

Answers to Referees. We would like to thank you for your efforts to improve our manuscript by your comments. Answers from the authors in blue text. Line references of the authors refer to manuscript prior to the present revision.

**Referee 2:**

This manuscript reports on measurements of aerosol sulfur in aerosol samples collected from the IAGOS-CARIBIC platform over a 16 year period. Analysis focuses on a new regression technique that the authors suggest can be used to infer both the gradient of sulfur concentration and the integrated burden in the lowermost stratosphere (LMS), starting at the dynamical tropopause and extending to 3 km above it. It is also suggested that this analysis provides an estimate of the relative contribution of stratospheric sulfate mixed downward and tropospheric sources on the sulfur concentration in the upper troposphere (UT), and how these contributions vary seasonally. Compared to the original version, changes in the data processing now produce more pronounced seasonality in the estimated contribution of S derived from tropospheric sources to the S burden in the UT. In particular, the analysis now suggests a quite strong peak which authors attempt to link to downward transport from the ATAL.

While the approach has changed in response to comments on first version, and some sections describing the data analysis are now more clear, I still find myself not very convinced. Step 1, doing forced linear regression (FLR) of S versus distance above the tropopause is relatively straight forward, except for one key point. The data “were grouped with respect to concentration levels of the different years, resulting in 4 to 5 groups of data for each season” (lines 179-180). The discussion about this step seems to justify at least 2 groups (background and influenced by volcanoes), and perhaps the volcanic group could be (was?) further divided into strong and weak influence. It seems that the authors recognized that there had to be more than 2 groups in each season to do step 2 regressions, but how these were created is not explained. Some objective method of creating these groups, perhaps based on mean, median, max measured S concentrations, or maybe based on time since eruptions were known to have perturbed the LMS burden (based on previous work by this group using largely the same data set) would seem essential, especially for anyone attempting to replicate the analysis or apply the technique to some future similar data set. As it reads now, it almost seems that the approach was to sort by year based on timing of volcanos, but then move years (or individual samples?) that didn’t work into another group until things looked better

It would be difficult to use a priori knowledge to classify the effects of a given volcano. We still miss accurate inputs to global models that describe the distribution and the evolution of volcanic injections to the stratosphere (<http://volmip.org/>). We did not have one volcanic group because the eruptions vary in strength and distribution. Fig.4 shows two examples where you can see that the “background” and “moderate tropical volcanism” groups are closely spaced, but still clearly separated, see e.g Friberg et al in Tellus 2014. The need for further groups of data is obvious (Fig. 4) to cover the strong influences of e.g. Kasatochi, Sarychev and Nabro.

To clarify, we have added (line 183): “The remaining data of each year were tested for systematical differences. Those years where the data overlapped in particulate sulfur – height above the tropopause space, were grouped together.”

The next step in the analysis is now basically clear conceptually: for each season the intercept and slope from 4 or 5 FLR in step 1 were regressed against each other, using weighted regression with the weighting based on estimated uncertainty in the FLR fits. The explanation of this weighting is not completely clear to me, but seems to generate uncertainty only in the FLR intercept (termed offset) when transferred to Fig 3 which shows results of the weighted regressions by season. Surely, the slopes from step 1 were also uncertain. It would seem that uncertainty in both slope and intercept from the FLR would propagate into uncertainty in the estimated slope and intercept from the weighted regression. Based on the small number of data points and the magnitude of the error bars in all 12 panels of Fig 3, I find the small error bars in Fig 5 very surprising (seems that error bars in 5 a should come directly from uncertainty in the slope from fits in Fig 3, while those in 5 b come from uncertainty in the intercepts in Fig 3).

Sorry but we do not understand this comment. Fig. 5a collects the slopes obtained from the regressions shown in Fig. 3, and Fig. 5b collects the intercepts from the same figure. The errors of the two quantities are computed and presented in Fig. 5. This is clearly explained in the manuscript.

In particular, the authors need to carefully consider how to propagate uncertainty from both initial measurement, uncertainty in  $Z$ , and uncertainty in slope and intercept of the FLR to the intercept values for the weighted regressions in Fig 3 and clearly explain how they do this, since they assert that the derived values represent the concentration of S in the UT that is derived from tropospheric sources and use these numbers for most of the remaining analyses in the paper. Lines 244 – 249 address how they attempted to account for the uncertainty from FLR, but not in very clear manner.

Of course the errors from step 1 very much affects the results in step 2. Imagine that we substantially increase the errors in step 1. The first visible sign would be that the data points would be more scattered in step 2, because the errors are larger. That would affect the step 2 regression and induce larger errors in the estimated  $C_{S,UT_{trop}}$  and  $k$ . The second way the results of step 1 enters step 2 is by using the errors from step 1 as weights, where data points with small variance affects the regression more than those with large variance. If you are still not convinced, try to draw alternative fits in fig 3, minding the size of the error bars, and you will see that the errors given for  $C_{S,UT_{trop}}$  and  $k$  are highly realistic.

The referee is right in that the slope estimates also have uncertainties. The average variance ratio was 0.19, i.e. the relative variance of the slope divided with that of the intercept from the step 1 FLR. The slope errors thus are small compared to those of the intercept, and were therefore neglected.

On line 237 the following was added: The relative variance of the slopes was much smaller than that of the offsets (average variance ratio of 0.19). For simplicity, the variance of the slope was therefore neglected in these regressions.

If the seasonality shown in Figures 5 and 7 is real (signal truly larger than uncertainty) that is an interesting finding.

Glad you found something interesting about the manuscript. Concerning the “if”: Please consider Fig. 7a where you find the S in UT of tropospheric origin together with the UT concentrations during “background” and “moderate influence from tropical volcanism”. The latter two sets of data are straightforward averages of measurements in the UT obtained without any regression. First look at the first half of the year. The average of the slightly volcanically influenced case clearly exceeds the background case as a result of the added volcanic aerosol. We know that the Junge layer exists also during background conditions and gives rise to transport of S in this half of the year. In Fig. 6c (or, if you suspect circular evidence, in Fig. 6a in Friberg et al., Tellus 2014) you can see that the tropical volcanism in 2005 – mid 2008 almost doubled the S column compared the background conditions (1999 – 2002), and in Fig. 7b you can see that approximately a factor of 2 appears between the two cases in the first half of the year. Consequently, the tropospheric component should be clearly below the background case during this time of the year, otherwise our results would disagree. In the fall, we see that the volcanic and background data almost coincide (Fig. 7a). Although the stratospheric concentrations are elevated they do not affect the UT concentration very much (Fig. 4b), since we are not in the main season of the deep Brewer-Dobson transport to LMS and UT. It is therefore no surprise that the tropospheric component is more important in the fall. The surprise for us was that we, in addition to the spring maximum, have a second maximum in S concentration of the UT in the fall, a result we obtained by direct averaging of the data taken in the UT. The results of the part of the analysis that you criticize fits very well into the frame obtained by simpler means. We have clearly explained the methods used and errors involved in the different computations. Further explanations have been added to this second revision of the manuscript, see above. There are no reasons to question our results.

Going to CALIOP data to try to explain the fall peak in tropospheric influence on S in UT is also a good idea, but I am not sure how much support the progressive descent of the ATAL in the top row of Fig 8 provides to the authors interpretation of CARIBIC UT/LS S. How much does the ATAL spread zonally over just a few months? Would the upper level anticyclone associated with the Asian monsoon not keep the enhanced aerosol more or less over Tibet?

Have a look in Figs. 8 f – j and Yu et al. in PNAS 2017 ([www.pnas.org/cgi/doi/10.1073/pnas.1701170114](http://www.pnas.org/cgi/doi/10.1073/pnas.1701170114)) for answers to your questions. Here is a quote from the Yu et al. paper: “Between August and December a large portion of ASM aerosol moves poleward following the lower branch of the Brewer–Dobson circulation. The aerosol is eventually flushed out of the stratosphere at high latitudes.” (ASM = Asian Summer Monsoon). We take the liberty to interpret “high latitudes” in the meaning high latitude compared to the initial position of the ATAL, because it is well established that the main transport out of the LMS occurs at

midlatitudes. With this clarification you see the full agreement between this statement and Fig. 8f-j as well as the explanations given in the manuscript.

Did many (any) of the CARIBIC flight pass through or near the 60-120E longitude band shown in Fig 8?

20% of the samples were taken east of 60° E. The entire ATAL region extends to approximately 15° E (J. Geophys. Res. Atmos., 120, doi:10.1002/2014JD022372.). 53% of our samples were taken east of 15° E. It should, however be pointed out that the effluents of the Nabro eruption occupied this region in 2011, see Fig. 3 in Bourassa et al. (Science 2012). In the manuscript we demonstrate the transport down to flight altitude, both by Caribic and Calipso measurements.

In response to this comment we clarified in line 443: “We therefore illustrate the subsidence using the eruption of the tropical volcano Nabro in the summer 2011 (Table 1).”, by adding ‘... 2011 (Table 1), with effluents that occupied approximately the same region as the ATAL, see Fig. 3 in Bourassa et al. (2012).’

In summary, I am still not convinced that the data analysis is robust enough to believe that from now forward every CARIBIC S measurement can be converted into an estimate of the S column 3 km above the tropopause (Fig 6), or that the source of S in the UT changes from almost completely stratospheric in FM to about 90% tropospheric in ON (Fig. 7). Authors need to use and describe an objective means to bin their data, and more clearly discuss how confident they are about the uncertainty in their derived estimates of “particulate sulfur concentration of tropospheric origin” (also known as the intercepts of the weighted regressions).

As pointed out several times already, we are very confident in the methodology used. We admit that a minor variance component was overlooked, but its effect is insignificant. The error estimates are obtained based on a solid methodology.

Minor editorial comments

Line 82 what is meant by “crust and particle-bound water”?

changed to: “crustal particles and enhanced signal caused by hygroscopic growth”

Lines 147-149 reword this to make it clear what is meant by “not temperature decrease in relation to surface of the earth or the degree of cloud processing” and how either of these is related to distance below the PV tropopause.

changed to: “... , and hence not an altitude-typical degree of cloud processing.”

Lines 178-194 This section is where the data binning and how 4 or 5 groups were objectively defined needs to be explained.

Please see the answer on this subject above.

Line 223 and equation 2, seems that year (y) should be group (g). If you really did the regression for 12 seasons and 16 years the number would be  $\gg 52$

Eqn. 2 is beyond the group stage. The slope depends on the year the sample was taken (mainly due to volcanism) and the season (air circulation).

Line 284 Fig 4--→Fig 4a

Thank you. Changed in the text.

Line 289 Fig 4--→Fig 4b

Thank you. Changed in the text.

Line 415-417 This sentence is describing the seasonality of “particulate sulfur concentration of tropospheric origin” not the concentration of S in UT. Figure 7 shows nearly no seasonal change in UT S concentration in background year, and a spring peak in the volcanic year.

Thank you. We have changed to: “... distinctive characteristics of the particulate sulfur concentration of tropospheric origin in the UT,”

#### **Referee 4:**

Review of Paper: Particulate sulfur in the upper troposphere and lowermost stratosphere – sources and climate forcing by Martinsson et al. 2017, submitted to Atmospheric Chemistry and Physics

Martinsson et al. studied sources regions of upper tropospheric sulfur using long-term IAGOS-CARIBIC data. The authors show that the relative contributions (either from transport below or mixed down from the stratosphere) in UT sulfur are dependent on seasons, volcanic activities. The paper has gone through the first round of review with substantial changes based on previous reviewers especially reviewer #1. In general the paper is in good shape, but some concerns (especially on tropospheric contributions) need to be addressed before publication.

a. In the abstract, please give more definition on UT and LMS, i.e. LMS is  $\sim 3$  km above dynamic tropopause; while UT is how many km below?

The UT in this study is defined by the measurement range of Caribic (99% of the aerosol samples were taken in the altitude range 8.8 – 12 km) and the position of the tropopause. Both the flight and tropopause altitudes vary for the different samples. The part of the LMS regularly probed by

the measurements also defined by the measurement altitude range and the variable tropopause altitude. We add the altitude range of the measurements in the abstract to clarify.

In line 11 we have added: "... stratosphere (LMS) of the northern hemisphere extratropics during monthly intercontinental flights at 8.8 – 12 km altitude of the IAGOS-CARIBIC platform in the time period 1999 – 2014."

We also changed in line 94 from "9 – 12 km" to '8.8 – 12 km' in order to provide the same resolution in data everywhere in the manuscript.

b. Line 25 "We find that tropospheric sources dominate during summer and the fall as a result of downward transport of the Asian tropopause aerosol layer (ATAL) formed in the Asian monsoon"

The submitted manuscript (Lines 23 – 24) reads: "We find that tropospheric sources dominate during the fall as a result of downward transport of the Asian tropopause aerosol layer (ATAL) formed in the Asian monsoon,". Hence no disagreement with the somewhat more detailed description in line 576.

Line 576 "The ATAL is transported downwards, and affects the extratropical tropopause region in August to December."

