Asia where concentrated dust and  
600 its variability may strongly influence cloud properties and therefore sulfate IRF. Figure  
601 S8 shows the anthropogenic sulfate IRF efficiencies that are calculated based on  
602 anthropogenic IRF from the present-day and preindustrial condition simulations. The  
603 values are similar to the incremental IRF efficiencies, further validating the robust  
604 results from the decomposed regional IRF with the sulfur tagging technique.

605

## 606 **6. Conclusions and discussions**

607 A sulfur tagging technique is implemented in Community Atmosphere Model  
608 (CAM) of the Community Earth System Model (CESM) and used in this study to  
609 examine source-receptor relationships of sulfate concentrations, DRF and IRF  
610 originating from sixteen regions/sectors (North America, Central America, South  
611 America, Europe, North Africa, Southern Africa, the Middle East, Southeast Asia,  
612 Central Asia, South Asia, East Asia, RBU, PAN, ROW, VOL, and DMS) for 2010–  
613 2014. The anthropogenic emissions came from the CEDS inventory developed for  
614 the CMIP6.

615 Near-surface sulfate concentrations are mostly contributed by local emissions in  
616 regions with high emissions, such as Eastern U.S., Southern Africa, South Asia, and  
617 Eastern China, where local source contributions exceed 80%. Over regions with  
618 relatively low SO<sub>2</sub> emissions, the near-surface sulfate concentrations are primarily  
619 attributed to non-local sources from long-range transport.



620 The source-receptor relationships have strong seasonal variations. The export of  
621 sulfate from Europe contributes to 16–20% of near-surface sulfate concentrations  
622 over North Africa, RBU and Central Asia in all seasons. Sulfate from the Middle East  
623 is effectively transported to the surrounding receptor regions and accounts for 15–24%  
624 of sulfate concentrations over North Africa, Southern Africa and Central Asia in DJF  
625 and SON, and 19% over South Asia in MAM. Sources in RBU account for 21–42% of  
626 sulfate concentrations over Central Asia, with a peak contribution in JJA. Northerly  
627 winds over East Asia in DJF and SON associated with East Asian winter monsoon  
628 transport sulfate from highly polluted Eastern China to Southeast Asia, accounting for  
629 about 50% of near-surface sulfate concentrations over Southeast Asia. East Asia  
630 also contributes 15% to the near-surface sulfate over RBU in JJA and 11% over North  
631 America in MAM. The transport of sulfate from South Asia contributes 11–24% of  
632 near-surface sulfate over Southeast Asia in DJF and MAM. Regional sulfate  
633 concentration efficiencies are higher over regions with dry atmospheric conditions  
634 and less export, suggesting that the lifetime of aerosols mainly driven by wet  
635 deposition, together with regional export, is important in determining the regional air  
636 quality.

637 The simulated global total sulfate DRF is  $-0.42 \text{ W m}^{-2}$ , with  $-0.31 \text{ W m}^{-2}$   
638 contributed by anthropogenic sulfate and  $-0.11 \text{ W m}^{-2}$  contributed by natural sulfate.  
639 DMS has the largest contribution to the global sulfate DRF, followed by East Asia,  
640 ROW and South Asia. In the Southern Hemisphere, DMS contributes 17–84% to the  
641 seasonal total sulfate DRF. In the Northern Hemisphere tropics, East Asia, South



642 Asia, ROW, and DMS exert similar contributions of 10–20%. East Asia has the  
643 largest contribution of 20–30% over the Northern Hemisphere mid- and high-latitudes,  
644 followed by South Asia and ROW.

645 Sulfate incremental IRF is estimated using an additional simulation in which sulfur  
646 emissions are reduced by 20%. The difference in cloud radiative forcing between the  
647 control simulation and this second simulation gives the sulfate incremental IRF of the  
648 last 20% of sulfur emissions, which is  $-0.44 \text{ W m}^{-2}$  globally. DMS has the largest  
649 contribution, explaining  $-0.23 \text{ W m}^{-2}$  of the global sulfate incremental IRF, because of  
650 the clean marine background conditions, followed by  $-0.06 \text{ W m}^{-2}$  from volcanic  
651 emissions.

652 The Middle East, North Africa, and Southern Africa have high global DRF  
653 efficiencies, due to both longer aerosol lifetimes (from low precipitation) and strong  
654 insolation. Regions in the Southern Hemisphere with low background aerosols have  
655 stronger global IRF efficiencies than those over the polluted Northern Hemisphere,  
656 because cloud properties are more susceptible to aerosol perturbations in a more  
657 pristine environment.

658 Note that, although simulated near-surface sulfate concentrations are in  
659 agreement with observed values at the IMPROVE sites over North America and at  
660 the EANET sites over part of East Asia and Southeast Asia, the model strongly  
661 underestimates sulfate concentrations by  $-54\%$  in China, compared to site  
662 observations from the CAWNET network. Comparison of column-integrated  $\text{SO}_2$   
663 between model simulation and OMI satellite data shows a possible overestimation of



664 SO<sub>2</sub> in the model. The simulated SO<sub>2</sub> near-surface concentrations, however, are  
665 underestimated by 25% compared to observations over thirteen sites in China,  
666 suggesting a large bias in satellite retrievals or too much SO<sub>2</sub> simulated at higher  
667 altitudes. The model SO<sub>2</sub> concentrations over downwind regions of China are  
668 underestimated by 45%, indicating that the transport of SO<sub>2</sub> from China is probably  
669 underestimated in the model. A less efficient transformation from SO<sub>2</sub> to sulfate could  
670 also lead to the underestimation of sulfate in the model. The underestimation of sulfate  
671 over China could lead to the underestimation of contributions from East Asia to remote  
672 sulfate concentrations, global DRF and incremental IRF, as well as their efficiencies.

