We thank the reviewers for their good summary and comments. Below is our point-to-point response. We use PxxLxx, Fig. xx, and Table xx for revised version and OPxxLxx, OLxx, OFig.xx, OTable xx for the original submission. Please check PxxLxx in the revised file, whose name is tagged “track”.

Review 1 (R1):

1. Relevance and novelty of this study:
R1#1. It would be helpful if the authors could state the reasons for focusing on sulfur species in a more straightforward way in the introduction section. While the introduction highlights the environmental impacts of sulfur species, other pollutants not discussed in this paper also affect the environment. Why should we focus on sulfur species? Is the bias of aerosol climate models predominantly due to sulfur? As we mentioned in P2L74-89, the sulfur cycle plays a key role in atmospheric quality, climate, and ecosystems. This importance is primarily due to the fact that the atmospheric sulfate component itself contributes to radiation forcing (RF) almost as much as all other major non-natural aerosol components, as concluded from 16 AeroCom model results (Myhre et al., 2013). More importantly, uncertainty in sulfate simulations in current climate models is a major contributor to biases in aerosol optical depth (AOD, Fig. 3 in Gliß et al., 2021) and radiative forcing (RF, Fig. 7 in Myhre et al., 2013). Above discussion has been added in P2L89-94.

![Fig. 3 in Gliß et al., (2021): Species contributions to total AOD for each model (annual global average).](image1)

![Fig. 7 in Myhre et al., (2013): Component and total RF from 16 AeroCom models. Solid lines inside the boxes show the model mean, dashed lines show the median. The box indicate one standard deviation and whiskers indicate the max and min of the distribution.](image2)

R1#2. I would appreciate it if the authors could highlight the novelty of their work. Has any other research analyzed the ATOM data specifically for sulfur species?
To date, there is one publication that extends the simplified DMS oxidation scheme used in Community Atmospheric Model Chemistry version 6 (CAM6-chem) using ATom data (Fung et al., 2022). However, a comparison of SO2, a key sulfur species in the sulfur cycle, between CAM6-chem and ATom has not been reported. We have summarized previous studies of aerosols (including sulfur species) related to ATom in P3L134-145.

R1#3. Is this the first study that utilizes ATOM measurements in comparison with AeroCom models?
Our work is the first study to use ATom measurements for comparison with the AeroCom models, focusing on all sulfur species simulated in current aerosol climate models. This work...
extends previous efforts that utilize ATom measurements to evaluate the organic carbon (Hodzic et al., 2020) and black carbon (Katich et al., 2018) of AeroCom models. See P3L134-137.

R1#4. If there have been studies on sulfur variability and sources over the ocean, what novel findings does this paper present?

To our knowledge, there are no such publications investigating the variability and sources of all major sulfur species in the ocean. Our study aims not only to reveal sulfur variability based on multiple measurements and model simulations, but also to tease out the underlying processes behind the variability by integrating analyses of simulated sulfur species in aerosol climate models. This work therefore provides direction for improvements in aerosol climate models. See P3L141-145.

For example, our study found that all AeroCom models overestimate DMS near the sea surface (Fig. 4). Is this overestimation due to the model DMS vertical gradient being steeper than the ATom data (Fig. 8) or the DMS emissions being larger than they should be? A unique approach taken in our study revealed DMS emissions as one of the causes by integrating the analysis of DMS, SO2, and SO4 in a model-ATom comparison (P11L740-747). By reviewing the DMS emission methods adopted by the five AeroCom models (Table 3), we further found that the calculation of the air-sea exchange flux formula plays a key role in determining DMS emissions (P9L589-648).

The AeroCom models also tend to underestimate SO2 and overestimate SO4 over remote oceans. This performance prompted modelers to explore potential faster chemical conversion of SO2 and different dry and wet removal of SO2 and SO4. One suggestion for modelers is that using interactive oxidant calculation is insufficient to solve the problem because there is no systematic bias in sulfate simulations (Fig. 6) between the three participating models using online oxidants and the other two models using offline oxidants (Table 2). See P8L469-550.

R1#5. The authors should make some comparison with other studies in a new section or talk about it in an existing section of the paper.

To date, there have been no sulfur budget reports focusing on the vast ocean. However, previous AeroCom studies have reported global sulfate atmospheric loading and its diversity across multiple AeroCom models using monthly and global mean column loadings. The newly added Table 5 summarizes these studies, including their reported global and annual sulfate multi-model mean (MMM) and diversity ($\delta$). $\delta$ is related to the standard deviation (std_dev) and is defined as $\delta = \text{std_dev} / \text{MMM} \times 100$ (%). The results of this work are lower than AeroCom-I but higher than AeroCom-II, which may be related to the different target years involved in these studies. One point to note is that the diversity of AeroCom-III models $\delta$ has not reduced since AeroCom-I, which was studied nearly 20 years ago. The nearly 30% diversity combined with a greater proportion of sulfate than other anthropogenic aerosols make accurate sulfate modeling very important. Furthermore, these previous AeroCom studies were almost purely model studies, with only Gliß et al., 2021 introducing AOD-related light fields from ground stations and satellite retrievals. We have added this discussion in Sect. 4 (P12L807-823).
Furthermore, if we improve sulfur simulation, what advantages can we expect? It would be helpful if the authors could briefly discuss the implications of the new findings in the conclusion section. Improved sulfur models are needed, given that the sulfur cycle plays a critical role in atmospheric quality, climate, and ecosystems, and that its bias in current aerosol climate models is one of the largest uncertainties in air quality and RF research (see response to R1#1). Through a comprehensive multi-model and multi-instrument comparison of remote ocean sulfur species in this work, shortcomings in model sulfur simulations are pointed out and potential directions for improvement are suggested (see response to R1#4). We have added a brief discussion of the implications of the new findings in the Conclusion section (P15L946-956), summarizing our response to R1#1#4#5.

