Impacts on cloud radiative effects induced by coexisting aerosols converted from international shipping and maritime DMS emissions

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Abstract. International shipping emissions (ISE), particularly sulfur dioxide, can influence the global radiation budget by interacting with clouds and radiation after being oxidized into sulfate aerosols. A better understanding of the uncertainties in estimating the cloud radiative effects (CRE) of ISE is of great importance in climate science. Many international shipping tracks cover oceans with substantial natural dimethyl sulfide (DMS) emissions. The interplay between these two major aerosol sources on cloud radiative effects over vast oceanic regions with relatively low aerosol concentration is an intriguing yet poorly addressed issue confounding estimation of the cloud radiative effects of ISE. Using an Earth system model including two aerosol modules with different aerosol mixing configurations, we derive a significant global net CRE of ISE (−0.153 W m⁻² with standard error of ±0.004 W m⁻²) when using emissions consistent with current ship emission regulations. This global net CRE would become much weaker and actually insignificant (−0.001 W m⁻² standard error of ±0.007 W m⁻²) if a more stringent regulation were adopted. We then reveal that the ISE-induced CRE would achieve a significant enhancement when lower DMS emission is prescribed in the simulations, owing to the sub-linear relationship between aerosol concentration and cloud response. In addition, this study also demonstrates that the representation of certain aerosol processes, such as mixing states, can influence the magnitude and pattern of the ISE-induced CRE. These findings suggest a re-evaluation of the ISE-induced CRE with consideration of DMS variability.

1 Introduction

Marine stratiform clouds have a strong cooling effect on the climate system. They cover about 30% of the global ocean surface (Warren et al., 1988), and can reflect more solar radiation back to space than the dark ocean surface at cloud-free conditions. On the other hand, low-altitude marine stratiform clouds form and develop near to the ocean surface (only several degrees cooler than ocean surface) and thus have limited impacts on the longwave radiation balance (Klein and Hartmann, 1993). Therefore, the annual-mean net radiative effect of cloud at the top of the atmosphere (TOA) is negative (i.e., cooling) and can be up to −20 W m⁻² on the global scale (Boucher et al., 2013). Consequently, even a few percent change in marine stratocumulus cloud cover can double or offset the anthropogenic global warming due to greenhouse gases.

Sulfate aerosols are efficient cloud condensation nuclei and control the formation of marine clouds and their micro- and macro-physical properties (McCoy et al., 2015). The international shipping-emitted sulfur dioxide from combustion of heavy fossil oil (Figure 1) can be oxidized to sulfate aerosols that can increase cloud droplet number concentrations, cloud liquid water path, and planetary albedo, resulting in more solar radiation being reflected back to space, exerting a cooling effect on the climate system (Capaldo et al., 1999; Devasthale et al., 2006; Lauer et al., 2007; Lauer et al., 2009). Although international shipping emissions (ISE) contribute only about 5% (5.4 Tg S yr⁻¹) to the total anthropogenic sulfur emissions (Corbett and Koehler, 2003; Endresen et al., 2005; Klimont et al., 2013), they dominate the sulfur concentration across much of the ocean,
such as the North Pacific Ocean (NPO) and the North Atlantic Ocean (NAO), as shown in Figure 2. Therefore, the radiative impact of the ISE via perturbing marine stratocumulus clouds could be large—especially because marine stratocumulus clouds are often collocated with busy shipping lanes (Neubauer et al., 2014). Note that although ISE also contain significant amount of black carbon and organic carbon aerosols, since this study mainly focuses on aerosol induced CRE instead of aerosol direct radiative effect, we focus on only primary and secondary sulfate aerosols due to their much higher hygroscopicity than those of black carbon and organic carbon aerosols (Pringle et al., 2010).

The estimated global annual mean of the ISE-induced net cloud radiative effect (CRE) at TOA has large uncertainties due to the complication in simulating clouds and aerosol–cloud interactions, ranging from −0.60 to −0.07 W m\(^{-2}\) (Capaldo et al., 1999; Lauer et al., 2007; Eyring et al., 2010; Righi et al., 2011; Peters et al., 2012; Partanen et al., 2013; Peters et al., 2013). Compared with net CRE, the net direct radiative effect (DRE) of shipping emissions is much weaker, with a magnitude of only −0.08 to −0.01 W m\(^{-2}\), approximately one tenth of the former (Endresen et al., 2003; Schreier et al., 2007; Eyring et al., 2010).

Besides shipping-emitted sulfur compounds, oceanic phytoplankton-derived dimethyl sulfide (DMS) is another significant component in the atmospheric sulfur cycle over oceans (Grandey and Wang, 2015; Mahajan et al., 2015; McCoy et al., 2015; Tesdal et al., 2016). DMS can be oxidized by hydroxyl radical or nitrate radical to produce sulfur dioxide and finally converted to sulfate aerosols (Boucher et al., 2003). The global total DMS emission is estimated to range from 8 to 51 Tg S yr\(^{-1}\) based on model simulation (Quinn et al., 1993; Dentener et al., 2006); this uncertainty range is itself substantially larger than the total sulfur emissions from shipping. The global annual mean of the DMS-induced net CRE at TOA ranges from −2.03 to −1.49 W m\(^{-2}\) determined by DMS climatology (Gunson et al., 2006; Thomas et al., 2010; Mahajan et al., 2015).