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677 *Data availability.* All the emissions datasets used in this study can be obtained from  
678 <https://pcmdi.llnl.gov/projects/input4mips>. The sulfate datasets are available from  
679 <http://vista.cira.colostate.edu/IMPROVE/> for IMPROVE sites, <http://www.eanet.asia>  
680 for EANET sites, and <http://www.emep.int> for EMEP sites. The OMI satellite-derived  
681 total column burden of SO<sub>2</sub> can be downloaded from  
682 [http://disc.sci.gsfc.nasa.gov/Aura/data-holdings/OMI/omso2e\\_v003.shtml](http://disc.sci.gsfc.nasa.gov/Aura/data-holdings/OMI/omso2e_v003.shtml). The  
683 CESM model is publically available at <http://www.cesm.ucar.edu/models/cesm1.2/>.  
684 Our model results can be made available through the National Energy Research  
685 Scientific Computing Center (NERSC) servers upon request.



686

687 *Competing interests.* The authors declare that they have no conflict of interest.

688

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947 **Table 1.** Seasonal emissions (units: Tg S season<sup>-1</sup>) of combustion (anthropogenic +  
948 biomass burning) SO<sub>2</sub> and DMS from the sixteen source regions/sectors in  
949 December-January-February (DJF), March-April-May (MAM), June-July-August (JJA),  
950 and September-October-November (SON) and annual total emissions (ANN).  
951

	NAM	CAM	SAM	EUR	NAF	SAF	MDE	SEA
DJF	8.313E-01	3.458E-01	3.284E-01	1.073E+00	1.519E-01	6.507E-01	8.388E-01	3.537E-01
MAM	7.016E-01	3.659E-01	3.677E-01	8.251E-01	1.529E-01	5.871E-01	8.421E-01	3.731E-01
JJA	8.761E-01	3.731E-01	4.740E-01	6.456E-01	1.534E-01	8.090E-01	8.398E-01	3.516E-01
SON	7.045E-01	3.550E-01	4.357E-01	7.829E-01	1.518E-01	6.641E-01	8.353E-01	3.517E-01
ANN	3.114E+00	1.440E+00	1.606E+00	3.327E+00	6.099E-01	2.711E+00	3.356E+00	1.430E+00
	CAS	SAS	EAS	RBU	PAN	ROW	VOL	DMS
DJF	3.156E-01	1.593E+00	5.043E+00	8.913E-01	1.266E-01	2.836E+00	3.106E+00	5.991E+00
MAM	2.720E-01	1.626E+00	4.406E+00	7.443E-01	1.352E-01	2.775E+00	3.175E+00	4.770E+00
JJA	2.300E-01	1.605E+00	4.084E+00	6.455E-01	1.597E-01	2.739E+00	3.175E+00	3.537E+00
SON	2.619E-01	1.594E+00	4.299E+00	6.940E-01	1.625E-01	2.813E+00	3.141E+00	3.918E+00
ANN	1.080E+00	6.418E+00	1.783E+01	2.975E+00	5.840E-01	1.116E+01	1.260E+01	1.822E+01

952

953



954 **Figure Captions**

955

956 **Figure 1.** (a) Tagged source regions (NAM: North America, CAM: Central America,  
957 SAM: South America, EUR: Europe, NAF: North Africa, SAF: Southern Africa, MDE:  
958 the Middle East, SEA: Southeast Asia, CAS: Central Asia, SAS: South Asia, EAS:  
959 East Asia, RBU: Russia/Belarus/Ukraine, PAN: Pacific/Australia/New Zealand and  
960 ROW: rest of the world) and (b) the respective percentage contributions to global  
961 annual mean combustion SO<sub>2</sub> emissions (anthropogenic + biomass burning) from the  
962 individual source regions.

963

964 **Figure 2.** Spatial distribution of annual mean emissions (g S m<sup>-2</sup> yr<sup>-1</sup>) of  
965 anthropogenic SO<sub>2</sub>, volcanic SO<sub>2</sub>, and DMS from the sixteen tagged source  
966 regions/sectors averaged over 2010–2014.

967

968 **Figure 3.** Spatial distribution (left panel) and scatter plot (right) between the simulated  
969 and observed annual mean near-surface sulfate concentrations (µg m<sup>-3</sup>) over years  
970 2010–2014. Observations are from IMPROVE (up pointing triangle), EMEP (square),  
971 EANET (down pointing triangle) for years 2010–2014, and CAWNET (circle) for years  
972 2006–2007, which are scaled to 2010–2014 based on the ratio of CEDS 2010-2014  
973 SO<sub>2</sub> emissions to 2006-2007 emissions over China (which is 0.92). Solid lines mark  
974 the 1:1 ratio and dashed lines mark the 1:2 and 2:1 ratio. Normalized mean bias  
975 (NMB) and correlation coefficient (R) between observation and simulation are shown



976 on the right panel.  $NMB = 100\% \times \sum (M_i - O_i) / \sum O_i$ , where  $M_i$  and  $O_i$  are the  
977 modeled and observed values at site  $i$ , respectively.

978

979 **Figure 4.** Spatial distribution of relative contributions (%) to annual mean  
980 near-surface sulfate concentrations from each of the tagged source regions/sectors.  
981 Relative contributions to global averaged sulfate from individual source  
982 regions/sectors is shown at the bottom right of each panel.

983

984 **Figure 5.** Relative contributions of non-local sources to seasonal near-surface sulfate  
985 concentrations (left panels) and wind fields over 850 hPa (right panels). Arrows with  
986 numbers show contributions (%) of a source region to sulfate over a receptor region.  
987 Only relative concentrations larger than 10% are shown.