These arguments are unclear to me, what does "downwards transport" mean? A recent paper "efficient transport of tropospheric aerosol into the stratosphere via the Asian summer monsoon anticyclone" shows that ATAL is mainly lifted above locally to the stratosphere and further transport to high latitudes. See their Figure 4.

First of all (same as in an answer to referee 2): Here is a quote from the Yu et al. paper: "Between August and December a large portion of ASM aerosol moves poleward following the lower branch of the Brewer–Dobson circulation. The aerosol is eventually flushed out of the stratosphere at high latitudes." (ASM = Asian Summer Monsoon). We take the liberty to interpret "high latitudes" in the meaning high latitude compared to the initial position of the ATAL, because it is well established that the main transport out of the LMS occurs at midlatitudes. With this clarification you see the full agreement between this statement and Fig. 8f-j as well as the explanations given in the manuscript.

c. The same paper above argues that in summer/fall seasons, convections transport tracer gases (SO<sub>2</sub>) and aerosol to UT. Do you think it may be more likely a reason for higher tropospheric contribution of UT sulfur rather than downward transport of ATAL?

As can be seen in Figs. 8b – d, this is part of the ATAL (T for tropopause in ATAL). Fig. 8c covers the same period as Fig. 4 (top) of Vernier et al. (J. Geophys. Res. Atmos., 120, doi:10.1002/2014JD022372.). Just as our Fig. 8c, that of Vernier demonstrates that a large fraction of the ATAL is found below the 380 K isentrope. In that fig. they do not present the tropopause, but in our Fig. 8c you can see that the ATAL reaches all the way to below the tropopause. In the poleward transport the lower branch Brewer-Dobson circulation, as well as transport poleward in the LMS, occurs along isentropes bending towards lower altitudes. In Figs. 8f-j this downward transport can be identified together with a cross-isentrope transport amplifying the downward motion of the air.

We clarify in line 441, replacing “... stratospheric circulation is directed downwards in the extratropics, but the ATAL ...” with ‘... poleward circulation along isentropes bending downwards, further amplified by an extratropical cross-isentrope, downward component, brings the ATAL down to lower altitudes. The ATAL ...’.

d. “SO<sub>2</sub> measurements by the satellite-based instrument MIPAS indicate a UT seasonal variation in the NH midlatitudes with low concentrations in December to March and the highest concentrations in June to September (Höpfner et al., 2015).”

In my impression MIPAS SO<sub>2</sub> product is of large uncertainties, and may not be meaningful in UT. Shown in Rollins et al. (2017, GRL), MIPAS overestimates the SO<sub>2</sub> concentration in TP by a factor of ~3 or so.

We agree that the MIPAS data have large uncertainties (as pointed out also by Höpfner et al. 2015). The large discrepancies between their and Rollins et al. (2017) data are important in a general sense. However, in this manuscript we do not make any use of the numbers, only the seasonal variation reported in the Höpfner et al paper published in Atmos. Chem. Phys. two years ago.

In response we have made the following changes (line 422) from “SO<sub>2</sub> measurements by the satellite-based instrument MIPAS indicate a UT seasonal variation in the NH midlatitudes with low concentrations in December to March and the highest concentrations in June to September (Höpfner et al., 2015).”

to ‘SO<sub>2</sub> measurements by the satellite-based instrument MIPAS have recently become available (Höpfner et al., 2015). These data have large uncertainties (Höpfner et al., 2015), and likely overestimate the SO<sub>2</sub> concentration (Rollins et al., 2017). However, here we only qualitatively use the seasonal variation. The MIPAS results indicate a UT seasonal variation in the NH midlatitudes with low concentrations in December to March and the highest concentrations in June to September (Höpfner et al., 2015).’

e. I found the discussions on SO<sub>2</sub>, CO, OH are difficult to understand. Seems authors claim that direct transport of SO<sub>2</sub> and sulfate aerosol from lower troposphere is not going to explain the seasonal variation on tropospheric contribution to UT sulfur, because they will be washed out or cloud-processed. And the authors suggest the concentrations of OH and SO<sub>2</sub> are responsible. Well, are we certain that new TP, SO<sub>2</sub> oxidation is OH-limited? Any know seasonal cycle on SO<sub>2</sub> emissions can explain the seasonal variation? I think some surface SO<sub>2</sub> can survive in UT especially in deep convection.

In the manuscript (lines 407 – 414) we discuss the seasonal transport paths from the surface to the UT. Lines 415 – 429 discuss chemical species. The seasonal variation of the common pollution-tracer CO is found to differ from that of particulate sulfur, and we provide one explanation in the strong seasonal cycle of the OH family which produces particulate sulfur and consumes CO. We also discuss the seasonal cycle of the particulate sulfur precursor gas SO<sub>2</sub> (from the referenced MIPAS results) and find a covariation between increases of the SO<sub>2</sub> and particulate sulfur of tropospheric origin. In this part of the manuscript we try to find the plausible scenario behind our observations, which includes the interpretation that deep convection transports SO<sub>2</sub> to the UT

(Lines 424 – 425). Lines 440 – 468 cover the effects of the ATAL, which is also a result of deep convection.

In response we have added (lines 427 – 428): “The increase in  $C_{S,UT_{trop}}$  during the spring and summer months coincides with increases in the convective activity as well as in the concentration of both  $SO_2$  and OH, thus offering a plausible explanation.” However, the quantitative understanding of the  $SO_2$  transport paths and UT seasonality requires further studies.

Co-Editor Decision: Reconsider after minor revisions (Editor review) (02 Aug 2017) by Kostas Tsigaridis

Comments to the Author:

Dear Authors,

I have another set of review reports available, including a new reviewer, to get one more opinion, since the reports so far did not help me converge to a decision. I believe that the main problem the manuscript has is that the methodology is still fairly opaque, as described in the new reports. Please take into accounts the new comments and submit a revised manuscript.

Please also take into account that one of the reviewers (in a private communication) mentioned that there are still unexamined assumptions about the stratospheric contribution relation with the slopes and intercepts, and a rigorous look at those is needed. The uncertainty in the tropopause determinations is also rather uncertain and the impact of this uncertainty needs to be discussed.

Kind regards,  
Kostas Tsigaridis

Dear Editor,

We have thoroughly answered the comments from the referees and tried to clarify and improve the legibility of the manuscript on all places we found ways to further improve the manuscript. We hope that you now find the manuscript suitable for publication.

Sincerely,  
The Authors

# Particulate sulfur in the upper troposphere and lowermost stratosphere - sources and climate forcing

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10 **Abstract.** This study is based on fine mode aerosol samples collected in the upper troposphere (UT) and the lowermost stratosphere (LMS) of the northern hemisphere extratropics during monthly intercontinental flights [at 8.8 – 12 km altitude](#) of the IAGOS-CARIBIC platform in the time period 1999 – 2014. The samples were analyzed for a large number of chemical elements using the accelerator-based methods PIXE (particle-induced X-ray emission) and PESA (particle elastic scattering analysis). Here the particulate sulfur concentrations, obtained by PIXE analysis, are investigated. In addition, the satellite-borne lidar aboard CALIPSO is used to study the stratospheric aerosol load. A steep gradient in particulate sulfur concentration extends several kilometers into the LMS, as a result of increasing dilution towards the tropopause of stratospheric, particulate sulfur-rich air by tropospheric air forming the extratropical transition layer (ExTL). Observed concentrations are related to the distance to the dynamical tropopause. A linear regression methodology handled seasonal variation and impact from volcanism. This was used to convert each data point to standalone estimates of a concentration profile and column concentration of particulate sulfur in a 3 km altitude band above the tropopause. We find distinct responses to volcanic eruptions, and that this layer in the LMS has a significant contribution to the stratospheric aerosol optical depth and thus to its radiative forcing. Further, the origin of UT particulate sulfur shows a strong seasonal variation. We find that tropospheric sources dominate during the fall as a result of downward transport of the Asian tropopause aerosol layer (ATAL) formed in the Asian monsoon, whereas transport down from the Junge layer is the main source of UT particulate sulfur in the first half of the year. In this latter part of the year, the stratosphere is the clearly dominating source of particulate sulfur in the UT during times of volcanic influence as well as background conditions.

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## 1 Introduction

30 The global mean surface temperature has increased considerably in the two last years (NOAA, Dec. 5, 2016), which followed on a 15-year period with slow temperature evolution. CMIP5 (coupled model intercomparison project) models predict stronger than observed temperature increase in this period (Fyfe et al., 2013; Fyfe et al., 2016). Reasons for these differences were sought, and the Interdecadal Pacific Oscillation connected with increased subduction and upwelling (England et al., 2014; Meehl and Teng, 2014), variations in volcanic aerosol (Solomon et al., 2011; Santer et al., 2014) and solar (Myhre et al., 2013) forcings were identified as main causes of the discrepancies. These

35 phenomena are all elements of natural climate variability, thus highlighting the importance of these influences for assessing the human impact on the climate (Ramanathan and Feng, 2008).

Air from the tropical troposphere containing aerosol precursor gases is lifted into the tropical stratosphere in the Brewer-Dobson circulation. Carbonyl sulfide (OCS) is the most abundant sulfur-containing gas in the atmosphere. When lofted into the stratosphere, OCS is converted to sulfur dioxide (SO<sub>2</sub>) at approximately 25 km altitude aided by  
40 UV radiation (Crutzen, 1976) and in a next step to sulfuric acid, giving rise to the background stratospheric aerosol layer. The upward tropical flow is connected with a downward flow in the extratropics, where stratospheric air flows back to the troposphere.

The stratospheric aerosol concentration varies strongly due to volcanic influence, from events like the strong 1991 Mt. Pinatubo eruption, inducing negative radiative forcing in excess of 1 W/m<sup>2</sup> (McCormick et al., 1995), to close to  
45 background conditions at around the turn of the millennium (Bauman et al., 2003; Martinsson et al., 2005). Overall, anthropogenic influence on stratospheric aerosol is small compared to that from volcanism (Neely et al., 2013). Recent volcanic eruptions, such as in 2008 and 2009 of the Kasatochi and Sarychev in the extratropics and in 2011 of the Nabro in the tropics, significantly altered the stratospheric aerosol (Vernier et al., 2011a; Bourassa et al., 2012). These eruptions also had a profound impact on the northern hemispheric lowermost stratosphere (LMS) (Martinsson et al.,  
50 2009) and the global aerosol optical depth (AOD) of the stratosphere (Andersson et al., 2015). After reaching the LMS, the stratospheric aerosol is transported to the upper troposphere (UT). A recent study indicates that volcanic aerosol also has an indirect climate effect by affecting the reflectance of cirrus clouds in the UT (Friberg et al., 2015).

The UT particulate sulfur, in addition, has tropospheric sources. In the tropics, convection lofts aerosol and aerosol precursor gases from low altitudes into the UT (Hess, 2005). During the monsoon season, primarily the Asian monsoon,  
55 appreciable amounts of aerosol and gaseous aerosol precursors are lifted to the tropopause region, extending from the UT to approximately 420 K potential temperature in the stratosphere forming the Asian tropopause aerosol layer (ATAL) (Vernier et al., 2011b). The ATAL is a seasonal and regional phenomenon, which according to atmospheric modelling is strongly connected with Asian pollution sources (Neely et al., 2014). In the extratropics warm conveyor belts (WCB) (Stohl, 2001) and deep convection are the main lofting channels, the latter especially in the summer  
60 season. Large amounts of pollutants are long-range transported across the Pacific and Atlantic oceans, where sources are affecting the background concentrations of various species over remote continents. This transport is most efficient above the boundary layer in the pressure interval 700–900 hPa (Luan and Jaeglé, 2013). WCBs reaching the UT have their maximum frequency in the winter (Eckhardt et al., 2004) and transport of boundary layer air into the LMS maximizes in the winter and spring (Skerlak et al., 2014), whereas deep convection dominates in the summer (Kiley  
65 and Fuelberg, 2006).

Aerosol particles in the UT and LMS contain a large fraction of sulfur compounds, mostly sulfates (Dibb et al., 2000, Martinsson et al., 2001, Xu et al., 2001; Kojima et al., 2004). These particles also contain a considerable fraction of carbonaceous aerosol (Murphy et al., 1998; Nguyen et al., 2008; Martinsson et al., 2009), whereof a minor fraction is black carbon (Schwarz et al., 2010; Friberg et al., 2014). In some periods, particularly during spring, crustal particles  
70 also are found in this part of the atmosphere (Papapiropoulos et al., 2002).