Most of the aforementioned studies addressed separately the impacts on CRE of shipping and DMS emissions, largely ignoring the potential nonlinearity in the response of cloud radiative effects to aerosol variations when these sulfate aerosols from two different sources often collocate in the marine atmosphere, such as NPO and NAO (Figure 1). The nonlinearity between DMS emission and the associated CRE was studied previously without taking into account the shipping emissions (Pandis et al., 1994; Russell et al., 1994; Gunson et al., 2006; Thomas et al., 2011). Here, to evaluate the CRE induced by both ISE and DMS emissions with a consideration of their interactions, we selected three regions for detailed analysis: the NPO and NAO where the ISE dominate the concentrations of sulfur dioxide and sulfate aerosols, and the Southern Ocean where DMS is the dominant source (Figure 2).

This study employs an Earth system model including an interactive aerosol model that simultaneously resolves both external and internal mixtures of sulfate, black carbon, and organic carbon aerosols. Aerosol mixing in this way can resolve aerosol activation process more realistically than either mixing all aerosol species internally or ignoring any mixing at all. By comparing the results with the default aerosol scheme that assumes internal mixing, we also quantify the impacts of various assumptions of aerosol mixing states on estimates of the CRE of ISE and DMS emissions. We further quantify the ISE-induced CRE based on various regulations of the International Maritime Organization (IMO) on the fuel sulfur content. Therefore, our findings have important implications for policy makers and future estimates of CRE induced by both ISE and DMS emissions.

2 Methods

2.1 Climate model

The Community Earth System Model version 1.2.2 (CESM1.2.2) is configured with the Community Atmosphere Model version 5.3 (CAM5.3). CAM5.3 includes a modal aerosol model with an option of 3 or 7 lognormal distributions of aerosol size (MAM3 or MAM7). In this study, a new modal aerosol model—the two-Moment, Multi-Modal, Mixing-state resolving Aerosol
model for Research of Climate (MARC; version 1.0.3 here) (Kim et al., 2008; Kim et al., 2014; Rothenberg and Wang, 2016, 2017; Grandey et al., 2018) is introduced and used to evaluate both the DRE and CRE of ISE. The details of MARC are described below. Aerosol DRE are represented by coupling between aerosols and radiation. Aerosol CRE are included by activating aerosols to work as cloud condensation nuclei and ice nuclei in the stratiform clouds (Morrison and Gettelman, 2008; Gettelman et al., 2010). Parameterization of aerosol activation is based on particle size and hygroscopicity of aerosols. Similar to other climate models, CAM5 does not directly include aerosol’s influence through microphysics on convective clouds, but it allows aerosols to influence convective clouds indirectly, such as by aerosol’s effect on circulation, surface evapotranspiration and so on.

### 2.2 MAM3

The default aerosol scheme in CESM 1.2.2, MAM3, has three modes, each with a lognormal size distribution: Aitken, accumulation, and coarse. Various aerosol species are internally mixed within each mode. Aitken mode is a mixture of sulfate, secondary organic carbon and sea salt; accumulation mode is mixture of sulfate, black carbon, primary organic carbon, secondary organic carbon, dust, and sea salt; coarse mode is a mixture of dust, sea salt, and sulfate (Liu et al., 2012). MAM3 tracks both the mass concentration and the number concentration of each aerosol mode.

### 2.3 MARC

MARC uses seven modes with different lognormal size distribution to represent the population of sulfate and carbonaceous aerosols: three modes for pure sulfate (nucleation or NUC, Aitken or AIT, and accumulation or ACC), one each for pure black carbon (BC) and pure organic carbon (OC), one mixture of BC–sulfate in core-shell structure (MBS), and one mixture of OC–sulfate (internal mixture; MOS). MARC predicts total particle mass and number concentrations while assuming the standard deviation within each of the seven modes to define at any given time the lognormal distribution of particle size. In addition, carbonaceous mass concentrations inside MBS and MOS are also predicted to allow the mass ratios between sulfate and carbonaceous compositions to evolve over time, changing the optical and chemical properties of the mixed aerosols. The emissions of mineral dust and sea salt that MARC uses are calculated by the land surface model and atmosphere model, respectively (Mahowald et al., 2006; Albani et al., 2014; Scanza et al., 2015). Mineral dust and sea salt are each represented by four bins with fixed sizes in MARC. For details of MARC aerosol mode size distribution and chemical parameters, please refer to Rothenberg and Wang (2017). Note that in MARC model, gas-phase sulfur compounds can be oxidized in both gaseous and aqueous phase to form sulfate that could enter aerosol phase in several pathways: (1) aerosol nucleation to form new nucleation mode sulfate aerosols; (2) condensation of gaseous sulfuric acid on both external sulfate and carbonaceous aerosols (the latter specifically ages carbonaceous aerosols to form sulfate-carbonaceous aerosol mixtures); and (3) evaporation of cloud and rain drops that resuspends aqueous sulfate to accumulation mode sulfate aerosol (Kim et al., 2008; Grandey et al., 2018; Rothenberg et al., 2018).