988

989 **Figure 6.** Relative contributions (%) of local emissions (inside the tagged regions) to  
990 near-surface sulfate concentrations. Contributions from natural source sectors are  
991 added to ROW here. Contributions less than 50% are shown in cold colors and those  
992 larger than 50% are shown in warm colors.

993

994 **Figure 7.** Relative contributions (%) to near-surface sulfate concentrations averaged  
995 over land and ocean of the Northern and Southern Hemisphere from emissions in the  
996 sixteen tagged source regions/sectors.

997



998 **Figure 8.** Seasonal and annual mean regional concentration efficiency of sulfate ( $\mu\text{g}$   
999  $\text{m}^{-3}$  ( $\text{Tg S yr}^{-1}$ ) $^{-1}$ ) of the sixteen tagged source regions/sectors. The efficiency is  
1000 defined as the local contribution to near-surface sulfate concentration divided by the  
1001 corresponding sulfur emissions from that region (seasonal emissions multiplied by 4).  
1002 Error bars indicate 1- $\sigma$  of mean values during years 2010–2014. The receptor region  
1003 of ROW is used to calculate efficiency of VOL and DMS.

1004

1005 **Figure 9.** Contributions to zonal mean sulfate direct radiative forcing ( $\text{W m}^{-2}$ ) from  
1006 emissions of the tagged regions/sectors shown in colors (left Y axis) and from global  
1007 total emissions shown in black (right Y axis). Only regions with maximum of zonal  
1008 mean sulfate direct radiative forcing stronger than  $-0.1 \text{ W m}^{-2}$  are shown here.

1009

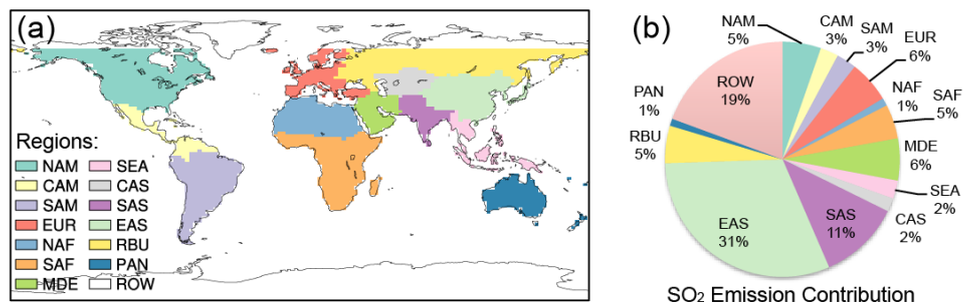
1010 **Figure 10.** Relative contributions (%) from emissions in the sixteen tagged  
1011 regions/sectors to sulfate direct radiative forcing over the Southern Hemisphere  
1012 high-latitudes ( $90^{\circ}\text{S}$ – $60^{\circ}\text{S}$ ), Southern Hemisphere mid-latitudes ( $60^{\circ}\text{S}$ – $30^{\circ}\text{S}$ ),  
1013 Southern Hemisphere tropics ( $30^{\circ}\text{S}$ –Equator), Northern Hemisphere tropics  
1014 (Equator– $30^{\circ}\text{N}$ ), Northern Hemisphere mid-latitudes ( $30^{\circ}\text{N}$ – $60^{\circ}\text{N}$ ), and Northern  
1015 Hemisphere high-latitudes ( $60^{\circ}\text{N}$ – $90^{\circ}\text{N}$ ).

1016

1017 **Figure 11.** Spatial distribution of responses of annual mean indirect radiative forcing  
1018 of sulfate (IRF,  $\text{W m}^{-2}$ ) to a 20% reduction in sulfur emissions (standard simulation –  
1019 simulation with 20% emission reduction). Regional contributions are calculated as a