2. Information Overload and Simplification:

R1#7. Some parts of the paper contain overwhelming information that may be simplified or moved to supplements. For instance, Section 3.1 allocates 28 lines (OL212-L239) to discuss three different sampling intervals, which may not be key information the readers need to know. This information (corresponding to OFig 2 (a)(d)(g)(i)) only builds up one-third content in OFigure 2, which makes the main point of OFigure 2 very hard to catch. The discussion of the three-time resolution AMS data (OL212-239) and their visualization in Fig. 2 have been moved to Supplementary Information. We have now made Fig. 2 in a manner consistent with Figs. 3-4.

R1#8. Another example starts from OL317 where the authors spent time explaining how the flag ‘-888’ is replaced by ‘0’ to represent the low values, which, although crucial for validating results, may not be necessary for most readers. OL317-329 have also been moved to Supplementary Information.

R1#9. Additionally, the division of 5 models into 3 groups from OL340-L354 seems superfluous and is never referenced throughout the paper. I would recommend a description without grouping the models. These sentences are changed to “Specifically, CAM-ATRAS and GEOS have good SO4 vertical gradients over the tropical and NH oceans, but their SO4 values are too low compared to measurements over the Southern Hemisphere (SH) free troposphere. The SO4 of IMPACT and OsloCTM3 decreases too slowly with altitude, as shown by their overestimated SO4 values at high altitudes globally. E3SM performed SO4 simulations among other models. However, the performance of these models’ SO4 vertical profiles cannot simply be explained by the way the oxidant is applied, because among the five models, CAM-ATRAS, IMPACT, and OsloCTM3 used interactive oxidant calculations, while E3SM and GEOS used archived oxidant data (Table 2). The complexity of the chemistry deserves more attention. Of the five models, OsloCTM3 and GEOS participated in the multi-model OH assessment (Nicely et al., 2000) and OsloCTM3 had a shorter methane lifetime (relative to OH) than GEOS.” See P8L469-550.

3. Layout and Readability of Figures:

R1#10. Due to the huge load of information that is shown, optimizing the layout of figures is crucial to enhance readability. For example, I would recommend relocating the legends in OFigures 5-8 and putting this information on the top/bottom or right side of the charts. The legends in Figs. 6-9 (OFigures 5-8) are now at the top of the charts.
Since you ‘use 10-s merged data where observations above DL throughout the main text unless otherwise stated’ (OL241), could you just show the results of 10-s data only on OFigure 2 and move the other to the supplement?

Done. Please also see response to R1#7.

Additionally, as Atom-1/2/3/4 are not following the order of the four seasons, I recommend adding notes on the seasons at a proper place in OFigures 9,10,11, and 13 to guide the readers when reading through the section about seasonal changes in the paper.

Done.

4. Conclusion:

The conclusion section is mostly a summary of the content. As mentioned earlier, the implication of the new findings can be stated in this section.

We have added this paragraph in the Conclusion section (P15L946-956):

“Even after two decades of development, the diversity of sulfate simulations from AeroCom-I to AeroCom-III has not decreased. However, accurate sulfate simulation in current climate models is crucial to reduce radiative forcing biases. Several potential directions for improving sulfur simulations are suggested above. More importantly, apart from the shortcomings of individual models, all modelers should focus on the calculation of the air-sea exchange flux formula, as it plays a key role in determining DMS emissions. Modelers also need to study DMS and SO2 vertical transport as well as SO4 wet deposition during long-distance transport, as model biases are greatest at high altitudes. One suggestion to modelers is that the use of online oxidant fields is insufficient to explain the model sulfate bias, as there was no systematic bias in the sulfate simulations between the models using interactive oxidants and the models using archival oxidants in this study. The complexity of chemistry deserves more attention.”

Please also see response to R1#4.

Technical comments:

1. Please standardize the color and font of the indices of panels in OFigure 2.

Indices of panels in Fig. 2 have been removed based on the suggestion of reviewer 3 (R3P7L305).

2. Please refine OFigure 12 to maintain the consistent style of other figures.

Done.

3. In the caption of OFigure 9, AMS should be orange instead of ‘red’.

Done.

4. Please replace the vertical bar in OFigure 13 with a straight line as the shape and color is misleading.

Done.

5. In OL467, please add a period after ‘4’.

Done.

Reviewer 2 (R2):

In section 2.2, the authors say that they perform source attribution using sensitivity analysis method. By sensitivity analysis I would assume a perturbation approach where two simulations are performed: one with the baseline emissions and another where emissions from a given source are changed. The difference between the simulated variables in the two simulations would communicate
the sensitivity of the simulated concentrations to changes in precursor emissions from a particular source.