### 2.4 Difference between MARC and MAM3

The most fundamental difference between MARC and MAM is that MARC includes both external and internal mixtures of aerosols in fifteen modes while MAM treats all aerosols as internal mixtures in three modes. As a result, the processes of many aerosol microphysical processes including gaseous condensation, new particle formation, and nucleation scavenging differ between these two models (Kim et al., 2008; Grandey et al., 2018; Rothenberg et al., 2018). “For instance, aerosol activation or nucleation scavenging in MARC and MAM3 is calculated based on competition for water vapor among various types (or modes)
of aerosols with different hygroscopicity. In this case, external sulfate modes and the mixture of BC and sulfate (MBS) with BC as core and sulfate as shell would have the same hygroscopicity as sulfate, while external BC and OC would have much lower hygroscopic values. Whereas, MAM calculates this process based on the volume weighted hygroscopicity of each mode based on all the aerosol constitutions within the mode. In that case, the change of individual aerosol species would not influence much the number of activated aerosol substantially (Rothenberg et al., 2018).

2.5 Radiation diagnostics

In the diagnostic mode of CESM-MARC, the DRE are diagnosed by calling the radiation scheme three times in each radiation time step. The first call does not include any aerosols, providing “clean-sky” diagnostics (Ghan, 2013). The second call includes only mineral dust and sea salt aerosols. The third call includes all aerosols. The first and third call are diagnostic, i.e. the radiation budget calculated from these two calls are only used to as model output, therefore they do not influence model integration in the next time step; while the second call is prognostic, i.e., the radiation budget from this call is passed to other model schemes to calculate associated model variables, such as temperature, surface evaporation and so on. Therefore, DRE of only dust and sea salt aerosols are prognostic while all other aerosols including ISE are diagnostic. Note that all radiation variables calculated in these three calls are stored in the model history files for further analyses. In the complimentary MAM3 simulations, the first radiation call (prognostic) includes all aerosols while the second call is a “clean-sky” diagnostic call, excluding all aerosols.

The DRE of aerosols is calculated by subtracting the TOA radiation in the “clean-sky” call (no direct aerosol–radiation interaction) from that in the call that includes all aerosols. The CRE is calculated by subtracting the TOA radiation at clear-sky from that at all-sky in the “clean-sky” call (Ghan, 2013). The calculation of the DRE and CRE of ISE takes a further step—subtracting the DRE and CRE in simulation without ISE from that with ISE. In this way, the DRE and CRE of ISE can be isolated and evaluated separately. Note that all radiative effects are calculated at TOA.

2.6 Experimental design

Three groups of simulations are designed to evaluate: (a) the DRE and CRE of ISE, and (b) the sensitivity of the ISE-induced CRE to both DMS emissions and aerosol mixing assumptions. CAM5 was run at a horizontal resolution of 1.875°×2.5° and 30 vertical layers with sea surface temperature (SST), sea ice, greenhouse gas concentrations prescribed at the level of year 2000. The aerosol emissions in year-2000 were used except for modified shipping and DMS emissions. The DMS emission is prescribed with a global annual average of 18.2 Tg S yr⁻¹ in DMS reference simulations (Dentener et al., 2006). Each simulation runs for 32 years driven by 12-month cyclic climatological sea surface temperature, with the first 2 years discarded as spin up. Since the SSTs were prescribed, a 2-year period of spin up should be enough for aerosol concentrations and other model components to reach an equilibrium state (e.g., Righi et al., 2011).

The first group uses CESM-MARC and includes four simulations, which share the same DMS reference emissions \((DMSRef)\) while differing in four various ISE of sulfur compounds (sulfur dioxide and sulfate) (Table 1). \(ShipZero\_DMSRef\) simulation is integrated excluding all aerosol and aerosol precursor emissions from ISE, i.e., sulfur dioxide, sulfate aerosol, organic carbon aerosol, and black carbon aerosol. The other three simulations—\(ShipLow\_DMSRef, ShipRef\_DMSRef,\) and \(ShipHigh\_DMSRef\)—include the standard emissions of carbonaceous aerosols (e.g., BC and OC) while use three various emission scenarios of sulfur compounds from ISE. The three emission scenarios are based on the assumptions of sulfur content of the heavy fuel oils for ocean-going ships. Currently, the average sulfur content is 2.7% (Corbett and Koehler, 2003; Endresen et al., 2005), which is equivalent to about 5.4 Tg S year⁻¹, referred to as \(ShipRef\). On the other hand, as of 2013 the high sulfur
fuel oil that has 3.5% sulfur content continued to be permitted outside the Emission Control Areas (Lauer et al., 2009; Winebrake et al., 2009), referred to as ShipHigh. However, the IMO has planned to lower the sulfur content to 0.5% outside the Emission Control Areas (Winebrake et al., 2009; Notteboom, 2010; International Maritime Organization, 2016) after 2020, referred to as ShipLow. In ShipLow and ShipHigh, the total global sulfur shipping emissions are 1.0 and 7.2 Tg S year$^{-1}$, respectively. These numbers related to annual sulfur emissions are generally estimated based on the total amount of heavy fuel burnt by ships and the associated emission rates, whose uncertainties were addressed by Peters et al. (2012). The differences between these three and the zero shipping emission scenarios represent how various regulations on marine fuel influence the ISE-induced CRE.