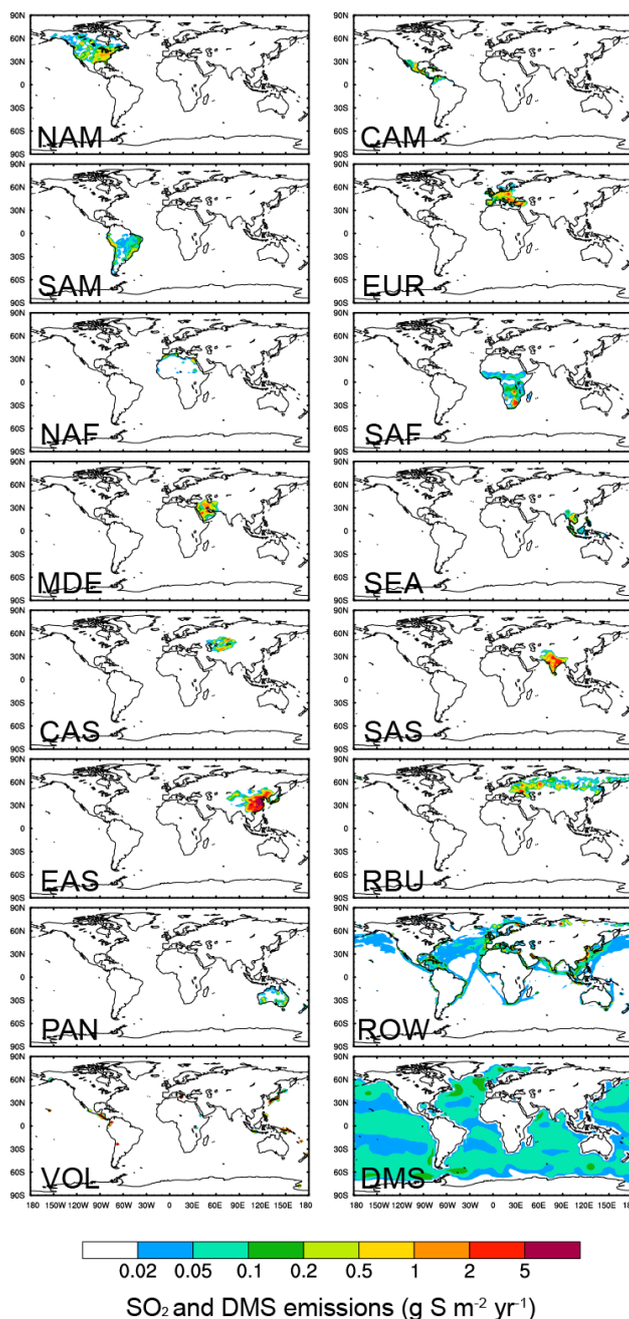


1020 scaled total incremental IRF in each grid cell by the ratio of source contribution to total  
1021 sulfate mass concentration reduction averaged from the surface layer to 850 hPa.  
1022 Regional mean contributions to global incremental IRF of sulfate are shown at the  
1023 bottom right of each panel.  
1024  
1025 **Figure 12.** Seasonal and annual mean global sulfate (a) direct and (b) indirect  
1026 radiative forcing efficiency ( $\text{mW m}^{-2} (\text{Tg S yr}^{-1})^{-1}$ ) of the sixteen tagged source  
1027 regions/sectors. The sulfate radiative efficiency is defined as the global sulfate  
1028 radiative forcing divided by the corresponding scaled annual sulfur emission  
1029 (seasonal emission multiplied by 4). Error bars indicate  $1\text{-}\sigma$  of mean values during  
1030 years 2010–2014.



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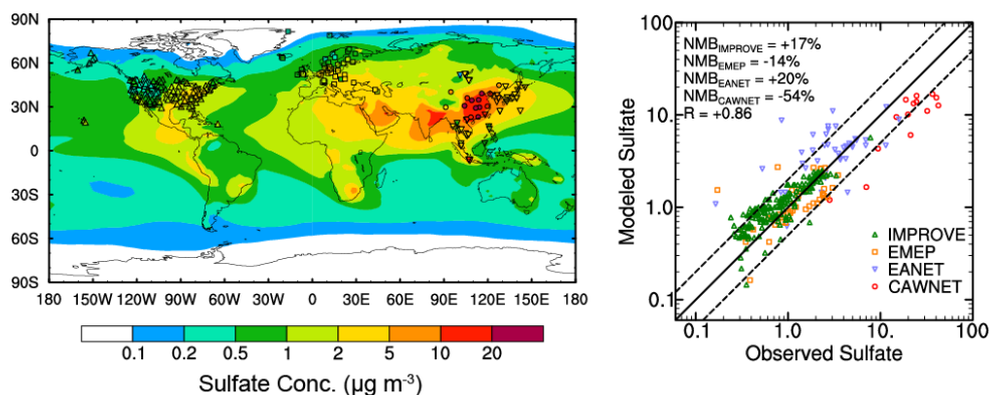
**Figure 1.** (a) Tagged source regions (NAM: North America, CAM: Central America, SAM: South America, EUR: Europe, NAF: North Africa, SAF: Southern Africa, MDE: the Middle East, SEA: Southeast Asia, CAS: Central Asia, SAS: South Asia, EAS: East Asia, RBU: Russia/Belarus/Ukraine, PAN: Pacific/Australia/New Zealand and ROW: rest of the world) and (b) the respective percentage contributions to global annual mean combustion SO<sub>2</sub> emissions (anthropogenic + biomass burning) from the individual source regions.



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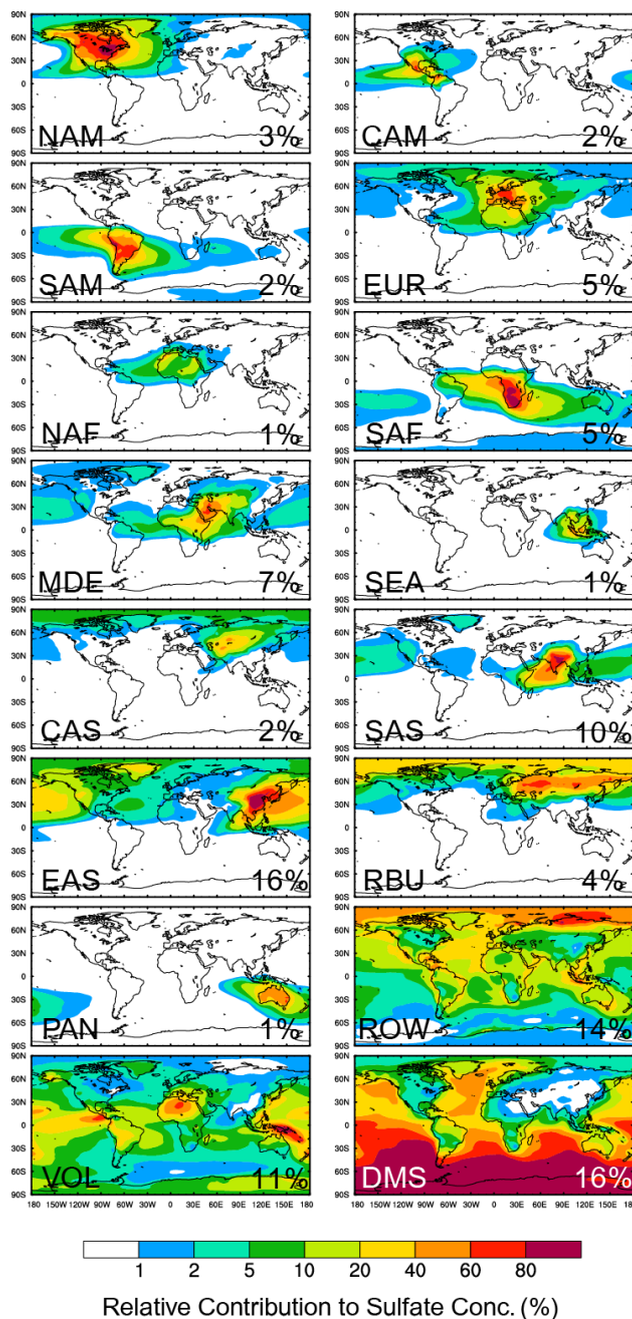
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1043 **Figure 2.** Spatial distribution of annual mean emissions (g S m<sup>-2</sup> yr<sup>-1</sup>) of  
1044 anthropogenic SO<sub>2</sub>, volcanic SO<sub>2</sub>, and DMS from the sixteen tagged source  
1045 regions/sectors averaged over 2010–2014.