In this study the concentration of particulate sulfur in the UT and LMS is investigated based on aerosol samples collected from the IAGOS-CARIBIC platform in the time period 1999 – 2014. In this period the UT and LMS aerosol was affected by volcanism from several eruptions as demonstrated in our previous studies based on this data set (Martinsson et al., 2009; Andersson et al., 2013; Friberg et al., 2014; Martinsson et al., 2014; Friberg et al., 2015; Andersson et al., 2015). The IAGOS-CARIBIC aerosol elemental concentration measurements from the LMS are taken in strong concentration gradients that are affected by mixing tropospheric air into the lowest part of the LMS. Each measurement flight results in a small number of samples, being insufficient to reconstruct the gradient. Therefore, we have frequently relied on concurrent IAGOS-CARIBIC measurements, mostly by relating the particulate sulfur measurements to ozone concentrations to express e.g. volcanic influence on the aerosol concentration (Martinsson et al., 2009). Satellite-based measurements do usually not provide specific chemical information about aerosol particles. This lack of chemical information in e.g. lidar measurements can cause a bias in LMS particle concentrations close to the extratropical tropopause from non-volcanic species such as crystal particles and particulate-bound water enhanced signal caused by particle hygroscopic growth. Here we present standalone estimates of the stratospheric sulfur aerosol in terms of mass concentration profiles and column concentrations based on a new method, which is then used to study the AOD of the lowest part of northern hemispheric LMS and its radiative forcing. This study also comprises a discussion on the relative importance of stratospheric and the tropospheric sources to the UT of particulate sulfur, seasonal dependences, and different modes of tropospheric transport involved.

## 2 Methods

### 2.1 Sampling, analysis and classification

This study is based on measurements of particulate sulfur taken from the IAGOS-CARIBIC platform (Brenninkmeijer et al., 2007; [www.caribic-atmospheric.com/](http://www.caribic-atmospheric.com/)), where the atmosphere is studied using modified passenger aircrafts (March 1999 – April 2002: Boeing 767-300 ER from LTU International Airways, and May 2005 – present: Airbus 340-600 from Lufthansa) during monthly sets of usually four intercontinental flights. A large number of trace gases and aerosol parameters are measured from that platform during flights in the altitude range 9-8.8 – 12 km, including gaseous and condensed water, O<sub>3</sub>, CO, NO/NO<sub>y</sub>, VOCs, greenhouse gases, halo-carbons, mercury, particle number concentrations, size distributions and elemental concentrations (Brenninkmeijer et al., 2007; Hermann et al., 2003; Schuck et al., 2009; Baker et al., 2010; Oram et al., 2012; Zahn et al., 2012; Martinsson et al., 2014; Dyroff et al., 2015; Slemr et al., 2016; Hermann et al., 2016).

Aerosol sampling from the IAGOS-CARIBIC platform in the time period 1999 – 2014 has resulted in 1198 samples analyzed for aerosol elemental concentrations. The measurements were mainly taken in the northern hemisphere (NH) extratropics and the tropics, while only a small fraction of the samples were taken in the southern hemisphere. Here the focus is on the extratropical LMS and UT of the NH. Aerosol particles with aerodynamic diameter in the range 0.08 – 2 μm were collected with a multi-channel impactor with a collection efficiency of 97% ± 4% (Nguyen et al., 2006). The typical time required to collect one sample is 100 minutes. Accelerator-based methods were used to analyze the collected samples with respect to elemental concentrations, using Particle-induced X-ray emission (PIXE) to

analyze the concentration of elements with atomic number larger than 15 (Martinsson et al., 2001). Concentrations of hydrogen, carbon, nitrogen and oxygen were investigated by particle elastic scattering analysis (PESA; Nguyen and Martinsson, 2007). Here the particulate sulfur concentrations are used. The accuracy of the analyses is estimated to 10% and the combined uncertainty in sampling and analysis is estimated to 12%. Further analytical details are found in Martinsson et al. (2014). Finally, the concentration of particulate sulfur is mostly given as a mixing ratio by normalization to STP (standard temperature (273.15 K) and pressure (101300 Pa)). When computing column concentration and AOD of the lowest 3 km of the LMS, the STP concentrations are converted to volume concentrations using the pressure and temperature of the measurement, and the altitude dependence in the LMS of the molar volume obtained from ECMWF (European centre for medium-range weather forecasts).

The dynamical tropopause (Gettelman et al, 2011) at the potential vorticity (PV) of 1.5 PVU (potential vorticity units;  $1 \text{ PVU} = 10^{-6} \text{ K m}^2 \text{ kg}^{-1} \text{ s}^{-1}$ ) was used to classify samples with respect to tropospheric and stratospheric air. The PV along the flight track was obtained from archived analyses from ECMWF with a resolution of  $1 \times 1$  degree in the horizontal and 91 vertical hybrid sigma-pressure model levels. The PV was interpolated linearly in latitude, longitude, log pressure and time to the position of the IAGOS-CARIBIC aircraft.

## 2.2 Altitude

The UT usually holds significantly lower particulate sulfur concentration than the LMS. Combined with bi-directional exchange of tropospheric and stratospheric air across the tropopause, this leads to a gradient of increasing concentration in the LMS from the tropopause. In addition, the concentration of particulate sulfur in the LMS varies due to the influence from volcanism (Martinsson et al., 2009), which has been shown to cause significant radiative forcing (Andersson et al., 2015). In order to study the particulate sulfur gradient in the LMS, fine aerosol elemental concentration measurements from the IAGOS-CARIBIC platform were used in relation to the distance between measurement position and the tropopause ( $Z$ ):

$$Z = Z_a - Z_{tp} \quad (1)$$

where  $Z_a$  and  $Z_{tp}$  are the altitudes of the aircraft and the tropopause.  $Z_{tp}$  refer to the dynamical tropopause of 1.5 PVU, which is a low limit to ensure that very little LMS air will be considered as tropospheric. The  $Z_{tp}$  was obtained from the ERA-Interim data of ECMWF, whereas the altitude of the aircraft was obtained from pressure measurement which was converted to altitude using the ECMWF data. The position of the tropopause was obtained using the aircraft as the starting position. If that position is in the UT, i.e. has a potential vorticity lower than 1.5 PVU, the tropopause is found by searching upwards in the potential vorticity field. Tropopause folds in a small number of cases induce multiple tropopauses in the vertical direction, complicating the analysis of the stratospheric samples. In order to handle that problem, searches were undertaken both upwards and downwards to find the tropopause closest to the aircraft. That distance is assigned a positive value irrespective of whether the tropopause is below or above the aircraft, because positive sign indicates stratospheric air.

The study of the aerosol concentration gradient deals primarily with the LMS. The dataset, however, contains observations both in the LMS and the UT and one sample sometimes contains particles from both regions. These concentrations are connected by the exchange across the tropopause. Figure 1 shows the samples that were taken in

the UT during the entire sampling time. It is clear that the dependence on the distance from the tropopause is non-existent ( $R^2 = 0.02$ ). Further evaluation (not shown) by normalization to seasonal average concentrations ( $R^2 = 0.01$ ) or to individual groups of concentration data that will be explained below ( $R^2 = 0.002$ ) did not reveal a dependence of the UT particulate sulfur concentration on the distance from the tropopause, either. It should be pointed out that the variability in distance to the tropopause is mainly caused by variability in the altitude of the tropopause, because the altitude of the measurement aircraft is fairly constant. This implies that the distance from the tropopause in Fig. 1 does not reflect the sampling altitude, and hence not ~~temperature decrease in relation to the surface of the earth or the an altitude-typical~~ degree of cloud processing. The distance to the tropopause for some summer measurements was very large, due to the seasonality of the position of the tropics. Approximately one third of all stratospheric air masses transported across the extratropical tropopause reach the 500 hPa level of the atmosphere, corresponding to approximately 5000 m transport, in 4 – 5 days (Skerlak et al., 2014). This illustrates that the exchange from the stratosphere goes deep in a rather short time. Based on these observations and arguments, the UT particulate sulfur concentration is considered independent of the distance from the tropopause under presented conditions, and thus indicative of the particulate sulfur concentrations of the air mixed into the LMS.

The time required to collect one sample was subdivided into ten time intervals of equal length, where  $Z$  was computed along the flight route. For samples taken in the LMS, the sample was represented by the average distance to the tropopause of the ten time intervals. Those time intervals when the sampling was undertaken in the UT, i.e. with  $Z < 0$ ,  $Z$  was set to zero, since no  $Z$  dependence of the UT particulate sulfur concentration could be identified. This implies that samples collected entirely in the UT are found at  $Z = 0$ . Some of the samples were collected both in the LMS and in the UT. In these cases, the average  $Z$  over the ten time intervals of each sample was computed with all UT parts of a sample set to  $Z = 0$ . That way all samples could be utilized to study the particulate sulfur concentration in the LMS.

### *2.3 Methodology to evaluate the S gradient around the tropopause*

The particulate sulfur concentration in the LMS is investigated by linear regression in two steps. To that end we need to consider that the LMS shows a seasonal dependence induced by variability in stratospheric circulation and exchange across the tropopause over the year. Within one season differences between years can be large, mainly due to varying influence from volcanism. Further, the onset of volcanic influence on the particulate sulfur concentration can cause large variability within the season in the same year, and patchiness of young volcanic clouds can further affect the data analysis.

This study was limited to samples taken north of  $30^\circ$  N, where in total 765 samples are available. The highest latitude of sampling was  $77^\circ$  N, and 90% of the samples (5% removed each side) were taken in the range  $32 - 64^\circ$  N. Most of the LMS samples (95%) were taken within 3000 m from the tropopause. In terms of altitude, the results of this study therefore can be considered representative of the range 0 – 3000 m above the tropopause.

A large number of measurements is needed to obtain high statistical significance. For that, data averaged over three months were used. However, to allow for exchange of data in the analyses in order to test the stability of the results, as well as catch smooth seasonal variations in the UTLS, a seasonal overlapping technique is used. In doing so three-

months-seasons shifting by one month are used, e.g. the season MAM is followed by a season AMJ where data from April and May are used in both “seasons”. Data from one season were grouped with respect to concentration levels of the different years, resulting in 4 to 5 groups of data for each season, and in total 52 groups of data from the 12 seasons to be analyzed. Some data were excluded from these analyses, which reduced the amount of data from 765 to 694. Of the excluded 71 samples, 60 pertained to periods when fresh volcanism induced strong patchiness in the particulate sulfur concentration. Eleven samples were considered outliers for other reasons, e.g. single samples affected by volcanism during a season or recent up-transport from strongly polluted regions. The remaining data of each year were tested for systematical differences. Those years where the data overlapped in particulate sulfur – height above the tropopause space, were grouped together. This way groups with varying degrees of volcanic influence were formed. Groups typical of “background” conditions were primarily based on data obtained during the 1999 – 2002 period characterized by low volcanic influence on the stratospheric aerosol (Bauman et al., 2003; Deshler et al., 2008) and data from mid-2013 to mid-2014 when the LMS is back to near-background conditions in the NH extratropics, see Table 1 for relevant volcanic eruptions. Another period that we will return to later is mid-2005 to mid-2008 when the stratosphere was affected by three tropical eruptions in 2005 – 2006: Manam, Soufriere Hills and Rabaul (Vernier et al., 2009), which also affected the NH LMS (Friberg et al., 2014). There was some variability during these years, implying that not always all of the years were grouped together. They could be grouped with other years, e.g. 2011 and 2013 often had similar concentrations and gradients during the winter and spring seasons. These groups were handled individually in the regression procedures described next, but in a context of the discussion section these groups of “moderate influence from tropical volcanism” were averaged to describe the UT aerosol along with the “background” group described above.

First linear vertical concentration gradients of particulate sulfur in the lower LMS was calculated. Ordinary linear regression (OLR) was undertaken for the 52 groups of data mentioned above of particulate sulfur concentration as a function of the altitude above the tropopause. Figure 2a shows the cumulative frequency of the coefficient of determination ( $R^2$ ) of the 52 OLRs undertaken.  $R^2$  spans 0.48 to 0.95, 72% of the groups having  $R^2$  exceeding 0.6 and 50% exceeding 0.66. The deviations from the OLR models consists of scatter that does not show any trends.

When investigating the variance of the dependent variable (the particulate sulfur concentration,  $C_s$ ) along the independent variable (altitude above the tropopause,  $Z$ ) it is clear that the variance of  $C_s$  increases with increasing  $Z$ . This heteroscedastic nature of the data, which is shared by most natural science data sets, could unfavorably affect in particular the offset of the OLR. In order to further investigate the effects of this problem two variable transformations of the dependent variable were tested: logarithmic and square root transformations. The logarithmic transformation turned the problem around to the other side, i.e. the logarithm of  $C_s$  has large variance for small  $Z$  and small variance for large  $Z$ . The square root transformation of  $C_s$ , on the other hand, shows rather constant variance along the  $Z$  axis, thus making this transformation more suitable for regression. The transformation  $C'_s = \sqrt{C_s}$  was applied to the data. As already pointed out,  $C_s$  and  $Z$  has a linear relation implying that the following expression should be minimized with respect to slope (a) and offset (b):  $(C'_s - \sqrt{aZ + b})^2$ . This results in rather tedious expressions that we solved numerically for all the 52 data groups. Figures 2b and c show comparisons between the slopes and the offsets obtained

by the square root transformed regression results and the OLRs. It is clear that the heteroscedastic nature of the data causes large deviations, especially in the offset.