The second group also uses CESM-MARC and is comprised of three pairs of simulations: (ShipRef_DMSZero, ShipZero_DMSZero), (ShipRef_DMSLow, ShipZero_DMSLow), and (ShipRef_DMSRef, ShipZero_DMSRef). Note that the pair of (ShipRef_DMSRef, ShipZero_DMSRef) simulations are also part of the first group. The annual emission of DMS is 18.2 Tg S year$^{-1}$ in the DMSRef simulations (Dentener et al., 2006; Liu et al., 2012), and half of that in the DMSLow simulations. DMS emission is excluded in the DMSZero simulations. Each pair of the simulations include ShipZero and ShipRef, the difference of which represents the ISE-induced impacts. The purposes of the DMSZero and DMSLow simulations are to quantify the sensitivities of the ISE-induced CRE (i.e., the difference of CRE in each pair of simulations) to DMS emission and the associated large uncertainty in DMS emission, respectively (Quinn et al., 1993; Dentener et al., 2006). DMS emission in the DMSLow simulation is 9.1 Tg S year$^{-1}$, which is close to the lower boundary of DMS emission estimates, i.e., 8 Tg S year$^{-1}$ (Quinn et al., 1993). Such sensitivities are examined by calculating the differences among the three pairs of DMS simulations.

The third group is the same as the second, but using the default MAM3 aerosol module of CAM5.3 in CESM. The purpose for designing the third group is to cross-validate the simulated DMS impacts on the ISE-induced CRE in the second group. One bonus of the third group is to quantify the impacts of using different aerosol modules with different aerosol mixing states on the simulated results. The anthropogenic emissions for MAM3, which are described by Liu et al. (2012), differ slightly from those for MARC. All of the experiments are summarized in Table 1.

3 Results

3.1 DRE of ISE

The all-sky DRE of various aerosol species from ISE is diagnosed as the difference between ShipRef_DMSRef and ShipZero_DMSRef and shown in Figure 3. The total ISE can cause a global negative (cooling) DRE of $-23.5$ mW m$^{-2}$, with the strongest negative (cooling) DRE in the areas with intense shipping tracks, such as mid-latitude areas in the Pacific Ocean and Atlantic Ocean, South China Sea, North Indian Ocean, and the Red Sea. The sulfate aerosols in the accumulation mode (i.e., ACC) contribute 89% to the global total DRE, followed by MOS aerosols with a contribution of 22%. Note that OC and MBS has a counteracting warming effect (Remember that all gas-phase and aerosol emissions from shipping have been removed in ShipZero scenarios). The contributions of other aerosol species are very limited and their magnitudes are smaller than 6%. The magnitude of the total cooling effect is within the range from $-50$ to $-10$ mW m$^{-2}$ estimated in previous studies (Endresen et al., 2003; Schreier et al., 2007). The meridional variations of global zonally-averaged total DRE show that the DRE has the strongest cooling effect of $-80$ mW m$^{-2}$ between 30°N and 40°N and becomes weaker towards both polar regions and can be ignored beyond 45°S and 60°N. The all-sky DRE of total aerosols in ShipLow_DMSRef and ShipHigh_DMSRef have similar patterns to those in ShipRef_DMSRef and have magnitudes of $+1.0$ and $-33.0$ mW m$^{-2}$, respectively. All of the calculated global DRE values except for BC are confident at the 90% level.
3.2 CRE of ISE under various shipping emission regulations

The CRE of ISE is much stronger than the DRE and shows different spatial patterns under various shipping emission regulations (Figure 4). At the reference level of shipping emissions (ShipRef_DMSRef), significant cooling CRE in SW is simulated in areas of intense shipping tracks, such as the mid-latitude Pacific Ocean and the Baffin Bay between Canada and Greenland, with a global average of $-0.218$ W m$^{-2}$. The LW CRE shows positive values in some small areas of high latitude, with a global average of $+0.065$ W m$^{-2}$. Consequently, the global net CRE (SW+LW) is $-0.153$ W m$^{-2}$ with a similar spatial pattern to that of SW. At the high level of shipping emissions (ShipHigh_DMSRef), the CRE changes to $-0.253$, $+0.073$, $-0.179$ W m$^{-2}$ for SW, LW, and net, respectively; more areas show significant changes than in ShipRef_DMSRef. Note that all of the above values are statistically significant above the 90% confidence level. However, at the low level of shipping emissions (ShipLow_DMSRef), fewer areas demonstrate significant changes than in ShipRef_DMSRef and ShipHigh_DMSRef and the global averages of the CRE are not significant at the 90% confidence level for SW and net. These results indicate that more stringent shipping emission regulation on sulfur content decided by the IMO to be applied after 2020 could effectively reduce or even largely eliminate the net CRE induced by ISE.

Further analyses demonstrate that the changes in CRE are caused by perturbations in both cloud water path (CWP; Figure S1) and column-integrated cloud droplet number concentrations (CDNC; Figure 5) induced by ISE. Figure S1 demonstrates significant increases in total CWP mainly over the NPO and NAO at the reference and high levels of ISE. The increases in total CWP is largely (87%) attributed to liquid CWP with the remaining contribution (13%) from ice CWP at the reference shipping emission level. Such increases in CWP could reflect more solar radiation to space and thus cause a cooling radiative effect at TOA, as shown in Figure 4. Note that very limited areas in North Pacific Ocean show significant increases in ice CWP, indicating that a small portion of surface shipping emissions could be vertically transported to very high altitude and form ice cloud. At the high level of shipping emissions, a larger increase in CWP is simulated, which is consistent with the cooler radiative effects. However, no significant changes are simulated in total, ice, or liquid CWP at the low level of shipping emissions. Associated with increases in CWP, the column-integrated cloud droplet number concentration (CNDC) also illustrates significant increases at all levels of shipping emission (Figure 5), which collocate with increases in CWP (Figure S1) and decrease in CRE (Figure 4) over the NPO and NAO. The sulfate aerosols from shipping emissions are highly efficient cloud condensation nuclei (CCN) and thus can increase the CDNC, which in turn affects CWP. Note that cloud area fraction does not exhibit any significant changes due to shipping emissions (not shown).