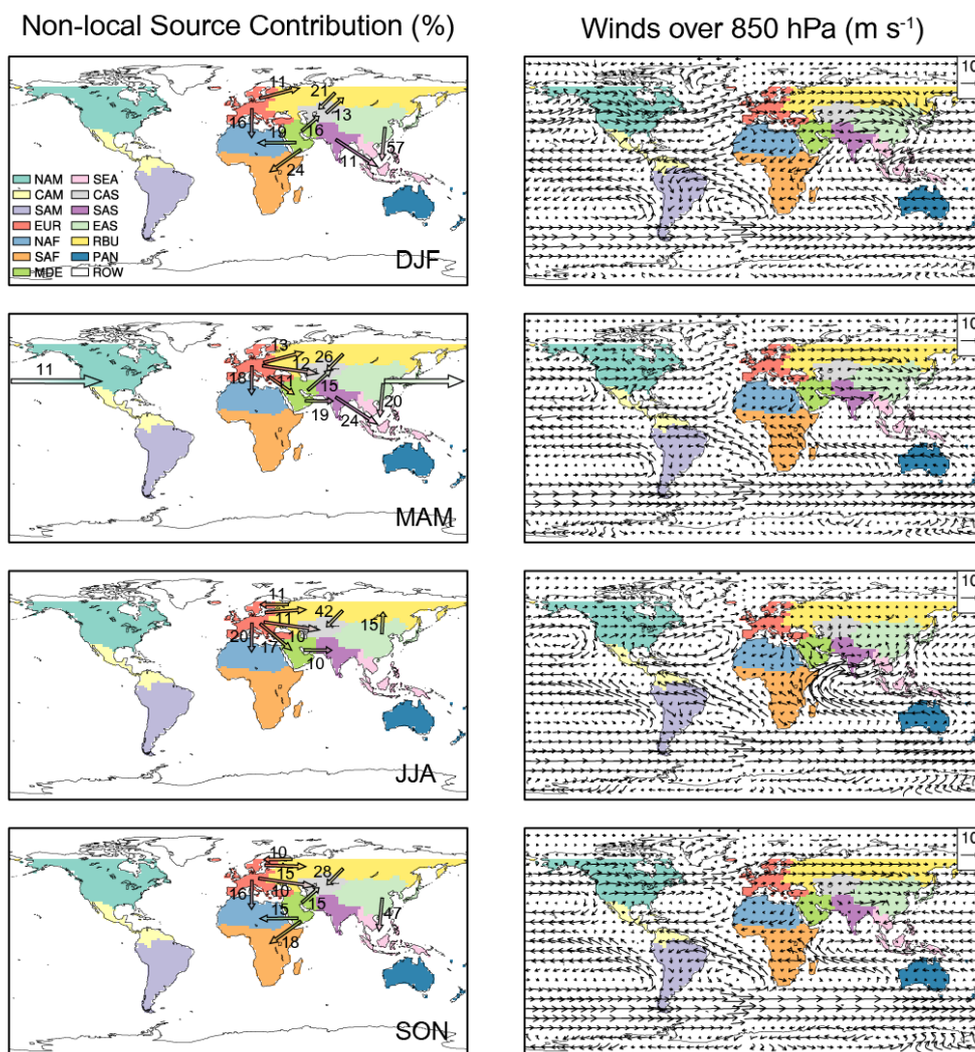
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**Figure 3.** Spatial distribution (left panel) and scatter plot (right) between the simulated and observed annual mean near-surface sulfate concentrations ( $\mu\text{g m}^{-3}$ ) over years 2010–2014. Observations are from IMPROVE (up pointing triangle), EMEP (square), EANET (down pointing triangle) for years 2010–2014, and CAWNET (circle) for years 2006–2007, which are scaled to 2010–2014 based on the ratio of CEDS 2010–2014  $\text{SO}_2$  emissions to 2006–2007 emissions over China (which is 0.92). Solid lines mark the 1:1 ratio and dashed lines mark the 1:2 and 2:1 ratio. Normalized mean bias (NMB) and correlation coefficient (R) between observation and simulation are shown on the right panel.  $\text{NMB} = 100\% \times \sum(M_i - O_i) / \sum O_i$ , where  $M_i$  and  $O_i$  are the modeled and observed values at site  $i$ , respectively.



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**Figure 4.** Spatial distribution of relative contributions (%) to annual mean near-surface sulfate concentrations from each of the tagged source regions/sectors. Relative contributions to global averaged sulfate from individual source regions/sectors is shown at the bottom right of each panel.