Yes, we need two simulated fields, one linked to baseline emission and the other linked to a given source, to diagnose source attribution. We have two methods to calculate the two fields in model simulation.

1. Run model twice for field \( F \). The first run \( F \) is linked to the baseline emission \( (F_{\text{base-emi}}) \) and the second run \( F \) is linked to a given source \( (F_{\text{tag-emi}}) \).

2. Run model once for fields \( F \) and \( F_{\text{tag}} \) (a newly added tracer) simultaneously. \( F \) is linked to the baseline emission and \( F_{\text{tag}} \) is linked to a given source.

The advantage of using the second method is obvious, simply for computational efficiency, especially when multiple tagged tracers are required. See newly added Sect. 2.3 “Tag-tracer study in GEOS”.

R2#2. However, section 5 mentions tagging method being used for source attribution within GEOS-Chem model. Tagging involves changing the chemical mechanism within the model to write out the concentration variables along with the label of the source which they originate from.

Each specific aerosol component in GEOS GOCART is simulated independently from the others, and the contribution of each emission type to the total aerosol mass is also not interfered by that of other emission types. Thus, additional aerosol tracers can easily be “tagged” according to emission source types. The GEOS GOCART module includes a tagged aerosol mechanism that allows GOCART to calculate and transfer multiple sets of aerosol fields (e.g., \( A \), \( A_{\text{tag}} \), etc) in a single job submission. See newly added Sect. 2.3 “Tag-tracer study in GEOS”.

R2#3. I would also like to point to Line 45 (abstract) where the authors write: “sensitivity studies by applying tagged tracers”. I recommend the authors to reframe this sentence by clarifying the exact source attribution approach and model (GEOS-Chem), to avoid possible confusion among readers.

Changed the sentence to “We identify their origins from anthropogenic versus natural sources with sensitivity studies by applying tagged tracers in GEOS model linking to emission types of anthropogenic, biomass burning, volcanic, and oceanic emissions.” See P1L44-47.

R2#4. Since you discuss the results from GEOS-Chem tagged simulation, please specify the tagging method in section 2.2. Cite the relevant documentation of GEOS-Chem version and other previous papers where this tagging method has been used previously if any, for attributing sulfur species to its source origins.

The description of the tagging method has been added in Sect. 2.3.

Tracer-tag technical in GEOS has been widely used in aerosol and gas studies (Bian et al., 2021; Nielsen et al., 2017; Strode et al., 2018) and in supporting various aircraft field campaigns such as ARCTAS, KORUS-AQ, ATom, CAMP2Ex, ACCLIP, and more. Such techniques are also adopted in other models such as GEOS-Chem model (Fisher et al., 2017; Ikeda et al., 2017) and CESM (Butler et al., 2018).

R2#5: Could you also say something about the contribution from shipping in the Pacific and Atlantic? Have you considered ship-based emissions also as anthropogenic emissions in your models? Would ship emissions which are also anthropogenic emissions be misread as anthropogenic emissions that are generally thought of as land-based emissions? Please make the necessary adjustments in their manuscripts at the locations wherever applicable (mostly sections 2.2 and section 5) and specify those changes in response to this comment.

Shipping emissions are concentrated between the tropics and mid-latitudes of the Northern Hemisphere over the Pacific and Atlantic Oceans. In our SO4 source analysis (Sect. 5), we discussed the contribution of various emission types (anthropogenic, biomass burning, volcanic, and marine) to atmospheric SO4
over remote oceans. Here shipping emission was considered as anthropogenic emissions. We did not further differentiate the location of anthropogenic emissions in this study. We specifically pointed this out in Sect. 2.2 (P5L231-233) and modified accordingly the Abstract P1L44-46, Introduction P3L130-132, main text P13L852-853, and Conclusion P15L929-930.

R2#6. Since you have used this tagged approach, I would also recommend that you specify the list of tags that you use. This could be either as a list in the text in section 2.2. Four tags we used in GEOS for this study relate to emission types of anthropogenic, biomass burning, volcanic, and oceanic emissions. This list is now given in Abstract P1L46-47 and Sect. 2.3.

R2#7: You could also add some future recommendations related to the scope of tagged simulations. With this approach we could also understand the contribution characteristics of emissions from various regions: both oceanic regions (Pacific, Atlantic, Sulfur Emission Control Areas etc.) and continental regions’ (Asia, North America, South America, etc) sulfur emissions. These recommendations could be added either in section 5, where the tagged model’s results are being discussed or in the concluding section (section 6).

Tags can be linked to various emission types (anthropogenic, biomass burning, etc.) and/or various emission regions (Pacific, Atlantic, Asia, North America, etc.), given that the contribution of each emission type/region to the total aerosol mass is also not interfered by that of other emission types/regions. The tagged simulation is a powerful tool that is widely used not only in aerosol components (Bian et al., 2021; Ikeda et al., 2017), but also in tracer gases such as CO (Fisher et al., 2017; Strode et al., 2018; Nielsen et al., 2017), CO2 (Lin et al., 2020), and O3 (Butler et al., 2018). This technique is particularly useful when supporting aircraft field campaigns (ARCTAS, KORUS-AQ, ATom, CAMP2Ex, ACCLIP, and etc) for flight planning due to its computational efficiency and its characteristic of tracking the origin of plumes linking to emission type and location. We have added this discussion in Sect. 2.3.