3.3 CRE of ISE under various DMS emissions

The biogenic emissions of DMS over oceans can be oxidized to sulfates and compete against shipping emitted sulfates for CCN and thus could influence the ISE-induced CRE. We find that the shipping emission-induced CRE exhibits significantly different patterns and global averages at different emission levels of DMS (Figure 6). With DMS emissions ranging from the reference level to low and zero levels, the magnitude of the ISE-induced negative CRE at SW increases from 0.218 to 0.457 and 2.435 W m$^{-2}$ on global scale, respectively; significant negative CRE is simulated over more areas in the SO. It is worth noting that the ISE-induced CRE can reach up to $-6$ W m$^{-2}$ when DMS emission is turned off, such as over NPO and NAO, which is a very large negative forcing even on the local scale. Since there are no comparable values in the literature, this large negative forcing warrants a detailed evaluation in future studies using different climate models. For CRE at LW, more areas with significant warming are seen in the SO, NPO, and NAO, with the global averages change from $+0.065$ to $+0.073$ and $+0.253$ W m$^{-2}$ when DMS emissions changes from the reference to low and zero levels, respectively. For net CRE, it shares the similar features with those at SW, but with smaller magnitudes.
The DMS emissions influence the ISE-induced CRE by perturbing the ISE-induced changes in CWP and column-integrated CDNC (Figures S2 and 7). The shipping emission-induced changes in the total and liquid CWP increase as DMS emission decreases, particularly over the SO, the NPO, and the NAO, while no significant changes are simulated in the ice CWP. The increased CWP is closely associated with the increases in the column-integrated CDNC, which changes from \(0.305 \times 10^9 \text{ m}^{-2}\) (2.5%, relative to climatological CDNC in the \(\text{ShipZero}_\text{DMSRef}\) simulation) to \(0.476 \times 10^9 \text{ m}^{-2}\) (3.9%) and \(0.999 \times 10^9 \text{ m}^{-2}\) (8.3%) on global scale as DMS emission decreases. These results imply that DMS emissions are the dominant sources of cloud seeds over remote oceans. The most prominent increases in CDNC are seen in the SO, NPO, and NAO. These results suggest important roles of DMS emissions playing in modulating the ISE-induced changes in cloud properties and radiation. Note that cloud area fraction does not exhibit any significant change (not shown).

As demonstrated in the above analysis, the impacts of DMS emissions on cloud response to shipping emissions are the most prominent over the SO, the NPO, and the NAO, so further analyses are performed over these three regions. Figure 8 shows cloud responses to shipping emission at different DMS emission levels over the three oceanic regions. Generally, cloud responses become weaker and weaker as DMS emission increases from zero (DMSZero) to low (DMSLow) and reference (DMSRef) level over all of the three regions. The most prominent change in cloud response is over the NPO, followed by the NAO and the SO, which is probably due to the higher contribution of shipping emission to the total sulfur dioxide and sulfate aerosols over the NPO than the NAO and the SO (Figure 2b and 2e). The removal of DMS emission (DMSZero) has a much stronger influence on cloud response to shipping emission than reducing DMS emission by half (DMSLow), indicating a strong non-linear competing effect for CCN between DMS and shipping emission.

### 3.4 CRE of DMS under various shipping emissions

Similar to DMS emissions’ impacts on the ISE-induced CRE and cloud properties, ISE could also influence the DMS emission-induced CRE and cloud properties. Generally, stronger cooling net CRE (−7.518 vs. −5.611 \text{ W m}^{-2}\) induced by DMS emissions are seen when shipping emissions are absent, particularly in areas of intense shipping tracks, such as over the NPO and the NAO (Figure 9). Such a net cooling CRE is mainly the result of SW CRE. Stronger cooling CRE is associated with larger increases in liquid and total CWP (Figure S3) and column-integrated CDNC (Figure S4) in simulations without shipping emissions than those with shipping emissions.

It is worth pointing out that DMS emissions have significant warming CRE at LW, particularly over mid- and high-latitude regions in the Southern Hemisphere and high-latitude regions in the Northern Hemisphere regardless of the presence of shipping emissions (Figure 9). Such a warming CRE could be attributed to increases in total cloud area fraction, which is further attributed to increases in the middle and low cloud area fraction in the high-latitude regions in both hemispheres (Figure S5). Our results indicate that DMS is a significant source to CCN in the extremely clean polar regions in both hemispheres.