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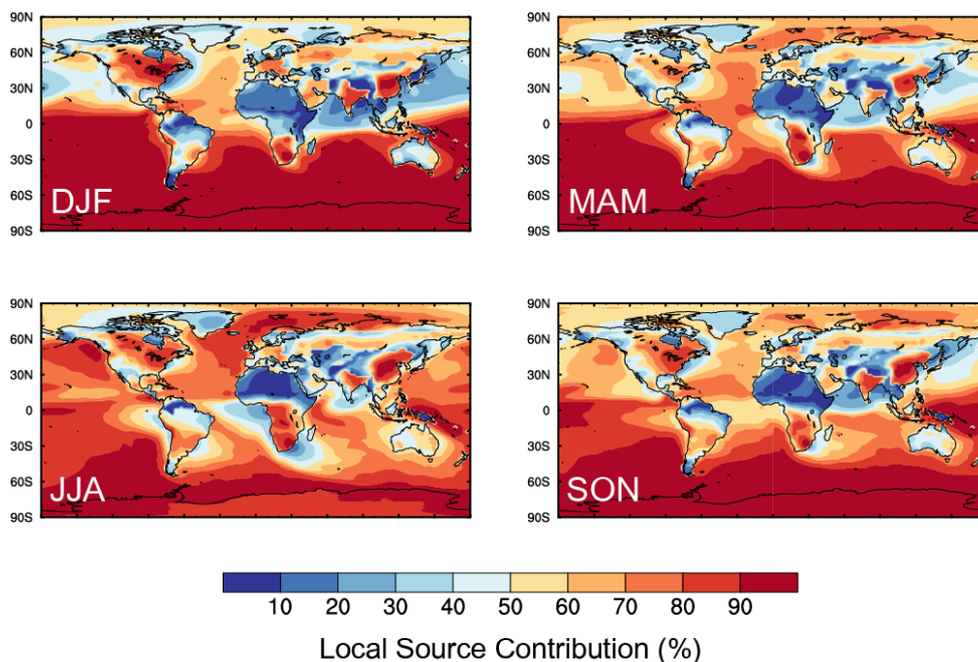
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1067 **Figure 5.** Relative contributions of non-local sources to seasonal near-surface sulfate

1068 concentrations (left panels) and wind fields over 850 hPa (right panels). Arrows with

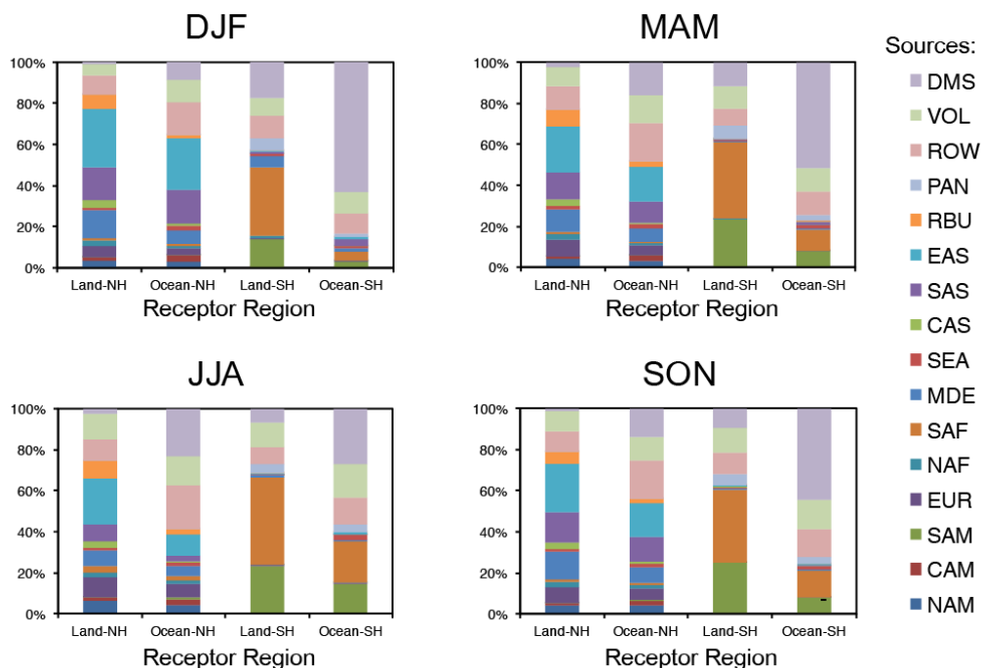
1069 numbers show contributions (%) of a source region to sulfate over a receptor region.

1070 Only relative concentrations larger than 10% are shown.



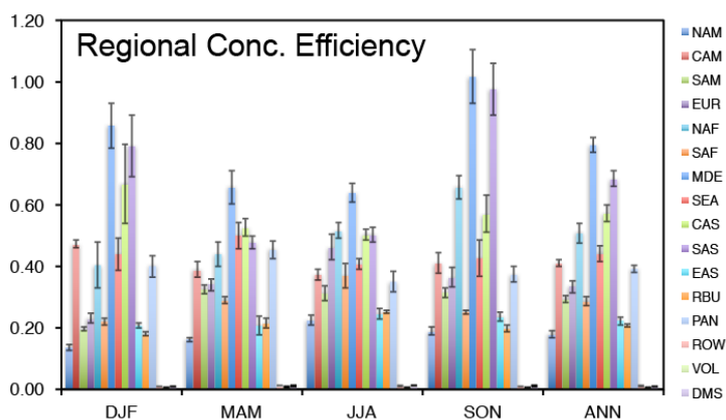
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**Figure 6.** Relative contributions (%) of local emissions (inside the tagged regions) to near-surface sulfate concentrations. Contributions from natural source sectors are added to ROW here. Contributions less than 50% are shown in cold colors and those larger than 50% are shown in warm colors.