215 Yet another transformation, based on forcing the regression to comply with the data for small  $Z$ , was investigated. To that end, the average concentration ( $C_{s,0}$ ) of the data points closest to the tropopause at the average distance  $Z_0$  from the tropopause was formed.  $C_{s,0}$  of the 52 data groups is on average based on 13 measurements, and the minimum was 5, and the average  $Z_0$  is 88 m and the largest is 273 m. The data were transformed linearly to place  $C_{s,0}$  and  $Z_0$  in the origin by forming  $C_s' = C_s - C_{s,0}$  and  $Z' = Z - Z_0$  followed by linear regression forced through the origin, i.e.  $C_s' = aZ'$ . Finally, the regression results are transformed back to the form  $C_s = aZ + b$ , where the slope  $a$  is not changed by the translation, and the offset is obtained by  $b = C_{s,0} - aZ_0$ . These results of the forced linear regressions are compared with the square root transformed regressions in Figures 2b and c. As can be seen, the forced linear regressions, in contrast to the OLRs, show only small deviations from the square root transformed data. Due to the more direct determination of the offset as well as the simplicity of forced linear regression compared with square root transformation, the forced linear regression method is chosen for the analyses. Thus, for each season ( $s$ ) and year ( $y$ ) in total 52 forced linear regressions were undertaken:

$$C_s(y, s, Z) = a(y, s)Z + b(y, s) \quad (2)$$

where  $a$  and  $b$  varies with season and the strength of the volcanic influence.

230 The  $S$  concentration at the tropopause and in the UT, expressed by  $b$  in eq. 2 is dependent on the stratospheric concentration which is affected by volcanism (Friberg et al., 2015). This means that the offset of the regressions is affected. This is expressed here by:

$$b(y, s) = C_{s,UT}(y, s) = a(y, s)k(s) + C_{s,UTtrop}(s) \quad (3)$$

where the first term expresses the contribution from stratospheric sources and the second that of tropospheric sources.

This results in the combined equation:

$$235 \quad C_s(y, s, Z) = a(y, s)Z + a(y, s)k(s) + C_{s,UTtrop}(s) \quad (4)$$

where  $k(s)$  reflects stratospheric influence on the UT particulate sulfur concentration and  $C_{s,UTtrop}(s)$  is the particulate sulfur concentration of tropospheric origin.

In order to obtain estimates of  $k$  and  $C_{s,UTtrop}$  a second regression for each season is undertaken where the offsets ( $b$ ) are related to the slopes ( $a$ ), see Fig. 3. The relative variance of the slopes was much smaller than that of the offsets (average variance ratio of 0.19). For simplicity, the variance of the slope was therefore neglected in these regressions.

240 The groups of data of a season differ in  $a$  and  $b$  mainly due to volcanic influence. A zero slope ( $a = 0$ ) would be obtained, should the stratospheric concentration become as low as the UT concentration. The UT aerosol of tropospheric origin ( $C_{s,UTtrop}$ ) thus can be estimated as the offset of the  $b - a$  regression. The slope of that regression shows how the offset  $b$  changes with increased slope  $a$ , hence expressing the sensitivity ( $k$ ) of the UT concentration to changes in stratospheric concentration. With access to these two parameters,  $C_{s,UTtrop}(s)$  and  $k(s)$ , the LMS concentration gradient and tropopause concentration can be estimated based on a single measurement of the particulate sulfur concentration.

The uncertainties of the forced regression results of a given season vary among the data groups. In order to account for this variability, weights are used in the regression between the offsets and slopes of a season. The weights are based on 70% double-sided student t distribution (t70%) estimates because some of the estimated  $C_{s,0}$  and  $Z_0$  rely on few observations. The t70% estimate of b ( $b = C_{s,0} - aZ_0$ , see above) is obtained by combining the upper t70% limit of  $C_{s,0}$  with the weakest t70% slope (a) and the strongest t70% slope at the lower limit multiplied with  $Z_0$ . The inverse of the squared t70% estimates of b obtained in this way are then used as weights in the second step regressions.

#### 2.4 Lidar data from the CALIPSO satellite

The evaluation of the particulate sulfur concentrations from the IAGOS-CARIBIC aircraft was aided by the use of lidar data from the CALIOP sensor aboard the CALIPSO (Cloud-aerosol lidar and infrared pathfinder satellite observation) satellite from NASA (Winker et al., 2010), performing 15 orbits per day covering the globe from 82° S to 82° N with a repeat cycle of 16 days during nights and days. The data evaluation was based on the methodology developed by Vernier et al., (2009). Here only the night-time data of the 532 nm wavelength lidar signals were used. The level 1 data of version 4-10 (averaged in a grid of 180 m vertically and 1x1 horizontally) were used to obtain scattering ratios, i.e. the ratio of the measured, combined air and aerosol scattering to the modeled air scattering based on molecule and ozone number concentrations from the GMAO (global modelling and assimilation office). Cloud pixels were removed by rejecting pixels of a depolarization ratio greater than 5%. This cloud mask was extended 360 m upwards to remove faint cloud residues, and pixels from underneath clouds were removed to avoid bias from cloud absorption, see Andersson et al., (2015) for further details. Removal of cloud pixels as well as periods of instrument failure, sometimes resulted in few observations in these pixels. In order to avoid statistical noise at least 20% of the maximum possible observation data was required for any given pixel (latitude, altitude). The pixels were generated by averaging in the longitude interval 60° E to 120° E, the main longitude region of the ATAL (Vernier et al., 2015).

### 3. Results

The linear regression methodology described in the previous section was applied to particulate sulfur concentrations ( $C_s$ ) as a function of distance from the tropopause ( $Z$ ) for seasons comprising three months. Data from different years of one season were grouped according to their concentrations, resulting in four or five groups differing with respect to the  $C_s - Z$  relation. The resulting groups of data from each season were modeled by forced linear regression. These model results are used in a second regression step to model seasonal influences from transport, which in turn are used to obtain the response of the LMS and UT particulate sulfur concentrations to changes induced mainly by volcanic eruptions.

This methodology was applied to all seasons, having a duration of three months. The final product of the regression methodology, i.e. the second regression step, of all twelve overlapping seasons is shown in Fig. 3. It is clear that slopes and offsets of the season groups obtained in the first step of forced linear regression, differing in particulate sulfur concentration related to distance from the tropopause, can readily be described by linear regressions for all twelve

three-month seasons. The relation between slopes and offsets has a very strong seasonal dependence. In some seasons small changes in the LMS sulfur concentration slope is connected with a strong change in tropopause sulfur concentration. This is most pronounced for the seasons centered in February, March and April. At the other end we find the seasons centered in September, October and November, where a change in the slope of the first regression step due to varying influence from volcanism has little effect on the tropopause concentration. To further illustrate the connection to the measurements, Fig. 4 shows the four data groups of the seasons most (Fig. 3: FMA) and least (Fig. 3: OND) susceptible to change in the tropopause concentration due to changed stratospheric concentration. In Fig. 4a the data group of least volcanic influence includes years 1999 – 2002 and 2014, and the most influenced years are 2009 and 2012 which are the springs after the Kasatochi (August 7, 2008) and Nabro (June 12, 2011) eruptions (we have no late winter/spring data after the most powerful eruption of the period studied (Sarychev, June 12, 2009) due to maintenance of the measurement aircraft). The season centered in November have the weakest volcanic influence years 1999 – 2001, 2012 and 2013 (Fig. 4b), whereas the group with strongest slope includes the three strongest eruptions of the period studied (Kasatochi, Sarychev and Nabro) a few months after respective eruption. Despite the very strong volcanic influence of the latter group, the tropopause concentration ( $Z = 0$ ) remains close to that of the closely distributed tropopause concentrations of the other groups of that season.

The sensitivity of the tropopause and UT concentration to changes in LMS concentration slope is denoted  $k$  in eq. 3 and 4, which thus is obtained for each regression depicted in Fig. 3. These results are collected in Fig. 5a. The salient features of the seasonal dependence can be described by two Gaussian distributions (Fig. 5a). The maximum sensitivity appears in the late winter and early spring when the seasonal variation in tropopause altitude has its maximum in rate of upward motion (Appenzeller et al., 1996). The maximum in rate of downward motion of the tropopause appears in the fall. This delays transport from the stratosphere to the troposphere, which is reflected by a low sensitivity  $k$ . This sensitivity can, in addition, be affected by the residence time of particulate sulfur in the UT. Interestingly, the downward transport in association with the Brewer-Dobson circulation from deeper, aerosol-rich stratospheric layers through the LMS takes place in the same season as the maximum in  $k$ , thus further enhancing the stratospheric influence on the UT by the term  $a(y,s) \times k(s)$  in eq. 4.

The offsets of the regression lines shown in Fig. 3 (when  $a = 0$ ) expresses the case when the stratospheric and tropospheric concentrations are equal, implying that this offset expresses the particulate sulfur concentration of tropospheric origin ( $C_{S,UTtrop}$ ) that is mixed into the LMS, see Fig. 5b. Two Gaussian distributions were used as fits to the seasonal variation of  $C_{S,UTtrop}$ . As for  $k$ , the seasonal variation of  $C_{S,UTtrop}$  is strong. The seasonal variation of  $C_{S,UTtrop}$  will be elaborated in the discussion section.

After obtaining  $C_{S,UTtrop}$  and  $k$ , the data that are needed for conversion of every measurement of the concentration to an estimate of the slope and offset of the LMS concentration are available. Thus, for each individual measurement ( $i$ ) in the LMS, consisting of the particulate sulfur concentration ( $C_{S,i}$ ) and the altitude  $Z_i$  above the tropopause, the slope and offset of equations 3 and 4 are obtained from:

$$a_i = \frac{C_{S,i} - C_{S,UTtrop}(s)}{Z_i + k(s)} \quad (5)$$

$$b_i = a_i k(s) + C_{S,UTtrop}(s) \quad (6)$$

The estimated slopes and offsets are shown in Figs. 6a and b, where the dots are individual measurements and the histogram monthly averages. Here measurements taken at an altitude of less than 50 m above the tropopause are not shown, because they were judged to have too small stratospheric character for an estimate of the concentration slope in the LMS. Both the slope and the tropopause concentration are affected by volcanism (Table 1), but the relative response of the slope is much stronger than that of the tropopause concentration, see e.g. the falls of 2008 and 2009 affected by the Kasatochi and Sarychev eruptions.

Observations at various altitudes above the tropopause are difficult to compare, due to the concentration gradient in the LMS. With the estimates of the tropopause concentration and the slope in the particulate sulfur LMS concentration, each measurement becomes an estimate of the total amount of particulate sulfur in the altitude interval investigated by integration of eq. 4. However, first the STP concentrations ( $C_{s,STP} = C_s = aZ + b$ ) need to be converted to volume concentrations ( $C_{s,v}$ ), which are related by  $C_{s,v} = C_{s,STP} Q_{STP} / Q_v(Z)$ , where  $Q$  are molar volumes. For that purpose, the altitude dependence of the molar volume from the tropopause up to 5 km above the tropopause was extracted from temperatures and pressures obtained from ECMWF for each sample. The molar volume can be expressed as  $Q_v(Z) = Q_v(0)e^{wZ}$ , where the tropopause is at  $Z = 0$  and  $w = 0.0001535 \text{ m}^{-1}$  obtained as the average of all samples. For a measurement the molar volume is  $Q_m$  obtained at distance  $Z_m$  from the tropopause, and the tropopause molar volume is computed by  $Q_v(0) = Q_m e^{-wZ_m}$ . Finally, the column concentration of particulate sulfur for the first 3 km above the dynamical tropopause of 1.5 PVU is obtained by integration of the volume concentration:

$$C_{s,col} = \int_0^{3000} \frac{Q_{STP}}{Q_v(Z)} C_{s,STP}(Z) dZ = \int_0^{3000} \frac{Q_{STP}}{Q_v(0)} e^{-wZ} (aZ + b) dZ \quad (7)$$

The particulate sulfur column is calculated for altitudes above the tropopause ( $Z$ ) in the range 0 to 3000 m, the upper limit because too few measurements (5%) were taken above that level. After integrating the column, it is also interesting to estimate the total amount of sulfur-connected aerosol. To that end, it was assumed that the aerosol consists of 75% sulfuric acid and 25% water, which is a commonly used stratospheric composition (Rosen, 1971; Arnold et al., 1998). This means that the total sulfur column was multiplied by a stoichiometric factor of  $h = 4.084$  to obtain the column of the sulfuric acid – water aerosol:

$$C_{A,col} = h C_{s,col} \quad (8)$$

The measurements were taken in the northern hemispheric latitudes higher than  $30^\circ$  with the highest latitude of  $77^\circ$ , with 90% of the data in the latitude range  $32 - 64^\circ \text{ N}$  and 70% of the data between  $37$  and  $57^\circ \text{ N}$ . The northern mid-latitudes sulfur aerosol columns are shown in Fig. 6c as monthly averages with standard errors (the few months with only one measurement available are shown without an error bar). The sulfur aerosol column of the LMS shows large variability primarily caused by volcanism. The lowest columns are found in the period 1999 – 2002, when the volcanic influence on the stratospheric aerosol was small, see Table 1 for relevant volcanic eruptions. The time period mid-2005 to mid-2008 was affected by tropical volcanism (Vernier et al., 2011a), which also caused elevated concentrations in the NH LMS (Friberg et al., 2014). The eruptions of the extratropical volcano Kasatochi in August 2008 placed two volcanic clouds in the stratosphere (Andersson et al., 2015), one in the LMS causing strongly elevated aerosol column of the LMS that ceased by November the same year, and the other above the LMS. The latter cloud was transported

355 downward, causing a rise of the lower LMS aerosol column in December 2008. After some influence from several  
eruptions of the extratropical volcano Redoubt in the spring of 2009, the eruption of Sarychev strongly affected the  
northern hemispheric stratosphere from June of 2009. The eruption of the Icelandic volcano Grimsvötn in May 2011  
had a strong and short impact on the northern LMS, which is reflected by a peak in June to July 2011 (Fig. 6c), before  
the tropical volcano Nabro reached the northern LMS in the early fall of the same year. After that eruption a gradual  
decrease of the aerosol load can be seen. The concentrations after mid-2013 approaches those of the period 1999 –  
360 2002, which was close to stratospheric background conditions.