The area-averaged cloud responses over the SO, the NPO, and the NAO, to DMS emissions at different shipping emission levels are shown in Figure 10. Cloud responses to DMS are stronger over the SO than over the NPO and the NAO regardless of the presence of shipping emissions due to the fact that DMS and shipping emissions respectively dominate the sulfur concentrations over the SO, and the NPO and the NAO (Figure 2). Moreover, cloud responses to DMS emissions become much stronger over all of the three oceanic regions when shipping emissions have been removed. However, such changes in cloud responses to DMS due to removal of shipping emissions (i.e., the slopes of the curves) are stronger over the NPO and the NAO than over the SO, which is caused by very limited shipping emissions over the SO. These results again indicate a strong non-linear competing effect for CCN between DMS and shipping emission.
3.5 Impacts of choice of aerosol module on the results

Besides the impacts of various ISE regulations and DMS emissions on the ISE-induced CRE, various assumptions about the aerosol mixing states could also have an impact. Figure 11 shows the same results as Figure 6 but using the MAM3 aerosol module instead of MARC. At the reference level of DMS emissions (DMSRef), the ISE-induced CRE are generally stronger in MAM3 (SW: −0.319, LW: +0.064, and Net: −0.255 W m⁻²; Figure 11) than in MARC for SW and net radiation (SW: −0.218, LW: +0.065, net: −0.153 W m⁻²; Figure 6). More areas with significant cooling CRE are simulated in MAM3 than in MARC, particularly in the Atlantic Ocean, West Pacific Ocean, and North Indian Ocean. At the low level of DMS emissions (DMSLow), both the global averages and spatial patterns of the CRE in MAM3 are very similar to those in MARC. The two aerosol modules show the biggest differences in CRE when DMS emissions are excluded (DMSZero). MARC simulates strong ISE-induced CRE over tropical regions and the subtropical and mid-latitude areas of the SO, while MAM3 gives no significant CRE over these regions. Generally, the ISE-induced CRE is stronger in MARC than in MAM3 when DMS emissions are excluded. The associated changes in CDNC and CWP due to ISE illustrates similar patterns to changes in CRE (Figures S6 and S7). A possible reason for such differences is the various mixing assumptions about sulfate and sea salt aerosols in MARC (external mixing) versus in MAM3 (internal mixing). Interested readers are referred to Grandey et al. (2018) for discussion of the differing radiative effects produced by MARC and MAM. To track down all the possible reasons for the differences in the ISE-induced CRE between the two aerosol schemes, more detailed analyses on a long chain of processes related to both aerosols and clouds are required, as done by Peters et al. (2014), which is out of the scope of this study and warrants more studies in the future.

By comparing Figure 8 with Figure 12 we also observe significantly different impacts of DMS emissions on cloud response to shipping emissions (i.e., the slopes of these curves). MAM3 simulates a weaker impact of DMS emissions on cloud response to shipping emissions than MARC, indicating a weaker non-linear competing effect for CCN between DMS and shipping emissions in MAM3 than MARC.

4 Conclusions and Discussion

Aerosols from ISE could exert significant cooling on the Earth’s climate system through aerosol–cloud and aerosol–radiation interactions. To reduce the pollution and climatic effects from this emission source, the IMO set various emission caps on sulfur content of marine fuel oil to be implemented in the future. Using a state-of-the-art climate model, we find that the newly proposed more stringent emission regulations of shipping emissions can effectively reduce the ISE-induced CRE. As demonstrated in our results, reducing sulfur contents from 3.5% to 2.7% and 0.5% could reduce both DRE (from −51.4 to −36.7 and −3.9 mW m⁻²) and CRE (from −0.179 to −0.153 and −0.001 W m⁻²) due to ISE, respectively. Although the ISE-induced CRE would be insignificant on a global scale if sulfur contents of ship fuels were reduced to 0.5%, over some regions significant CRE can still be detected—e.g., high latitude regions of the eastern Pacific Ocean. Therefore, implementation of cleaner fuels in shipping sector, such as natural gas, could be a potential solution for completely eliminating sulfate-induced CRE.

More importantly, we find that the magnitude and regional spatial pattern of the ISE-induced CRE are highly sensitive to natural DMS emissions. With DMS emissions reducing from 18.2 to 9.1 Tg S yr⁻¹ and zero, the ISE-induced net CRE changes from −0.153 to −0.384 and −2.182 W m⁻², respectively. On the other hand, the DMS-induced net CRE changes from −5.611 to −7.518 W m⁻² when shipping emissions at the reference level are removed in the simulations. It is worth noting that DMS is a significant source to CCN in the extremely clean polar regions in both hemispheres. The strong interactions of CRE between DMS and shipping emissions can be attributed to the nonlinearity in the responses of cloud processes to aerosols, particularly the
aerosol activation parameterizations (Abdul-Razzak et al., 1998). In a relatively clean environment, activated aerosol number concentration increases as ambient aerosol number concentration increases until reaching a peak at a specific aerosol number concentration, after which it decreases as ambient aerosol number concentration increases unless ambient vapor concentration is drastically increased. In other words, the fraction of activated aerosols decreases as ambient aerosol concentration increases (Figure S8). From the perspective of simulation, this nonlinearity in aerosol activation strongly suggests a reevaluation of CRE induced by shipping and DMS emissions as well as a reevaluation of parameterizations of aerosol–cloud interactions in the general circulation models. From the perspective of field measurements of aerosol–cloud relationships, it warrants careful attention when selecting measurement locations—shipping emissions-related measurements should be collected along intense shipping tracks while in areas with as little DMS emissions as possible to avoid contamination from DMS, and vice versa. Moreover, locations containing both shipping and DMS emissions should also be identified and sampled, in order to investigate non-linear interactions between the emissions.