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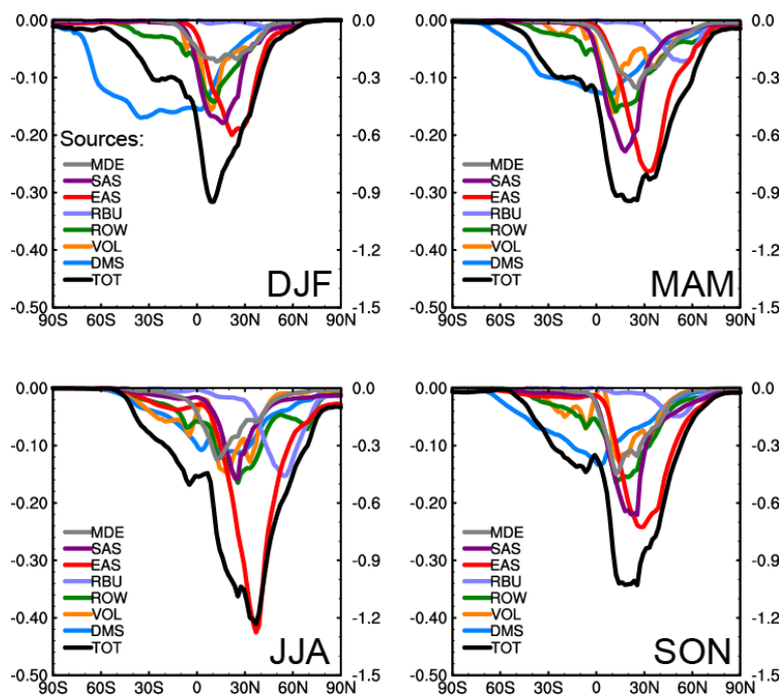
**Figure 7.** Relative contributions (%) to near-surface sulfate concentrations averaged over land and ocean of the Northern and Southern Hemisphere from emissions in the sixteen tagged source regions/sectors.



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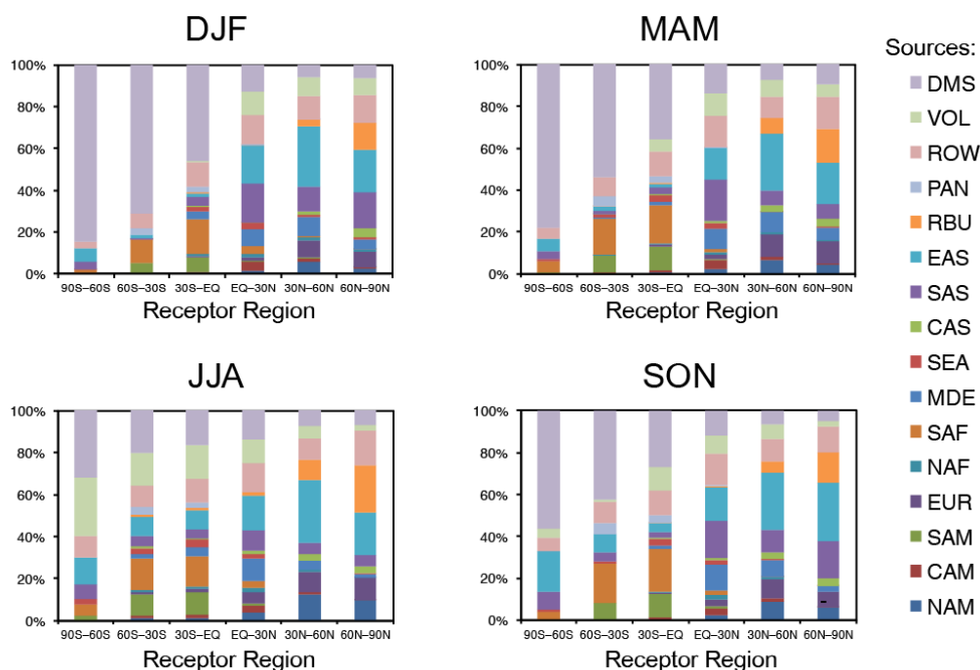
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1087 **Figure 8.** Seasonal and annual mean regional concentration efficiency of sulfate ( $\mu\text{g}$   
1088  $\text{m}^{-3}$  ( $\text{Tg S yr}^{-1}$ )<sup>-1</sup>) of the sixteen tagged source regions/sectors. The efficiency is  
1089 defined as the local contribution to near-surface sulfate concentration divided by the  
1090 corresponding sulfur emissions from that region (seasonal emissions multiplied by 4).  
1091 Error bars indicate 1- $\sigma$  of mean values during years 2010–2014. The receptor region  
1092 of ROW is used to calculate efficiency of VOL and DMS.  
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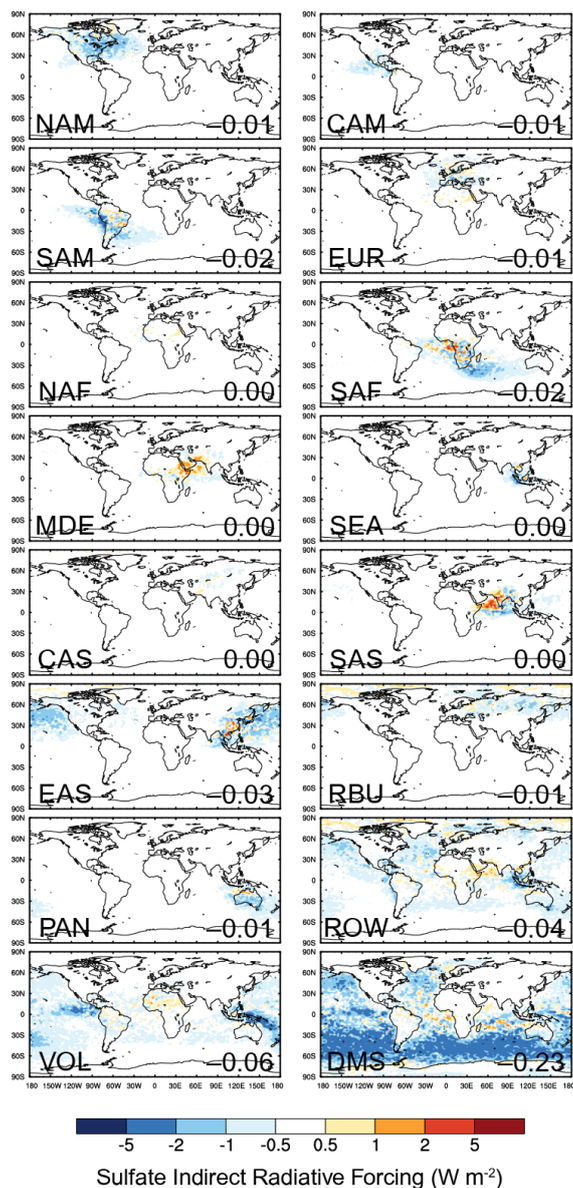
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**Figure 9.** Contributions to zonal mean sulfate direct radiative forcing ( $\text{W m}^{-2}$ ) from emissions of the tagged regions/sectors shown in colors (left Y axis) and from global total emissions shown in black (right Y axis). Only regions with maximum of zonal mean sulfate direct radiative forcing stronger than  $-0.1 \text{ W m}^{-2}$  are shown here.