Reviewer 3 (R3)

R3#1. The authors lack to highlight what is new in this study compared to previous literature. This is evident in the Introduction, where there is no mention of results from previous/recent studies using the ATom measurements nor using the AeroCom models. This is a major concern because the manuscript, as it is, looks like a comparison exercise with no follow-up message.

Thanks for this comment. In the introduction, we have added a paragraph (P3L134-141) summarizing relevant previous/recent studies using the ATom measurements and the AeroCom models.

R3#2. Lack of “what’s new”. The manuscript does not convey the message of what is new compared to previous literature. The Introduction needs to contain references to previous literature using the Atom measurements and the AeroCom models for studies of sulfur-containing species over land and ocean. I think that one paragraph for ATom and one for AeroCom would suffice. The Abstract and the Conclusions must also state what is new in the study.

Yes. We have added a paragraph reviewing the literature and noting "what's new" in P3L134-145 (see also reply to R3#1).

Our abstract now includes this sentence (P1L47-49): “Our work presents the first assessment of AeroCom sulfur study using ATom measurements, providing directions for improving sulfate simulations, which remain the largest uncertainty in radiative forcing estimates in aerosol climate models.”
The Conclusions now include this paragraph (P15L946-956): “Even after two decades of development, the diversity of sulfate simulations from AeroCom-I to AeroCom-III has not decreased. However, accurate sulfate simulation in current climate models is crucial to reduce radiative forcing biases. Several potential directions for improving sulfur simulations are suggested above. More importantly, apart from the shortcomings of individual models, all modelers should focus on the calculation of the air-sea exchange flux formula, as it plays a key role in determining DMS emissions. Modelers also need to study DMS and SO2 vertical transport as well as SO4 wet deposition during long-distance transport, as model biases are greatest at high altitudes. One suggestion to modelers is that the use of online oxidant fields is insufficient to explain the model sulfate bias, as there was no systematic bias in the sulfate simulations between the models using interactive oxidants and the models using archival oxidants in this study. The complexity of chemistry deserves more attention.”

This study can be relevant if, for instance, it is the first using the ATom airborne measurements. If this was the case, it must be emphasized considerably more in the Abstract, the Introduction (like OP1L93-97 but much better/stronger) and the Conclusions (and throughout the text).

No, ATom measurements have been used in various studies. The ATom aerosol studies include NPF in the tropics (Williamson et al., 2019), fine aerosol lifetimes (Gao et al., 2022), and various aerosol components such as sea salt (Bian et al., 2019), OA (Hodzic et al., 2020), smoke (Schill et al., 2020), mineral dust (Froyd et al., 2022), and DMS chemistry (Fung et al., 2022). See P3L134-140.

This study can be relevant also if a comparison between AeroCom models and ATom measurements has not been done before. In this case also, the authors should emphasize the originality of such intercomparison.

Our work is the first study to use ATom measurements for comparison with the AeroCom models, focusing on all sulfur species simulated in current aerosol climate models. This work extends a previous effort that utilizes ATom measurements for comparison with AeroCom models for organic aerosol (Hodzic et al., 2020). See new paragraph in P3L134-145.

R3#3. Conclusions. Based on the major comment above, the Conclusions need to be strengthened. Once the novelty of the study will be identified (again, first intercomparison?), the Conclusions must be modified accordingly. Here follow the two main conclusion points that need to be addressed:

1) It seems to me that the authors conclude that the dominant source of atmospheric sulfur (SO2 in particular) over oceans is anthropogenic. It is well known that the largest contributions to the sulfur budget (SO2 included) is due to human activities. Hence, despite the short lifetime of the sulfur compounds, it does not seem so striking that the major source of atmospheric sulfur is anthropogenic even over oceans (e.g., Chin et al., 2000). Thus, I have the impression that this conclusion does not add anything significantly new to the current knowledge. In order to make this conclusion more appealing, I suggest increasing its relevance by emphasizing its unicity. I couldn’t find any other modeling study asserting that the sulfur over the oceans is mostly caused by human activities. This means that the authors are the first proposing that conclusion (as far as my literature research goes), and that is certainly worth mentioning. However, I suggest to carefully check the literature for modeling studies looking at the atmospheric sulfur over the oceans. If the authors find a relevant study, it would be interesting to compare their findings with yours.

The authors also could not find other modeling study asserting that the sulfur over the oceans is mostly caused by human activities. We changed the conclusion sentence of “On the other hand,
our study shows that anthropogenic emissions remain a major source of sulfate aerosols
generated over remote oceans ...” to “On the other hand, our study is the first asserting that
anthropogenic emissions remain a major source of sulfate aerosols generated over remote
oceans ...” (P15L941-944).