Finally, we find that two different aerosol schemes, with different representations of aerosol mixing state, could produce a large difference (about 67%) in the ISE-induced global CRE. Generally, MARC aerosol module shows stronger nonlinear cloud response to DMS and shipping emissions than MAM3. Overall, numerical studies on the uncertainties in the shipping emission-induced CRE due to various ISE regulations, aerosol interactions, and aerosol mixing states can provide useful information for policy makers and have implications for future projections of anthropogenic climate change.

Besides the above-mentioned contributors to the uncertainty in estimating the CRE induced by shipping emissions, spatial resolution of the model is another significant source of this uncertainty. Possner et al. (2016) found that the ship-induced shortwave CRE could increase by a factor of two as model spatial resolution decreases from 1 km to 50 km. With higher spatial resolution, models can resolve fine-scale dynamical processes and feedbacks, such as interaction between aerosol and cumulus clouds (Malavelle et al., 2017). Though model resolution-induced uncertainty is not the focus of this study, it should be taken into account when interpreting the spread of shipping-induced CRE in studies of multi-model comparison.

Though we employed a state-of-the-art climate model in this study, it is not without caveats, given that neither MARC nor MAM (including both MAM3 and MAM7 in CAM5) aerosol modules treat nitrate aerosols because of the high computational expense for related aerosol-gaseous chemistry and aerosol thermodynamics calculations (Liu et al., 2012). Lack of treating nitrate aerosols could result in uncertainties in our results based on the fact that both mass of nitrate aerosols emitted from international shipping (e.g., Righi et al., 2011) and their hygroscopicity values (e.g., Kawecki and Steiner, 2018) are very similar to those of sulfate aerosols, and thus nitrate aerosols could have non-negligible competing effects on CRE with sulfate aerosols. Despite that some of the results from this study could be used to qualitatively project the potential outcome, a quantitative assessment should be facilitated to address this topic with an improved model.

Acknowledgments

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The authors declare no competing interests.

Data and code availability

The MARC source code is available via https://github.mit.edu/marc/marc_cesm/ and also archived with DOI 10.5281/zenodo.1117370, along with documentation on how to install and run the model. The commit 23e08fe was used in this study. All the analysis code and model output data analyzed are available via https://drive.google.com/drive/folders/1GHjrpvO06mzC8iFyT0Eif0PRiS5cReV?usp=sharing.

References


<table>
<thead>
<tr>
<th>Aerosol Modules</th>
<th>Experiments</th>
<th>DMS emissions</th>
<th>Ship emissions</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>MARC</td>
<td>DMS</td>
<td>DMSRef</td>
<td>ShipZero</td>
<td>DMS emission (Tg S yr⁻¹):</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ShipLow</td>
<td>DMSZero: 0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ShipRef</td>
<td>DMSLow: 9.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ShipHigh</td>
<td>DMSRef: 18.2</td>
</tr>
<tr>
<td></td>
<td>DMSLow</td>
<td>ShipZero</td>
<td></td>
<td>Ship emission (Tg S yr⁻¹):</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ShipRef</td>
<td></td>
<td>ShipZero: 0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ShipRef</td>
<td>ShipLow: 1.0 (0.5%)</td>
</tr>
<tr>
<td></td>
<td>MAM3</td>
<td>DMS</td>
<td>DMSRef</td>
<td>ShipRef: 5.4 (2.7%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ShipZero</td>
<td>ShipHigh: 7.0 (3.5%)</td>
</tr>
<tr>
<td></td>
<td>DMSZero</td>
<td>ShipZero</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>ShipRef</td>
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</tr>
<tr>
<td></td>
<td>DMSLow</td>
<td>ShipZero</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>ShipRef</td>
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</tr>
</tbody>
</table>

Notes: in ShipZero experiments, emission rates of all gas-phase and aerosol species from shipping emissions are set to zero; while in ShipLow, ShipRef, and ShipHigh experiments, all shipping emission rates (such as OC and BC) are set to year-2000 values except for emission rates of sulfur compounds (i.e. SO₂ and SO₄) which are modified. The percent for ship emission in the last column stands for the proportion of sulfur content in the heavy fuel oils by mass.