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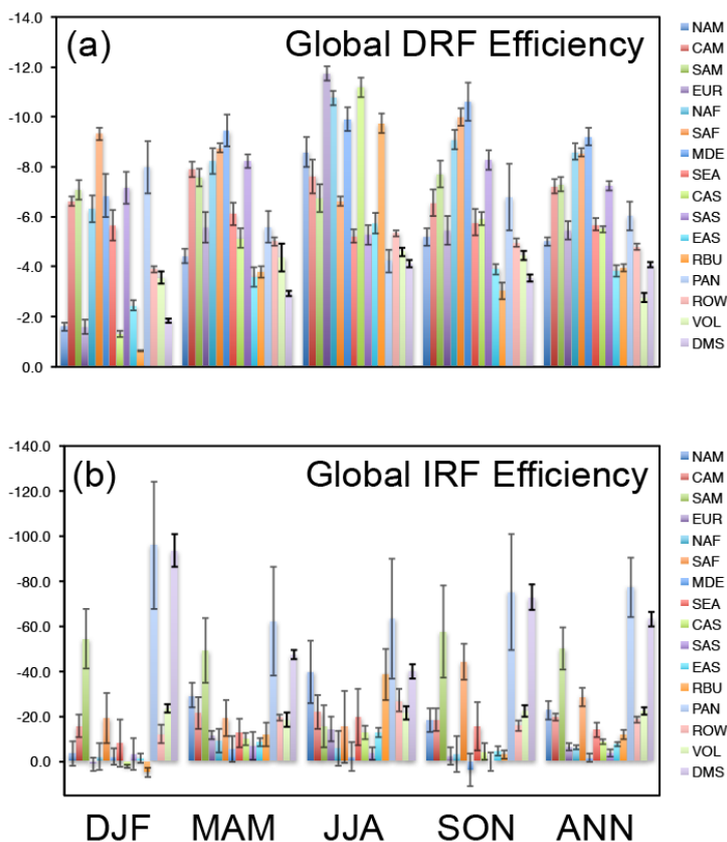
**Figure 10.** Relative contributions (%) from emissions in the sixteen tagged regions/sectors to sulfate direct radiative forcing over the Southern Hemisphere high-latitudes (90°S–60°S), Southern Hemisphere mid-latitudes (60°S–30°S), Southern Hemisphere tropics (30°S–Equator), Northern Hemisphere tropics (Equator–30°N), Northern Hemisphere mid-latitudes (30°N–60°N), and Northern Hemisphere high-latitudes (60°N–90°N).



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1115 **Figure 11.** Spatial distribution of responses of annual mean indirect radiative forcing  
 1116 of sulfate (IRF,  $\text{W m}^{-2}$ ) to a 20% reduction in sulfur emissions (standard simulation –  
 1117 simulation with 20% emission reduction). Regional contributions are calculated as a  
 1118 scaled total incremental IRF in each grid cell by the ratio of source contribution to total  
 1119 sulfate mass concentration reduction averaged from the surface layer to 850 hPa.  
 1120 Regional mean contributions to global incremental IRF of sulfate are shown at the  
 1121 bottom right of each panel.



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1124 **Figure 12.** Seasonal and annual mean global sulfate (a) direct and (b) indirect  
1125 radiative forcing efficiency ( $\text{mW m}^{-2} (\text{Tg S yr}^{-1})^{-1}$ ) of the sixteen tagged source  
1126 regions/sectors. The sulfate radiative efficiency is defined as the global sulfate  
1127 radiative forcing divided by the corresponding scaled annual sulfur emission  
1128 (seasonal emission multiplied by 4). Error bars indicate 1- $\sigma$  of mean values during  
1129 years 2010–2014.