The LMS aerosol contains a significant fraction of carbonaceous material (Martinsson et al., 2009), mainly organic in  
nature (Friberg et al., 2014), which will add to the aerosol columns of Fig. 6c and affect the refractive index of the  
particles. However, this work is dealing with the sulfurous fraction, the main fraction of the stratospheric aerosol. To  
put the results presented in Fig. 6c into perspective, the AOD is estimated using a simplified aerosol (which thus likely  
365 is an underestimation of the AOD). Thus, the “standard” stratospheric 75% sulfuric acid – 25% water composition, the  
particle density 1.669 g/cm<sup>3</sup> and the refractive index 1.44 will be used. Furthermore, particle size distribution  
measurements from IAGOS-CARIBIC have been taken since 2010 (Hermann et al., 2016). The changes of the size  
distribution induced by the moderate 2011 eruptions of Grimsvötn and Nabro were small (Martinsson et al., 2014),  
and agree well with previous measurements (Andersson et al., 2015) of the stratospheric background particle size  
370 distribution by Jäger and Deshler (2002). Thus, for the estimation of the AOD the latter particle size distribution was  
used for the entire time period studied. For fixed composition and particle size distribution the AOD is obtained as a  
fixed relation to aerosol column:  $AOD = fC_{A,col}$ , where  $f$  contains the relations between mass and area/extinction, in  
this case  $f = 3.29 \cdot 10^{-6} m^2/\mu g$ . Finally, converting the AOD to radiative forcing (RF) using the global average relation  
(Hansen et al., 2005; Solomon et al., 2011) of:

$$375 \quad RF = -25 \times AOD; \quad \text{in } Wm^{-2} \quad (9)$$

to obtain an estimate of the climate influence of the sulfate aerosol of the lower LMS. The peak AOD of the Kasatochi  
and Sarychev eruptions is approximately 0.006, corresponding to -0.15 W/m<sup>2</sup> in regional radiative forcing of the lowest  
3000 meters of the northern hemisphere LMS. Although no detailed comparisons will be made here, we find that the  
AOD and radiative forcing obtained from the particulate sulfur measurements show similar tendencies as satellite-  
380 based measurements (Andersson et al., 2015). The findings presented here also corroborates the findings of Andersson  
et al. (2015) on the importance of the LMS for the total stratospheric AOD and radiative impact.

#### 4 Discussion

The results presented here are based on measurements in the extratropical UT and the LMS of the NH, where the latter  
includes the extratropical transition layer (ExTL). Bi-directional exchange across the tropopause affects strong  
385 gradients in the ExTL for species having clearly different stratospheric and tropospheric concentrations (Hoor et al.,  
2002), such as particulate sulfur (Martinsson et al., 2005). In the previous section, the UT concentration, the gradient  
in the LMS and column amount of particulate sulfur in the ExTL were investigated, with its seasonal dependence (Fig.  
5) and the influence from volcanism (Fig. 6). In the processing of the data to obtain these results, in particular one

feature stands out: the seasonal dependence of the particulate sulfur concentration from tropospheric sources that is mixed into the ExTL (Fig. 5b). It shows a broad maximum from August to December, peaking in September to November, and a deep minimum in the late winter and early spring (February and March).

Let us now compare these concentrations in the UT of tropospheric origin ( $C_{S,UTrop}$ ) with average concentrations of particulate sulfur in the UT for two cases: “background conditions” dominated by data from mid-1999 to mid-2002 and “moderate influence from tropical volcanism” dominated by data from mid-2005 to mid-2008, see section 2.3 for details. The seasonal dependence of these two categories are shown in Fig. 7a together with  $C_{S,UTrop}$ . It is clear that, in line with the findings of Friberg et al. (2015), the UT particulate sulfur concentration is affected by volcanism. In Fig. 7a we see that the main differences in UT concentrations of the two cases appears from January to July, coinciding with the season of transport down from the Junge layer into the LMS and the shrinkage of the LMS due to tropopause upward motion (Appenzeller et al., 1996; Gettelman et al., 2011). In the period September to November the influence from volcanism on the UT particulate sulfur concentrations is small, compare the background and moderate volcanism cases (Fig. 7a). Comparing these cases to the concentration of particulate sulfur found to be of tropospheric origin it is clear that there is a strong agreement between all three categories in the fall months, whereas differences are large during the remainder of the year. The UT particulate sulfur concentration of stratospheric origin can be estimated by subtracting  $C_{S,UTrop}$  from the two cases of UT concentrations. The results are shown in Fig. 7b, where peak stratospheric influences of 19 and 36 ng/m<sup>3</sup> STP are found in the spring, and minimum contributions of approximately 1 ng/m<sup>3</sup> STP in the fall for the “background” and “moderate volcanism” cases. Summing the observations up (Fig. 7c), a clear seasonal dependence in the fraction of the UT particulate sulfur concentration originating in the stratosphere was found, from close to 100% in late winter/spring to approximately 10% in the fall. On a yearly average the fraction of the UT particulate sulfur that originates in the stratosphere is approximately 50% during background conditions and 70% during moderate influence from tropical volcanism.

The tropospheric source of UT particulate sulfur could be transported from the planetary boundary layer either in the form of particulate sulfur or precursor gases, in the latter case primarily sulfur dioxide (SO<sub>2</sub>). In fall, winter and spring, with maximum in the winter, vertical transport by WCBs from the boundary layer to the UT is strong (Eckhardt et al., 2004), whereas in the summer deep convection is the most important mode (Hess, 2005; Kiley and Fuelberg, 2006) in the extratropics. For sulfate, the most common chemical form of particulate sulfur, the concentration usually shows a rapid decline with altitude in the troposphere (Heald et al., 2011) associated with formation of precipitation. Cloud processing also tends to strongly reduce SO<sub>2</sub> concentrations with altitude, where the relative availability of SO<sub>2</sub> and hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) is important for the SO<sub>2</sub> lifetime in the cloud.

We have found some very distinctive characteristics of the particulate sulfur concentration of tropospheric origin in the UT, with low concentrations in February and March, increasing concentrations during the summer and maximum in the sources of tropospheric origin in the fall. Carbon monoxide (CO) is often used as a tracer of air pollution. Zbinden et al. (2013) found winter/spring maximum in the CO concentration in the UT of the NH in contrast to the tropospheric component of the UT particulate sulfur. This contrast can, at least in part, be explained by the oxidizing capacity in the UT. The summer abundance of the hydroxyl (OH) radical (Bahm and Khalil, 2004) induces a decline in CO

425 (Bergamaschi et al., 2000; Osman et al., 2016). High abundance of this radical, on the other hand, and availability of  
SO<sub>2</sub> can lead to production of particulate sulfur. SO<sub>2</sub> measurements by the satellite-based instrument MIPAS have  
recently become available (Höpfner et al., 2015). These data have large uncertainties (Höpfner et al., 2015), and likely  
overestimate the SO<sub>2</sub> concentration (Rollins et al., 2017). However, here we only qualitatively use the seasonal  
430 variation. The MIPAS results indicate a UT seasonal variation in the NH midlatitudes with low concentrations in  
December to March and the highest concentrations in June to September (Höpfner et al., 2015). Deep convection  
provides a rapid route upwards in the atmosphere, favoring transport of short-lived species like SO<sub>2</sub> (TF-HTAP-2010;  
Dickerson et al., 2007). Low abundance of both SO<sub>2</sub> and OH in the NH UT thus could explain the weak tropospheric  
contribution to the UT particulate sulfur during late winter and early spring. The increase in  $C_{S,UTrop}$  during the spring  
and summer months coincides with increases in convective activity as well as in the concentration of both SO<sub>2</sub> and  
435 OH, thus offering a plausible explanation. However, the quantitative understanding of the SO<sub>2</sub> transport paths and UT  
seasonality requires further study. Still, we need to consider the late maximum in the fall of the tropospheric source of  
UT particulate sulfur.

The Asian monsoon is an important feature of the global circulation in June to September, which has been found to  
reach deep into the lower stratosphere. This intrusion of tropospheric air into the tropical transition layer (Sunilkumar  
440 et al., 2017) has been manifested by gas phase components including water and ozone (Gettelman et al., 2004; Randel  
and Park 2006) and hydrogen cyanide (Randel et al., 2010). Later an aerosol layer extending from the UT to the  
potential temperature of 420 K in the lower stratosphere was found at 14 to 18 km altitude (Vernier et al., 2011b;  
Thomason and Vernier, 2013; Vernier et al., 2015). Figs. 8a-e show monthly means of the scattering ratio obtained  
from the CALIPSO sensor CALIOP for the months June to October 2013, when the volcanic influence was low.  
445 Formation of the ATAL can be identified in July (Fig. 8b) with maximum intensity in August (Fig. 8c). In September  
still a weakened ATAL can be identified (Fig. 8d), whereas in June and October the scattering in the ATAL region at  
14 – 18 km altitude is very weak (Figs. 8a and e).

The altitude range of the ATAL is above the measurement altitudes of IAGOS-CARIBIC (9 – 12 km). However, the  
stratospheric poleward circulation along isentropes bending downwards, further amplified by an extratropical cross-  
450 isentrope, downward component, brings the ATAL down to lower altitudes, is directed downwards in the extratropics,  
but the ATAL is too weak to be traced by CALIOP in the downward transport (Figs. 8 a-e). We therefore illustrate  
the subsidence using the eruption of the tropical volcano Nabro in the summer 2011 (Table 1), with effluents that  
occupied approximately the same region as the ATAL, see Fig. 3 in Bourassa et al. (2012). The effluents of this  
eruption were rapidly transported to the north in the TTL. Figs 8f-j actually includes two volcanic eruptions, besides  
455 the tropical volcano Nabro, the Icelandic volcano Grimsvötn injected a volcanic cloud to the tropopause region at  
midlatitudes which is visible in June and July 2011. The time series of the Nabro volcanic cloud in Figs. 8f-j  
unambiguously demonstrates the transport down to the IAGOS-CARIBIC flight altitudes in the season of the ATAL.  
In Fig. 6c we see a clear increase in the particulate sulfur column in September 2011, following the decline of the June-  
July 2011 short peak from the of Grimsvötn eruption in May 2011, thus demonstrating agreement between CALIOP  
460 and IAGOS-CARIBIC measurements of particulate sulfur. The volcanic cloud from Nabro initially resided at  
somewhat higher altitude than the ATAL, implying that the ATAL can be expected to reach IAGOS-CARIBIC flight

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altitudes somewhat earlier in the year. The transport in this season contains little particulate sulfur during periods without fresh volcanic aerosol, compare in Fig. 7 “tropical volcanism” (this category of Fig. 7 does not include the Nabro eruption, see section 2.3) with “background” and the particulate sulfur of tropospheric origin.

465 The downward transport allows delayed detection of the ATAL in the fall from altitudes above the measurement altitude range of IAGOS-CARIBIC in the same way as the spring detection of aerosol transported from the Junge layer, with or without volcanic influence, as shown in Fig. 7. A difference between these two seasons is that in the spring the strong source of particulate sulfur is stratospheric, whereas in the fall the source is tropospheric extending from the UT into the stratosphere. We therefore conclude that the UT particulate sulfur concentration of tropospheric origin that starts to increase rapidly in August, peaking in September to November, and is back at low concentration in January (Fig. 5b) most likely is caused by the ATAL formation from the Asian monsoon.

470 Model studies find similar UT – stratosphere distribution of the ATAL as the experimental studies, as well as that Asian pollution sources strongly contribute to the ATAL, and that sulfate is an important component of that aerosol layer (Neely et al., 2014). Besides this component, primary and secondary organic aerosol constituents are predicted to be important components of the ATAL (Yu et al., 2015). The present study is, as far as we know, the first observation of a chemical component of the ATAL. In agreement with modeling results we find that particulate sulfur is a component of the Asian tropopause aerosol layer. This component affects the tropopause region from July to September at 14 to 18 km altitude (Fig. 8) and the extratropical tropopause region from August to December (Fig. 5b).