Figure 1. Spatial patterns of annual means of sulfur emission (g S m⁻² yr⁻¹) from (a) international shipping and (b) natural DMS in the simulation at the reference emission level (i.e., ShipRef_DMSRef). The numbers below each panel are the global total annual emissions. Three regions are selected for further analysis: The North Pacific Ocean (NPO; 20°N–60°N, 140°E–240°E), the North Atlantic Ocean (NAO; 20°N–60°N, 300°E–360°E), and the Southern Ocean (SO; 20°S–60°S, 0°E–360°E).
Figure 2. Spatial patterns of (a) annual mean concentrations of total SO$_2$ (units: parts per billion by volume; ppbv), (b) and (c) respectively the contributions of shipping emission and natural DMS to total SO$_2$ in the lowest model layer. (d)–(f) are the same as (a)–(c), but for sulfate aerosols. These results are from MARC simulations and calculated as the differences between the simulations with the international shipping and DMS emissions at the reference and zero levels (i.e., Ship$_{Ref}$ DMS$_{Ref}$ minus Ship$_{Zero}$ DMS$_{Ref}$ and Ship$_{Ref}$ DMS$_{Ref}$ minus Ship$_{Ref}$ DMS$_{Zero}$).
Figure 3. Simulated direct radiative effect (DRE; units: mW m\(^{-2}\)) of ISE at TOA by MARC. The DRE is calculated as the difference between simulation results with and without ISE (i.e., ShipRef_DMSRef minus ShipZero_DMSRef) and averaged over the 30-year period of simulations at all-sky conditions. Panels (a)–(f) show the spatial patterns of DRE due to ISE with the global mean differences and the associated significant levels indicated by the numbers below each panel and panel (g) is the meridional variations of zonal mean DRE for various aerosol types from ISE and their total effects. The expansions of the abbreviations can be found in Section 2.3. The black dots represent grid points that are statistically significant above the 90% confidence level based on the two-tailed Student’s \(t\)-test.
Figure 4. Spatial patterns of MARC simulated cloud radiative effect (CRE; units: W m\(^{-2}\)) at TOA of ISE with various shipping emission levels. The CRE is calculated as the differences of radiation flux at TOA and at all-sky conditions between the simulation without shipping emissions and three simulations with the same DMS emissions at the reference level but various shipping emission levels (i.e., low, reference, high) in short-wave (SW), long-wave (LW), and net (SW+LW) and averaged over the 30-year simulation period. The numbers below each panel are the global means, standard deviation across the 30-year period, and the confidence level. The red dots represent grid points that are statistically significant above the 90% confidence level based on the two-tailed Student’s t-test.
Figure 5. Spatial patterns of MARC simulated column-integrated cloud droplet number concentration ($\times 10^9 \text{ m}^{-2}$) response to international shipping emissions. The responses are calculated as the differences of cloud droplet number integrated through the whole atmospheric columns between the simulation without shipping emissions and three simulations with the reference shipping emission and various DMS emissions (i.e., zero, low, and reference) over the 30-year simulation period. The numbers below each panel are the global means, standard deviation across the 30-year period, and the confidence level. The black dots represent grid points that are statistically significant above the 90% confidence level based on the two-tailed Student’s $t$-test.
Figure 6. Spatial patterns of MARC simulated cloud radiative effect (CRE; units: W m\(^{-2}\)) at TOA of ISE at various DMS emission levels. The CRE is calculated as the differences of radiation flux at TOA and at all-sky conditions between the simulation without shipping emissions and three simulations with the same shipping emissions at the reference level but various DMS emission levels (i.e., zero, low, and reference) in short-wave (SW), long-wave (LW), and net (SW+LW) and averaged over the 30-year simulation period. The numbers below each panel are the global means, standard deviation across the 30-year period, and the confidence level. The black dots represent grid points that are statistically significant above the 90% confidence level based on the two-tailed Student’s \(t\)-test.
Figure 7. Spatial patterns of MARC simulated column-integrated cloud droplet number concentration ($\times 10^9$ m$^{-2}$) response to international shipping emissions. The responses are calculated as the differences of cloud droplet number integrated through the whole atmospheric columns between the simulation without shipping emissions and three simulations with the reference shipping emission and various DMS emissions (i.e., zero, low, and reference) over the 30-year simulation period. The numbers below each panel are the global means, standard deviation across the 30-year period, and the confidence level. The black dots represent grid points that are statistically significant above the 90% confidence level based on the two-tailed Student’s $t$-test.

Figure 8. Impacts of DMS emissions on cloud responses to international shipping emissions. (a) Cloud radiative effects at TOA (W m$^{-2}$), (b) column-integrated cloud water path (g m$^{-2}$), and (c) column-integrated cloud droplet number ($\times 10^9$ m$^{-2}$). The green, purple, and blue curves respectively represent quantities area-averaged over the North Pacific Ocean (NPO), the North Atlantic Ocean (NAO), and the Southern Ocean (SO), which are shown as red boxes in Figure 1a. These results are from MARC simulations.
Figure 9. Spatial patterns of MARC simulated cloud radiative effect (units: W m\(^{-2}\)) of DMS emissions at various shipping emission levels. The CRE is calculated as the differences of radiation flux at TOA and at all-sky conditions between the simulation without DMS emissions and two simulations with the same DMS emissions at the reference level but various shipping emission levels (i.e., zero and reference) in short-wave (SW), long-wave (LW), and net (SW+LW) and averaged over the 30-year simulation period. The numbers below each panel are the global means, standard deviation across the 30-year period, and the confidence level. The black dots represent grid points that are statistically significant above the 90% confidence level based on the two-tailed Student’s \(t\)-test.

Figure 10. Impacts of ISE on cloud responses to DMS emissions. (a) Cloud radiative effects at TOA (W m\(^{-2}\)), (b) column-integrated cloud water path (g m\(^{-2}\)), and (c) column-integrated cloud droplet number (\(\times\)10\(^9\) m\(^{-2}\)). The green, purple, and blue curves respectively represent quantities area-averaged over the North Pacific Ocean (NPO), the North Atlantic Ocean (NAO), and the Southern Ocean (SO), which are shown as red boxes in Figure 1a. These results are from MARC simulations.
Figure 11. Same as Figure 6, but using a different aerosol module, namely MAM3 rather than MARC.

Figure 12. Same as Figure 8, but using a different aerosol module, namely MAM3 rather than MARC.