2) Another conclusion consists in the description of the significant differences between the
observations and the models. This is somehow discussed in the text with the attribution of the
differences to emissions (DMS, e.g., OP9L379-394), chemical conversion (SO2 to SO4, e.g.,
OP10L449-451) and transport (SO2, e.g., OP8L365-367) depending on the model, with a nice
final discussion (OP13L584588). However, these parts (especially OP13L584588) can be explored
further with a more detailed discussion of the models. Here’s a few ideas for further discussion:
1) what are the common aspects among the models? and their differences? Can we discern
something from similarities/differences 2) It is possible/feasible to perform additional sensitivity
tests to assess model performances (not for this manuscript but as material for discussion)? 3)
What would be the added value of satellite data in this comparison (again, for future studies):
would we learn something more thanks to the denser temporal and spatial sampling compared
to airborne measurements)?

Thanks for these excellent suggestions.

Regarding recommendation 1: We examined the DMS emission derivation methods used in five
AeroCom models (see the newly added Table 3 and the corresponding discussion in P9L573-581) and
found that the parameterization of air-sea exchanges in DMS flux calculations is very important P9L589-
648). We also recognized that using interactive oxidant calculation is insufficient to account for the
model sulfate bias (see P8L473-550).

Regarding recommendation 2: Yes. Several sensitivity tests have been suggested in Conclusion P14L910-
915.

Regarding recommendation 3: In-situ aircraft measurements provide intensive and extensive fields that
can be used to improve model simulations. However, aircraft measurements are limited by space and
time. On the other hand, satellite data coupled with global long-term measurements can provide an
independent assessment of model improvements. In this way, we can use the model as a bridge
between aircraft and satellite measurements. Denser temporal and spatial sampling of aircraft
measurements can provide tracer vertical profiles, which are critical for modeling tracer global
distribution and assisting satellite AOD/AI retrievals.

Additionally, aircraft measurements can provide detailed component measurements that cannot
currently be retrieved from satellite measurements. More intensive temporal aircraft measurements can
also provide tracer diurnal cycles, which are important for air quality studies.

3) Within the Conclusions, the authors can also dare to provide some outlook for future model
developments by suggesting ways to improve the current models (e.g., update the chemical scheme,
increase horizontal/vertical resolution, identify and try to improve relevant parametrization for
transport).

Several potential directions for improving sulfur simulations were suggested in Conclusion P14L910-915.
We also added the following sentences (P15L949-956) to the Conclusion. “More importantly, apart from
the shortcomings of individual models, all modelers should focus on the calculation of the air-sea
exchange flux formula, as it plays a key role in determining DMS emissions. Modelers also need to study
DMS and SO2 vertical transport as well as SO4 wet deposition during long-distance transport, as model biases are greatest at high altitudes. One suggestion to modelers is that the use of online oxidant fields is insufficient to explain the model sulfate bias, as there was no systematic bias in the sulfate simulations between the models using interactive oxidants and the models using archival oxidants in this study. The complexity of chemistry deserves more attention.”

R3#4. Tracer-tag. It would be very helpful if the authors added more detail to the methodology of the tracer-tagging concept used to obtain the source origins shown in OFig. 13. My understanding is that these sensitivity tests were conducted by the GEOS developers and made available to the authors. Even though the authors did not perform the experiments, the concept of tagged data needs to be explicitly discussed (currently it is superficially mentioned in Section 2.2) so the non-expert reader can understand the logic behind such sensitivity test.

The tracer-tag concept is now described in Sect 2.3 “Tag-tracer study in GEOS”. Please also see response to R2#1 - #4 and R2#6 - #7.

R3#5. Figures. The figures are generally too busy – there is a lot of information that is not always entirely discussed. For example, the AMS measurements at 1s and 60s in OFig. 2 are barely discussed. In that regard, the authors state that they use 10s merged data for AMS in the text unless otherwise stated (OP6L214-242). Based on that, I suggest removing the 60s and 1s lines from OFigure 2 (maybe moving the 1s, 10s, 60s discussion to a separate paragraph in the methods?). This way, the authors could merge the observations and models together in the same panels (like OFig. 3) and make OFig. 2 easier to read.

The information of 60-s and 1-s in OFig. 2 and corresponding discussion has been moved to the Supplementary Information.

Because of the complexity of the figures, the discussion is often confusing in the sense that the authors do not specify any panel when presenting the results. For example, the analysis of Fig. 2 (OP5L212-P6L264) never mentions one panel specifically. This makes it difficult to discern which flight (ATOM1-ATOM4) is considered.

Since the discussion follows the number of ATom deployments given in the panels, we removed the panel indices from Fig. 2.

In addition, the box-and-whisker panels are seldom mentioned, and they can be probably removed from there (and maybe merged and moved to Section 2 or moved to the supplement?). As a general rule, if a panel/figure is not discussed, it should not be shown.

Discussion L350-355 is based on the information shown in the box-and-whisker panels.

For OFigure 3, the statistics are briefly mentioned in OP6L280-282. Please provide more discussion of the statistics shown in OFig. 3 or consider moving the figures/panels to the supplement, given the large number of figures in the manuscript.