## 5 Conclusions

480 Particulate sulfur (usually sulfate) in the upper troposphere (UT) and the lowermost stratosphere (LMS) obtained from the IAGOS-CARIBIC platform was investigated at northern midlatitudes in the time period 1999 – 2014, which covers several tropical and extratropical volcanic eruptions. The study is based on the use of linear regression models, where individual measurements in the strong gradient of the extratropical transition layer (ExTL) can be converted to an estimate of the column of particulate sulfur in a 3000 m layer above the dynamical tropopause (here defined at 1.5 potential vorticity units). The obtained time series in particulate sulfur column concentration shows distinct response to extratropical volcanism and delayed elevation of the column concentration following tropical eruptions. Assuming the stratospheric background particle size distribution and composition (75% sulfuric acid and 25% water) the AOD and radiative forcing were estimated, e.g. the peak value following the 2009 Sarychev eruptions were estimated to 0.006 and -0.15 W/m<sup>2</sup>, respectively. These estimates refer mainly to the ExTL, i.e. lowest part of the LMS, thus highlighting the importance of the lowest part of the stratosphere for the overall climate impact of volcanism.

495 As part of this investigation the sources of UT particulate sulfur were explored. A distinct pattern emerges where tropospheric sulfur sources dominate the supply of particulate sulfur to the UT in the fall, whereas stratospheric sources dominate in January to July, the main season of transport from the Junge layer into the LMS. The smallest contributions from the troposphere are found in February and March in conjunction with low sulfur dioxide (SO<sub>2</sub>) and oxidant concentrations. As the concentrations of these species increase, the UT particulate sulfur concentration of tropospheric origin increases somewhat in April to July. The particulate sulfur concentration shows a threefold increase during the

fall, with maximum concentration in September to November. Making use of lidar data from the CALIPSO satellite together with the in situ measurements we find that the Asian tropopause aerosol layer (ATAL) resulting from the Asian monsoon is the cause of the increase. The ATAL is formed at 14 – 18 km altitude and extends from the UT to approximately 420 K potential temperature in the stratosphere, with main extension in July to September. The ATAL is transported downwards, and affects the extratropical tropopause region in August to December. As far as we know, this is the first measurement of a chemical species in particles connected with the ATAL. The stratospheric and tropospheric contributions to the UT particulate sulfur concentrations thus have strong and opposite seasonal dependences. On annual average the stratospheric contribution to the UT particulate sulfur is estimated to 50% during stratospheric background conditions. During influence from moderate tropical volcanism the stratospheric fraction rises to 70%. The particulate sulfur concentration in the UT is thus to a large degree governed by the stratosphere and volcanism.

#### **Data availability**

Data are available upon request to the corresponding author.

#### **510 Competing interests**

The authors declare that they have no conflict of interest.

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#### **References**

- 520 Andersson, S. M., B. G. Martinsson, J. P. Vernier, J. Friberg, C. A. M. Brenninkmeijer, M. Hermann, P. F. J. van Velthoven, and A. Zahn, Significant radiative impact of volcanic aerosol in the lowermost stratosphere, *Nat. Commun.* 6:7692 doi:10.1038/ncomms8692, 2015.
- Andersson S. M., Martinsson B. G., Friberg J., Brenninkmeijer C. A. M., Rauthe-Schöch A., Hermann M., van Velthoven P. J. F. and Zahn A., Composition and evolution of volcanic aerosol from eruptions of Kasatochi, Sarychev and Eyjafjallajökull in 2008 – 2010 based on CARIBIC observations. *Atmos. Chem. Phys.* 13, 1781-1796, doi:10.5194/acp-13-1781-2013, 2013.
- 525

- Appenzeller C., J.R. Holton, and K.H. Rosenlof, Seasonal variation of mass transport across the tropopause, *J. Geophys. Res.*, 101, 15071-15078, 1996.
- 530 Arnold F., J. Curtis, S. Spreng and T. Deshler: Stratospheric aerosol sulfuric acid: First direct in situ measurement using a novel balloon-based mass spectrometer apparatus, *J. Atmos. Chem.*, 30, 3-10, 1998.
- Bahm K. and M.A.K. Khalil, A new model of tropospheric hydroxyl radical concentrations, *Chemosphere*, 54, 143-166, 2004.
- Baker, A. K., Slemr F. and Brenninkmeijer C. A. M., Analysis of non-methane hydrocarbons in air samples collected aboard the CARIBIC passenger aircraft, *Atmos. Meas. Tech.*, 3(1), 311-321, 2010.
- 535 Bauman J.J., P.B. Russel, M.A. Geller and P. Hamill: A stratospheric aerosol climatology from SAGE II and CLAES measurements: 2. Results and comparisons, 1984-1999, *J. Geophys. Res.*, 108(D13), 4383, doi:10.1029/2002JD002993, 2003.
- Bergamaschi P., R. Heim, M. Heimann and P.J. Crutzen, Inverse modeling of the global CO cycle 1. Inversion of CO mixing ratios, *J. Geophys. Res.*, 105, 1909-1927, 2000.
- 540 Bourassa A. E., A. Robock, W.J. Randel, T. Deshler, L.A. Rieger, N.D. Lloyd, E.J. Llewellyn and D.A. Degenstein, Large volcanic aerosol load in the stratosphere linked to Asian monsoon transport. *Science* 337, 78-81, 2012.
- Brenninkmeijer C. A. M., Crutzen P., Boumard F., Dauer T., Dix B., Ebinghaus R., Filippi D., Fischer H., Franke H., Frieß U., Heintzenberg J., Helleis F., Hermann M., Kock H. H., Koepfel C., Lelieveld J., Leuenberger M., Martinsson B. G., Miemczyk S., Moret H. P., Nguyen H. N., Nyfeler P., Oram D., O'Sullivan D., Penkett S., Platt U., Pukek M., Ramonet M., Randa B., Reichelt M., Rhee T. S., Rohwer J., Rosenfeld K., Scharffe D., Schlager H., Schumann U., Slemr F., Sprung D., Stock P., Thaler R., Valentino F., van Velthoven P., Waibel A., Wandel A., Waschitschek K., Wiedensohler A., Xueref-Remy I., Zahn A., Zech U. and Ziereis H., Civil aircraft for the regular investigation of the atmosphere based on an instrumented container: the new CARIBIC system. *Atmos. Chem. Phys.* 7, 4953-4976, 2007.
- 545 Brühl C., J. Lelieveld, H. Tost, M. Höpfner and N. Glatthor., Stratospheric sulfur and its implications for radiative forcing simulated by the chemistry climate model EMAC, *J. Geophys. Res.* 120, 2103-2118, doi:10.1002/2014JD022430, 2015.
- Carn, S. A., and F. J. Prata, Satellite-based constraints on explosive SO<sub>2</sub> release from Soufrière Hills Volcano, Montserrat, *Geophys. Res. Lett.*, 37, doi: 10.1029/2010GL044971, 2010.
- 555 Clarisse, L., D. Hurtmans, C. Clerbaux, J. Hadji-Lazaro, Y. Ngadi, and P.-F. Coheur, Retrieval of sulphur dioxide from the infrared atmospheric sounding interferometer (IASI), *Atmos. Meas. Tech.*, 5(3), doi: 10.5194/amt-5-581-2012, 2012.
- Crutzen P.J., The possible importance of CSO for the sulfate layer of the stratosphere. *Geophys. Res. Lett.* 3, 73-76, 1976.
- 560 Deshler T., A review of global stratospheric aerosol: Measurements, importance, life cycle, and local stratospheric aerosol, *Atmos. Res.*, 90, 223-232, 2008.
- Dibb J. E., Talbot R. W. and Scheuer E. M., Composition and distribution of aerosols over the North Atlantic during the subsonic assessment ozone and nitrogen oxide experiment (SONEX), *J. Geophys. Res.*, 105, 3709– 3717, doi:10.1029/1999JD900424, 2000.

- 565 Dickerson R. R., C. Li, Z. Li, L.T. Marufu, J.W. Stehr, B. McClure, N. Krotkov, H. Chen, P. Wang, X. Xia, X. Ban, F. Gong, J. Yuan and J. Yang, Aircraft observations of dust and pollutants over northeast China: Insight into the meteorological mechanisms of transport, *Journal of Geophys. Res.*, 112(D24S90), doi:10.1029/2007JD008999, 2007.
- Dyroff C., A. Zahn, E. Christner, R. Forbes, A.M. Tompkins and P.F.J. van Velthoven, Comparison of ECMWF analysis and forecast humidity data with CARIBIC upper troposphere and lower stratosphere observations, Q. J. R. Meteorol. Soc. 141:833-844, doi:10.1002/qj.2400, 2015.
- 570 Eckhardt S., A. Stohl, H. Wernli, P. James, C. Forster and N. Spichtinger, A 15-year climatology of Warm Conveyor Belts, *J. Climate*, 17, 218-237, 2004.
- England M. H., S. McGregor, P. Spence, G.A. Meehl, A. Timmermann, W. Cai, A.S. Gupta, M.J. McPhaden, A. Purich and A. Santoso, Recent intensification of wind-driven circulation in the Pacific and the ongoing warming hiatus. *Nat. Clim. Change* 4, 222-227, 2014.
- 575 Friberg, J., B. G. Martinsson, S. M. Andersson, C. A. M. Brenninkmeijer, M. Hermann, P. F. J. van Velthoven, and A. Zahn, Sources of increase in lowermost stratospheric sulphurous and carbonaceous aerosol background concentrations during 1999-2008 derived from CARIBIC flights, *Tellus B*, 66, 23428, <http://dx.doi.org/10.3402/tellusb.v66.23428>, 2014.
- 580 Friberg J., B.G. Martinsson, M.K. Sporre, S.M. Andersson, C.A.M. Brenninkmeijer, M. Hermann, P.F.J. van Velthoven and A. Zahn, Influence of volcanic eruptions on midlatitude upper tropospheric aerosol and consequences for cirrus clouds, *Earth and Space Science*, 2, doi:10.1002/2015EA000110, 2015.
- Fyfe J. C., Gillett N. P. & Zwiers F. W. Overestimated global warming over the past 20 years. *Nat. Clim. Change* 3, 767-769, 2013.
- 585 Fyfe J.C., Meehl G.A., England M.H., Mann M.E., Santer B.D., Flato G.M., Hawkins E., Gillett N. P., Xie S.-P., Kosaka Y. and Swart N.C., Making sense of the early-2000s warming slowdown. *Nat. Clim. Change* 6, 224-228, 2016.
- Gettelman A., Kinnison D.E., Dunkerton T.J. and Brasseur G.P., Impact of monsoon circulations on the upper troposphere and lower stratosphere. *J. Geophys. Res.*, 109, D22101, doi:10.1029/2004JD004878, 2004.
- 590 Gettelman, A., P. Hoor, L. L. Pan, W. J. Randel, M. I. Hegglin, and T. Birner. The extratropical upper troposphere and lower stratosphere. *Rev. Geophys.* 49, doi: 10.1029/2011RG000355, 2011.
- Hansen J, et al., Efficacy of climate forcings, *J. Geophys. Res.*, 110, D18104, doi:10.1029/2005JD005776, 2005.
- Haywood, J. M., A. Jones, L. Clarisse, A. Bourassa, J. Barnes, P. Telford, N. Bellouin, O. Boucher, P. Agnew, and C. Clerbaux, Observations of the eruption of the Sarychev volcano and simulations using the HadGEM2 climate model, *J. Geophys. Res.*, 115(D21), doi: 10.1029/2010JD014447, 2010.
- 595 Heald C.L. et al., Exploring the vertical profile of atmospheric organic aerosol: comparing 17 aircraft field campaigns with a global model, *Atmos. Chem. Phys.*, 11, 12673-12696, doi:10.5194/acp-11-12673-2011, 2011.
- Hermann M., Heintzenberg J., Wiedensohler A., Zahn A., Heinrich G. and Brenninkmeijer C. A. M., Meridional distributions of aerosol particle number concentrations in the upper troposphere and lower stratosphere obtained by Civil Aircraft for Regular Investigation of the Atmosphere Based on an Instrument Container (CARIBIC) flights, *J. Geophys. Res.*, 108(D3), 4114, doi:10.1029/2001JD001077, 2003.
- 600

Hermann M., A. Weigelt, D. Assmann, S., Pfeifer, T. Müller, T. Conrath, J. Voigtländer, J. Heintzenberg, A. Wiedensohler, B.G. Martinsson, T. Deshler, C.A.M Brennkmeijer and A. Zahn, An optical particle size spectrometer for aircraft-borne measurements in IAGOS-CARIBIC, *Atmos. Meas. Tech.* 9, 2179-2194, doi:10.5194/amt-9-2179-2016, 2016.