The discussion has been modified as “Statistics indicate lower model SO2 medians than observed (box-and-whisker in Fig. 3), especially during ATom-1. However, the model means are comparable or even higher than those observed, indicating that the models simulate unobserved episode events. Consequently, the simulated mean/median ratio is higher than the observed value. Across the four ATom deployments, ATom-4 has much better model observation consistency.” See P7L385-389.
Figure 4 also contains a large number of panels – (and the box-and-whisker panels are again shortly 
mentioned). As my comment above, please provide more discussion or move the panels that are not 
discussed in the Supplement. Furthermore, the addition of panels c,f,i,l breaks the continuity with 
respect to the previous figures, which contain only PDFs and statistics. These panels should belong to a 
separate figure (and paragraph) that discusses the vertical information from the observations/model 
comparison (maybe merged/moved to Section 3.2?).

The discussion of P8L435-441 is based on the statistics shown by box-and-whisker panels. We pointed 
out this in the revised version.

The OFig. 4 has been separated into two figures: one contains PDFs and statistics (Fig. 4) and the other 
contains original panels c,f,i,l (Fig. 5).

A final note on the box-and-whisker panels: if the authors think that the statistics should be shown, 
maybe it can be shown as tables depending on the sulfur species instead of panels in the figures.

To avoid the graph being too busy, we removed the values of median and mean from the graph. 
However, we do think that summarizing the various statistics in charts can help readers navigate the 
data more easily.

For OFigures 9-11, I would suggest to average together the horizontal regions that do not differ much 
one from another (e.g., Fig. 9a, A40S-20S with A70S-40S) and adapt the discussion accordingly.
The sulfur species can differ much between A40S-20S and A70S-40S depending on the season. For 
example, the Atlantic free tropospheric SO2 over the A40S-20S in ATom-4 is much higher than over the 
A70S-40S (Fig. 10d). Such performance is also visible in SO4 (Fig. 9d) but to a much smaller extent. 
However, this behavior is not shown across the entire vertical column of DMS (Fig. 11d), indicating that 
a marine source is not the cause. Considering that A40S-20S is located in the downwind area of South 
America (SA), SA pollution showed an impact on SO2 in the marine atmosphere of A40S-20S. See 
discussion in P11L719-723.

R3 Minor Comments.
• OP1L1: “[... and ecosystems.”. Could you add one or two sentences on how the sulfur cycle plays 
the key role (e.g., pollution, acid rain)?

Is this typo for P1L34? Changed the sentence to “The sulfur cycle plays a key role in atmospheric air 
quality, climate, and ecosystems, such as pollution, radiative forcing, new particle formation, and 
acid rain.”

• OP2L62-63: “wreak havoc”. Could you replace this expression with something like “devastate, 
destroy”?

Done.

• OP2L83-84: “the TUT and above are observation-sparse regions”. I would rephrase this part as 
something like “...observations in the TUT region and above are sparse”.

Done.

• OP2L91: Is it possible to specify what “DC–8” means?

DC-8 means Douglas DC-8 jetliner. See change in P3L116.

• OP3L123: “(i.e., “0.2-12 km) (Thompson et al., 2021)“ --> (i.e., 0.2-12 km, Thompson et al., 2021).

Done.

• OP4L148-150: I recommend rephrasing this sentence to something like “Two instrument were used 
[...]: the California [...] and the NOAA [...] (Table 1)”. 
• OP4L177: “(CMIP6) (Feng et al., 2020)” → (CMIP6, Feng et al., 2020).
  Changed the sentence to “The suggested emissions are the Coupled Model Intercomparison Project Phase 6 Community Emissions Data System (CEDS, Hoesly et al., 2018) ….”. See P5L224-227.

• OP4L178: “(…. System (GFAS))” → (… System, GFAS).
  Done.

• Section 3.1: I suggest homogenizing the discussion here. As it is, OFig. 3 is less discussed than OFig. 2 and OFig. 4. The authors could enhance the discussion for OFig. 3 with the addition of a couple of paragraphs on SO2 or reduce the discussions of the OFigs. 2 and 4.
  The discussion of Figs. 2 and 4 has been reduced in Sect. 3.1.

• OP6L252-264: The Atom flights are not discussed separately for OFig. 2 (SO4). I suggest providing some discussion about the different flights to maintain consistency with the following discussions of OFigs. 3 and 4 (where the different flights are mentioned).
  We added discussion of different flights in P7L363-366. We also reframed the paragraph by discussing two measures first. This style is consistent with the style used in SO2 and DMS (i.e., Figs. 3-4) discussions.

• OP7L305: “OFig. 4c,f,i,l”. This is the only place of Section 3.1 where specific panels are mentioned. Since no other panel of OFigs. 2, 3 or 4 is explicitly discussed, I suggest 1) either discussing explicitly the remaining panels or 2) removing the explicit mention to OFig. 4c,f,i,l and move these panels into another figure (as suggested in my major comment about OFigures above).
  Following the reviewer’ suggestion, we have removed the indices of the panels in Figs. 2-4 and discussed the figures following ATom-# given in the figures. We also extracted OFig. 4c,f,i,l and put them in Fig. 5 to maintain continuity among Figs. 2-4.

• OP7L309: I suggest removing “Apparently”.
  Done.

• OP7L317: I suggest rephrasing “...is uniquely having a....” to “...... is unique because it has a ......”
  Done.

• OP8L344-345: I suggest rephrasing “Despite that improvements are needed, ...” to “Despite the need for improvements, the models are generally able to capture the shape of the SO4 profile.”
  Done.

• OP8L346-354: I find the introduction of the model Groups 1, 2 and 3 a little confusing (also, these groups are not mentioned again throughout the manuscript). I suggest keeping the model names as there are only 2 models per group and Group 2 consists only in E3SM.
  Removed the group categories, see P8L469-473. Also see R1#9 and R3#3.

• OP8L358: I suggest rephrasing “.... an order of magnitude, but around ....” to “.... an order of magnitude around ....”.
  Done.

• OP8L359: The authors say that SO2 is better in IMPACT in the NH and CAM-ATRAS and OsloCTM3 in the SH, but compared to what? Please specify the reference (I suppose that would be GEOS and E3SM?).
  Compared to other AeroCom models not mentioned in the sentence, see P9L553-555.

• OP8L360-361: Please be more specific with regard to how the models should improve the SO2 simulations (too large/little concentrations wrt observations?). I understand that the differences are thoroughly discussed in the following sentences, but it would be nice to have a preliminary “hint”.
Yes. The sentence is changed to “The tropical Pacific appears to be an interesting region, with all models except GEOS failing to capture observed local SO2 sources.” See P9L556-557.

- OP8L369-370. “All models [...] ATom-1 observation”. I do not quite understand what the authors mean with this sentence. Could you please explain it?
  The sentence is changed to “All other models show lower SO2 at the surface than in the lower free troposphere, which is inconsistent with the observed profiles.” See PL9565-566.

- OP8L371-375. Since this part refers to a figure in the supplement (OFig. S5), I suggest reducing this discussion here.
  The following sentence (OP8L372-375) was moved to the Supplementary Information: "Modeled SO2 volume mixing ratios are generally lower compared to the CIMS observations for most altitudes and latitudinal bins in ATom-1 to -3, which may be partially owing to the CIMS measurement issue discussed in Sect. 3.1."

- OP9L384: “The parameterization...”. Do the authors refer to the parameterization of Nightingale et al., (2000)? If yes, I suggest rewriting the beginning of the sentence with something like “That/This parameterization ...”.
  The paragraph has been rewritten to response R3#3, so this comment is no long relevant in the revised version.

- OP9L396: “... than the observed one,” I suggest providing an example of a panel showing this steeper gradient (e.g., OFig. 7 54N-90N Atlantic). It would be nice to add panel numbers/letter to these vertical profiles to facilitate the navigation during the discussion.
  We added "(e.g., Fig. 8 A54N-90N)" to P10L650-651. However, if we add panel numbers to these vertical profiles, we will end up mentioning only one number in the text, which goes against the reviewer's suggestion (see P7L305 above).

- OP9L399: I suggest rephrasing “tease out” with “obtain”.
  Done.

- OP9L396-410: Very nice discussion!
  Thanks.

- OP9L414: I suggest rephrasing “These behaviors are inconsistent with...” with “These patterns do not agree with...”.
  Done.

- Section 3.3: Concerning the definition of the altitude ranges, I suggest keeping the same nomenclature both in the text and in the figures. Specifically, in the text the authors use words like “free troposphere” or “boundary layer (BL)”, while in the figures they show only altitude ranges in kilometers (0-1.5, 1.5-6, .... km). My suggestion is to add the name of the layer to the range in kilometer in the Figures (e.g., “0-1.5 km (BL)”, etc.).
  Done.

- OP10L422: I suggest explaining what the Figures contain. Something like “In order to analyze [...], Figs 9-11 shows histograms of XX concentrations as a function of altitude (rows) and latitudinal band (columns)”.
  The sentence has been changed to “In order to analyze model performance on a regional and seasonal basis, Figs. 10-12 show histograms of SO4, SO2, and DMS concentrations as a function of altitude (rows) and latitudinal band (columns).” See P10L676-678.

- OP10L432-433: “The most high [...] (NH spring).” Could the authors specify the altitude range here?
  The altitude range for free-troposphere refers to 1.5 – 12 km, see change in P10L684-685.
• OP10L438: I suggest rephrasing “more polluted” with “tends to simulate higher SO4 concentrations”.
Done.

• OP10L438-440: “SO4 concentrations [...] the Supplement.” I suggest expanding this part by moving here the relevant parts of the Supplement.
The relevant discussion of the Supplement (see below) has been moved to the main text.

P11L711-715: “For example, in summer and winter, the CAM-ATRAS model gave the highest estimates of atmospheric SO4 in the oceanic BL, but the IMPACT and OsloCTM3 models gave the highest estimates of atmospheric SO4 in the free troposphere (Fig. S9). All models except the GEOS model generally overestimate SO4 in the atmosphere.”

P11L727-732: “For example, the E3SM model gives significantly higher SO2 compared with the measurements and other models in BL (Fig. S10). Unlike the case of SO4, all models tend to underestimate SO2 in the free troposphere, with some exceptions such as the GEOS model in the North Pacific mid-high-latitude winter (ATom-2) and the CAM-ATRAS and IMAPCT models in the South Atlantic mid-latitude autumn (ATom-4).”

P11L745-747: “The overestimation of the DMS multi-model median in Fig. 12 is clearly attributable to the contribution of all models shown in Fig. S11, with the models CAM-ATRAS and OsloCTM3 being more prominent.”