605

Hess, P. G., A comparison of two paradigms: The relative global roles of moist convective versus nonconvective transport, *Journal of Geophysical Research* 110(D20): D20302, 2005.

Hoor P., H. Fischer, L. Lange and J. Lelieveld, Seasonal variations of a mixing layer in the lowermost stratosphere as identified by the CO-O<sub>3</sub> correlation from in situ measurements, *J. Geophys. Res.*, 107, doi:10.1029/2000JD000289, 2002.

610

Höpfner M., C.D. Boone, B. Funke, N. Glatthor, U. Grabowski, A. Günther, S. Kellmann, M. Kiefer, A. Linden, S. Lossow, H.C. Pumphrey, W.G. Read, A. Roiger, G. Stiller, H. Schlager, T. von Clarmann and K. Wissmüller, Sulfur dioxide (SO<sub>2</sub>) from MIPAS in the upper troposphere and lower stratosphere 2002-2012, *Atmos. Chem. Phys.*, 15, 7017-7037, doi:10.5194/acp-16-7017-2015, 2015.

615

Jäger H. and T. Deshler, Lidar backscatter to extinction, mass and area conversions for stratospheric aerosols based on midlatitude balloonborne size distribution measurements, *Geophys. Res. Lett.*, 29, 1929, doi:10.1029/2002GL015609, 2002.

Kiley, C. M., and H. E. Fuelberg, An examination of summertime cyclone transport processes during Intercontinental Chemical Transport Experiment (INTEX-A), *Journal of Geophysical Research*, 111(D24S06), 2006.

620

Kojima, T., Buseck P. R., Wilson J. C., Reeves J. M. and Mahoney M. J., Aerosol particles from tropical convective systems: Cloud tops and cirrus anvils, *J. Geophys. Res.*, 109, D12201, doi:10.1029/2003JD004504, 2004.

Luan Y. and L Jaeglé, (2013), Composite study of aerosol export events from East Asia and North America. *Atmos. Chem. Phys.* 13, 1221-1242, doi:10.5194/acp-13-1221-2013.

625

Martinsson B. G., Papaspiropoulos G., Heintzenberg J. and Hermann M., Fine mode particulate sulphur in the tropopause region from intercontinental commercial flights. *Geophys. Res. Lett.* 28, 1175 – 1178, 2001.

Martinsson, B. G., Nguyen H. N., Brennkmeijer C. A. M., Zahn A., Heintzenberg J., Hermann M., and van Velthoven P. F. J., Characteristics and origin of lowermost stratospheric aerosol at northern midlatitudes under volcanically quiescent conditions based on CARIBIC observations, *J. Geophys. Res.*, 110, D12201, doi:10.1029/2004JD005644, 2005.

630

Martinsson B. G., Brennkmeijer C. A. M., Carn S. A., Hermann M., Heue K.-P., van Velthoven P. F. J. and Zahn A., Influence of the 2008 Kasatochi volcanic eruption on sulfurous and carbonaceous aerosol constituents in the lower stratosphere. *Geophys. Res. Lett.*, 36, L12813, doi:10.1029/2009GL038735, 2009.

635

Martinsson, B. G., J. Friberg, S. M. Andersson, A. Weigelt, M. Hermann, D. Assmann, J. Voigtländer, C. A. M. Brennkmeijer, P. J. F. van Velthoven, and A. Zahn (2014), Comparison between CARIBIC aerosol samples analysed by accelerator-based methods and optical particle counter measurements, *Atmos. Meas. Tech.*, 7, doi: 10.5194/amt-7-2581-2014.

McCormick M.P., L.W. Thomason and C.R. Trepte, Atmospheric effects of the Mt Pinatubo eruption. *Nature* 373, 399-404, 1995.

640

- Meehl, G. A. & Teng, H. CMIP5 multi-model hindcasts for the mid-1970s shift and early 2000s hiatus and predictions for 2016–2035. *Geophys. Res. Lett.* 41, 1711-1716, 2014.
- Murphy, D. M., Thomson D. S. and Mahoney M. J., In situ measurements of organics, meteoritic material, mercury, and other elements in aerosols at 5 to 19 kilometers, *Science*, 282(5394), doi: 10.1126/science.282.5394.1664, 1998.
- 645 Myhre, G. et al. Anthropogenic and Natural Radiative Forcing. In: *Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change*. Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 2013.
- Neely III R.R., O.B. Toon, S. Solomon, J.P. Vernier, C. Alvarez, J.M. English, K.H. Rosenlof, M.J. Mills, C.G. Bardeen, J.S. Daniel and J.P. Thayer, Recent anthropogenic increases in SO<sub>2</sub> from Asia have minimal impact on stratospheric aerosol. *Geophys. Res. Lett.*, 40, 999-1004, doi:10.1002/grl.150263, 2013.
- 650 Neely III R.R., Yu P., Rosenlof K.H., Toon O.B., Daniel J.S., Solomon S. and Miller H.L., The contribution of anthropogenic SO<sub>2</sub> emissions to the Asian tropopause aerosol layer. *J. Geophys. Res. Atmos.*, 119, 1571-1579, doi:10.1002/2013JD020578, 2014.
- 655 Nguyen H. N., Gudmundsson A. and Martinsson B. G., Design and calibration of a multi-channel aerosol sampler for studies of the tropopause region from the CARIBIC platform. *Aerosol Science and Technology* 40, 649-655, 2006.
- Nguyen H. N. and Martinsson B. G., Analysis of C, N and O in aerosol collected on an organic backing using internal blank measurements and variable beam size. *Nucl. Instr. and Meth. B* 264, 96-102, 2007.
- 660 Nguyen H. N., Martinsson B. G., Wagner J. B., Carlemalm E., Ebert M., Weinbruch S., Brenninkmeijer C. A. M., Heintzenberg J., Hermann M., Schuck T., van Velthoven P. F. J. and Zahn A., Chemical composition and morphology of individual aerosol particles from a CARIBIC flight at 10 km altitude between 50° N and 30° S. *J. Geophys. Res.*, 113, D23209, doi:10.1029/2008JD009956, 2008.
- NOAA National Centers for Environmental Information, State of the Climate: Global Analysis for October 2016, published online November 2016, retrieved on December 5, 2016
- 665 from <http://www.ncdc.noaa.gov/sotc/global/201610>.
- Oram, D. E., Mani F. S., Laube J. C., Newland M. J., Reeves C. E., Sturges W. T., Penkett S. A., Brenninkmeijer C. A. M., Röckmann T. and Fraser P. J., Long-term tropospheric trend of octafluorocyclobutane (c-C<sub>4</sub>F<sub>8</sub> or PFC-318), *Atmos. Chem. Phys.*, 12, doi: 10.5194/acp-12-261-2012, 2012.
- 670 Osman M.K., D.W. Tarasick, J. Liu, O. Moeini, V. Thouret, V.E. Fioletov, M. Parrington and P. Nédélec, Carbon monoxide climatology derived from the trajectory mapping of global MOZAIC-IAGOS data, *Atmos. Chem. Phys.*, 16, 10263-10282, doi:10.5194/acp-16-10263-2016, 2016.
- Papasiropoulos G., Martinsson B. G., Zahn A., Brenninkmeijer C. A. M., Hermann M., Heintzenberg J., Fischer H. and van Velthoven P. F. J., Aerosol elemental concentrations in the tropopause region from intercontinental flights with the CARIBIC platform. *J. Geophys. Res.* 107(D23), 4671, doi:10.1029/2002JD002344, 2002.
- 675 Prata, A. J., and C. Bernardo, Retrieval of volcanic SO<sub>2</sub> column abundance from atmospheric infrared sounder data, *J. Geophys. Res.*, 112, doi: 10.1029/2006JD007955, 2007.

- Ramanathan V. and Y. Feng, On avoiding dangerous anthropogenic interference with the climate system: formidable challenges ahead, *Proc. Natl. Acad. Sci. USA*, 105, 14245-14250, 2008.
- 680 Randel W.J. and Park M., Deep convective influence on the Asian summer monsoon anticyclone and associated tracer variability observed with the Atmospheric Infrared Sounder (AIRS). *J. Geophys. Res.*, 111, D12314, doi:10.1029/2005JD006490, 2006.
- Randel W.J., Park M., Emmons L., Kinnison D., Bernath P., Walker K.A., Boone C. and Pumphrey H, Asian monsoon transport of pollution to the stratosphere. *Science* 238, 611-613, 2010.
- 685 [Rollins A.W., et al., The role of sulfur dioxide in stratospheric aerosol formation evaluated by using in situ measurements in the tropical lower stratosphere. \*Geophys. Res. Lett.\*, 44, 4280-4286, doi:10.1002/2017GL072754, 2017](#)
- Rosen J.M., The boiling point of stratospheric aerosols, *J. Appl. Meteorol.*, 10, 1044-1045, 1971
- Santer, B. D. et al. Volcanic contribution to decadal changes in tropospheric temperature. *Nat. Geosci.* 7, 185-189, 2014.
- 690 Schuck T. J., Brenninkmeijer C. A. M., Slemr F., Xueref-Remy I. and Zahn A., Greenhouse gas analysis of air samples collected onboard the CARIBIC passenger aircraft, *Atmos. Meas. Tech.*, 2(2), 449-464, 2009.
- Schwarz, J. P., Spackman J. R., Gao R. S., Watts L. A., Stier P., Schulz M., Davis S. M., Wofsy S. C. and Fahey D. W., Global-scale black carbon profiles observed in the remote atmosphere and compared to models, *Geophys. Res. Lett.*, 37, doi: 10.1029/2010GL044372, 2010
- 695 Skerlak B., M. Sprenger and H. Wernli, A global climatology of stratosphere-troposphere exchange using the ERA-Interim data set from 1979 to 2011, *Atmos. Chem. Phys.* 14, 913-937, doi:10.5194/acp-14-913-2014, 2014.
- Slemr F., A. Weigelt, R. Ebinghaus, H.H. Kock, J. Bödewadt, C.A.M Brenninkmeijer, A. Rauthe-Schöch, S. Weber, M. Hermann, J. Becker, A. Zahn and B.G. Martinsson, Atmospheric mercury measurements onboard the
- 700 CARIBIC passenger aircraft, *Atmos. Meas. Tech.* 9, 2291-2302, doi:10.5194/amt-9-2291-2016, 2016.
- Solomon S., Daniel J. S., Neely III R. R., Vernier J. – P., Dutton E. G. and Thomason L. W., The persistently variable “background” stratospheric aerosol layer and global climate change. *Science*, 333, 866 – 870, 2011.
- Stohl A., A 1-year Lagrangian “climatology” of airstreams in the northern hemisphere troposphere and lowermost stratosphere, *J. Geophys. Res.*, 106, 7263-7279, 2001
- 705 Sunilkumar S.V., Muhsin M., Venkam Ratnam M., Parameswaran K., Krishna Murthy B.V. and Maria Emmanuel, Boundaries of tropopause layer (TTL): A new perspective based on thermal and stability profiles. *J. Geophys. Res. Atmos.*, 122, doi:10.1002/2016JD025217, 2017.
- TF-HTAP-2010: Task Force on Hemispheric Transport of Air Pollution: Hemispheric Transport of Air Pollution 2010, *Air Pollut. Stud.* 17, edited by: Dentener, F., Keating, T., and Akimoto, H., UNECE, Geneva, Switzerland, available at: <http://www.htap.org/>, 2010.
- 710 Thomas, H. E., I. M. Watson, S. A. Carn, A. J. Prata, and V. J. Realmuto, A comparison of AIRS, MODIS and OMI sulphur dioxide retrievals in volcanic clouds, *Geomatics, Natural Hazards and Risk*, 2(3), doi: 10.1080/19475705.2011.564212, 2011
- Thomason L.W. and Vernier J.-P., Improved SAGE II cloud/aerosol categorization and observations of the Asian tropopause aerosol layer: 1989-2005. *Atmos. Chem. Phys.* 13, 4605-4616, doi:10.5194/acp-13-4605-2013, 2013.
- 715