• OP10L429-440: There is no mention of the difference between Pacific and Atlantic regions. I suggest adding a couple of sentences discussing this separation.

We added following sentences in P10L692-709: “Compared to observations, model tends to simulate higher SO4 concentrations in the free tropospheric atmosphere. Both observations and simulations show that the SO4 in the Pacific is higher than that in the Atlantic during the NH high-latitude autumn (ATom-3) and the NH mid-latitude spring (ATom-4). The differences between observations and simulations are generally larger in the Atlantic than in the Pacific, particularly in the SH.”

• OP10L443-444: “..... and pollution affects ....” It is not clear to me what the authors mean with pollution. Could the authors explain?

Changed “pollution affects” to “this high SO2 region extends to”. See P11L718-719.

• OP10L444: I suggest changing “Areas where free tropospheric SO2 pollution is relatively polluted ....” to “Areas where free tropospheric SO2 concentrations are relatively large .....”.
Done.

• OP10L442-451: there is no mention of ATom 2 in this paragraph. I suggest including one sentence discussing the most relevant pattern for ATom 2.
The discussion for ATom-2 has been added in P11L720-723. “For instance, free troposphere appears to be more polluted than other regions in the NH Pacific during ATom-2 and in the SH mid-latitude Atlantic during ATom-4, but not in the BL, implying a potential source of horizontal transport.”

• OP10L454: I suggest rephrasing “.... when the hemisphere is in spring” with “.... during springtime”.
Done.

• OP10L458-459: This sentence concerning the BL sounds a bit redundant with part of the sentence before (P10L457, discussing the free troposphere) except for the point (3). I suggest merging the discussions together to avoid repetition.
Removed this sentence (OP10L457-458) to avoid repetition: “... suggesting a potential slower vertical transport or faster DMS chemical loss in models.”
• OP10L460: I suggest rephrasing “The convoluted effort can be somehow alleviated by ....” with “Additional insights can be obtained by ....”.
  Done.
• OP10L463: I suggest replacing “giving” with “because”.
  Done.
• OP10L453-465: Also here, very nice discussion.
  Thanks.
• OP11L468: I suggest adding a sentence or two explaining the need for this Section (why it is important to look at the sulfur budget in model in light of what was shown before and what will come in Section 5).
  We added following two sentences there: “Budget analysis is a simple and basic method that has been widely used to document the underlying performance of a model. This analysis allows us to evaluate the AeroCom-III sulfur simulations against previous AeroCom-I and -II studies and reserves a record for future model evaluations.” See P11L750-768.
• OP11L489: I suggest rephrasing “… are pretty much the same as they should be” with “… are as expected”.
  Changed the sentence to “source and deposition are pretty much the same as expected”.
• OP12L524: I suggest replacing “… is at its most active” with “is the largest”.
  Done.
• OP12L525: I suggest removing “and event”.
  Done.
• OP12L530: I suggest removing “where and”.
  Done.
• OP12L536 “… continental areas” I just have a question: is there any anthropogenic source of sulfur over the oceans (ships?). If yes, it would be nice to mention it.
  Done. See P13L852-853.
• OP12L554: “It has a clear…” What does “It” refer to? Tropospheric SO4? I suggest the authors to specify here.
  Changed “It has a clear …” to “There is a clear …”.
• OP12L556: “There are two …” This sentence needs to be introduced. Something like “Concerning volcanic sources, emissions from volcanoes are of two types.”
  Done.
• OP12L556-P13L561: The two types of volcanic contributions need to be separated more clearly. I suggest something like “One type is the volcanic degassing […]. The other type consists in the volcanic eruptions […].”
  Done. See P14L882-886.
• OP13L560: I suggest removing “eruption”.
  Done.
• OP13L571: I suggest adding “airborne” before “ATom”.
  Done.
• OP13L571-573: This nice sentence is the goal and “what’s new” in the manuscript. I suggest expanding this in the Introduction with references to previous studies (and a similar sentence in the Abstract). See my major comments on this aspect.
Yes. We have added a paragraph reviewing previous relevant research in the Introduction section (P3L134-145). We have also added Table 5 and corresponding discussion (P12L807-823) comparing the diversity of global sulfate atmospheric loads from multiple AeroCom models since Aerocom-I. See also responses to R1#5 and R3#2.

- OP13L590: “over remote oceans” Again, this is the goal and should be stated more clearly in the Introduction.
- These key words have been mentioned already in Abstract (P1L40), the Introduction (P2L105-106, L121, L128-129), and model experiment design (P5L250).
- OP13L599: “… this proportion is increased …” Do the authors know the amount of the increase?
- This SO4 increased by explosive eruptions varies in location and timing.
- OFigure 9: In the caption, I suggest adding the seasons corresponding to the different Atom flights. Done.
- OFigure 10: The contribution from biomass burning (BB) is hardly discussed in Section 5 (which is logical!). Therefore, I suggest renaming BB to something like OTHER and state that it includes biomass burning (and any other sulfur source) and that it will not be discussed in Section 5 because its contribution is negligible compared to AN, VOL and OCN. Done. See P13L856-857.