- Vernier, J. P., J. P. Pommereau, A. Garnier, J. Pelon, N. Larsen, J. Niesen, T. Christensen, F. Cairo, L. W. Thomason, T. Leblanc and I. S. McDermid, Tropical stratospheric aerosol layer from CALIPSO lidar observations. *J. Geophys. Res.* 114, doi: 10.1029/2009JD011946, 2009.
- 720 Vernier, J.-P., L. W. Thomason, J.-P. Pommereau, A.E. Bourassa, J. Pelon, A. Garnier, A. Hauchecorne, L. Blanot, C. Trepte, D. Degenstein and, F. Vargas, Major influence of tropical volcanic eruptions on the stratospheric aerosol layer during the last decade, *Geophys. Res. Lett.*, 38, L12807, doi:10.1029/2011GL047563, 2011a.
- Vernier J.-P., Thomason L.W. and Kar J., CALIPSO detection of an Asian tropopause aerosol layer. *Geophys. Res. Lett.*, 38, L07804, doi:10.1029/2010GL046614, 2011b.
- 725 Vernier J.-P., Farlie T.D., Natarajan M., Wienhold F.G., Bian J., Martinsson B.G., Crumeyrolle S., Thomason L.W. and Bedka K.M., Increase in upper tropospheric and lower stratospheric aerosol levels and its potential connection with Asian pollution. *J. Geophys. Res. Atmos.*, 120, doi:10.1002/2014JD022372, 2015.
- Winker D.M., et al., The CALIPSO mission – A 3D view of aerosols and clouds. *Bull. Am. Meteorol. Soc.*, 91, 1211-1229, 2010.
- 730 Xu, L., Okada K., Iwasaka Y., Hara K., Okuhara Y., Tsutsumi Y. and Shi G., The composition of individual aerosol particle in the troposphere and stratosphere over Xianghe (39.45\_N, 117.0\_E), China, *Atmos. Environ.*, 35, 3145–3153, doi:10.1016/S1352-2310(00)00532-X, 2001.
- Yu P., Toon O.B., Neely R.R., Martinsson B.G., and Brenninkmeijer C.A.M., Composition and physical properties of the Asian tropopause aerosol layer and the North American tropospheric aerosol layer. *Geophys. Res. Lett.*, 42, doi:10.1002/2015GL063181, 2015.
- 735 Zahn A., J. Weppner, H. Widmann, H. Schlote-Holubek, B. Burger, T. Kühner and H. Franke, A fast and precise chemiluminescence ozone detector for eddy flux and airborne application, *Atmos. Meas. Tech.* 5, 363-375, doi:10.5194/amt-5-363-2012, 2012.
- Zbinden R.M., V. Thouret, P. Ricaud, F. Carminati, J.-P. Cammas and P. Nédélec, Climatology of pure tropospheric profiles and column contents of ozone and carbon monoxide using MOZAIC in the mid-northern latitudes (24° N to 50° N) from 1994 to 2009, *Atmos. Chem. Phys.*, 13, 12363-12388, doi:10.5194/acp-13-12363-2013, 2013.
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**Tables**

**Table 1.** Most significant volcanic eruptions for the aerosol concentration in the northern hemisphere LMS in the time period studied.

<b>Volcano</b>	<b>Date</b>	<b>Lat./Long.</b>	<b>SO<sub>2</sub> (Tg)</b>
Manam	27 Jan 2005	4° S / 145° E	0.1 <sup>a</sup>
Soufriere Hills	20 May 2006	17° N / 62° W	0.2 <sup>b</sup>
Rabaul	7 Oct 2006	4° S / 152° E	0.2 <sup>a</sup>
Jebel at Tair	30 Sep 2007	16° N / 42° E	0.1 <sup>c</sup>
Okmok	12 Jul 2008	53° N / 168° W	0.1 <sup>c</sup>
Kasatochi	7 Aug 2008	52° N / 176° W	1.7 <sup>c</sup>
Redoubt	23 Mar 2009	60° N / 153° W	0.1 <sup>d</sup>
Sarychev	12 Jun 2009	48° N / 153° E	1.2 <sup>e</sup>
Grimsvötn	21 May 2011	64° N / 17° W	0.4 <sup>f</sup>
Nabro	12 Jun 2011	13° N / 42° E	1.5 <sup>f</sup>

745 <sup>a</sup>Prata and Bernardo, 2007

<sup>b</sup>Carn and Prata, 2010

<sup>c</sup>Thomas et al., 2011

<sup>d</sup>Brühl et al., 2015

<sup>e</sup>Haywood et al., 2010

750 <sup>f</sup>Clarisse et al., 2012

Figures

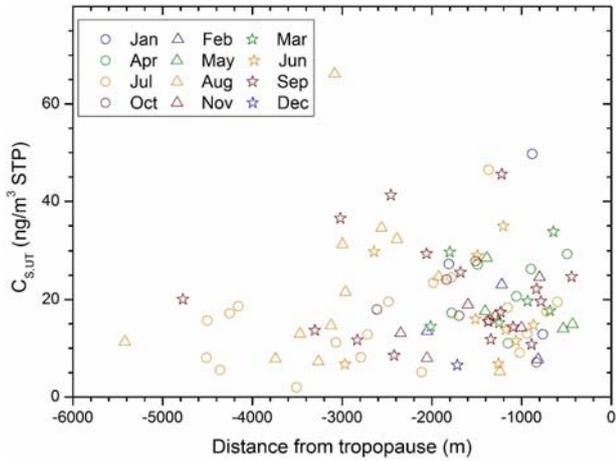


Figure 1: Upper tropospheric particulate sulfur concentration related to distance from the tropopause, where the symbols indicate sampling month.

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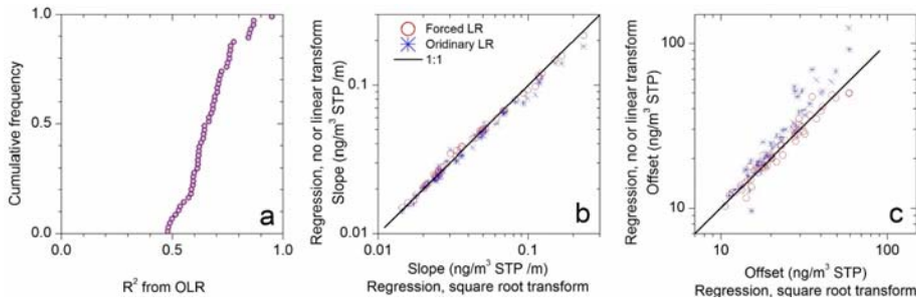


Figure 2: a) Cumulative frequency of the coefficient of determination ( $R^2$ ) of ordinary linear regression (OLR) between particulate sulfur concentration and distance from the tropopause of the 52 data groups used in this study. b) and c) Comparison of slope (Fig. 2b) and offset (c) between a square root transformed dependent variable and both OLR and forced linear regression.

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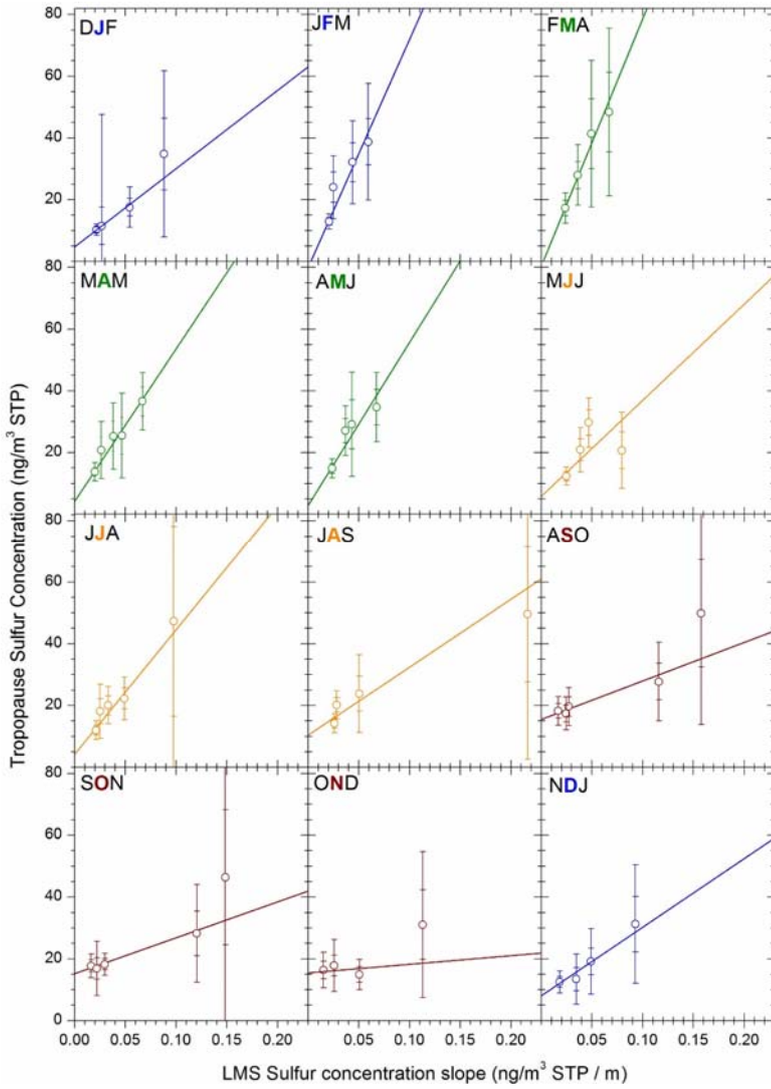
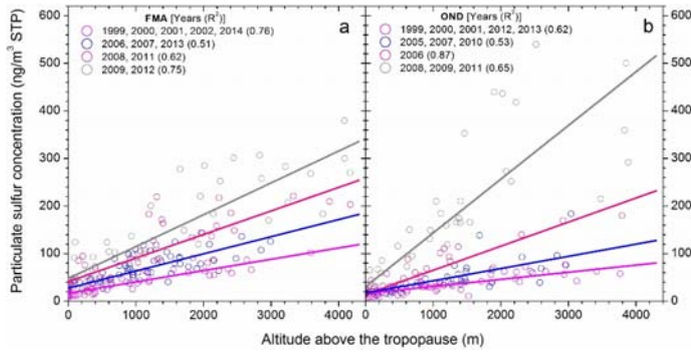
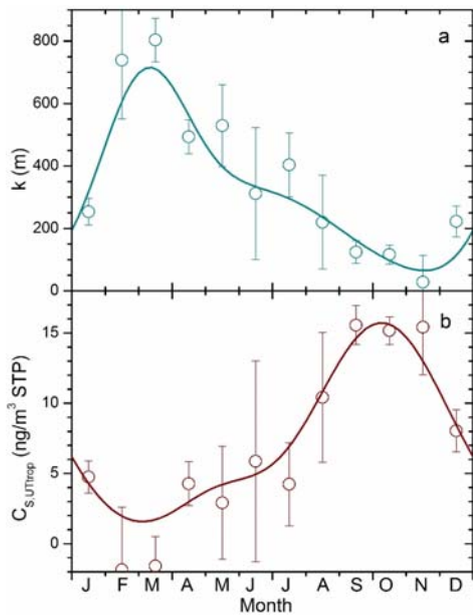


Figure 3: Weighted regression between slopes and offsets of all groups of each season. The offset indicates the tropopause (and UT) concentration of tropospheric origin ( $C_{S,UTrop}$ ) and the slope of the fit ( $k$ ) expresses the sensitivity of the UT concentration to changes in the stratospheric concentration slope (a). The error bars show and student-t 70% and 95% confidence interval, respectively.

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770 **Figure 4:** Examples of first step forced linear regression between the particulate sulfur concentration and the altitude above the tropopause for the data groups of two seasons: a) FMA (February, March and April) and b) OND (October, November and December). The legend shows which years are included in the respective data groups followed by the coefficient of determination in paranthesis.



775 **Figure 5:** Seasonal variation of (a) the sensitivity of the UT particulate sulfur concentration to the concentration slope in the LMS ( $k$ ), and (b) the UT particulate sulfur concentration of tropospheric origin ( $C_{S,UTrop}$ ).

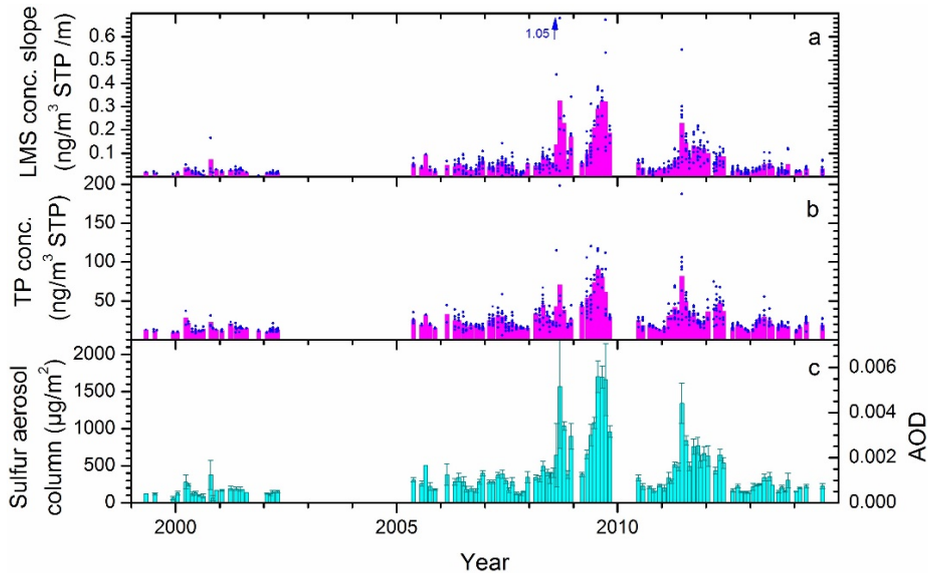


Figure 6: Estimated a) slopes and b) offsets based on individual measurements (dots) and monthly averages (magenta bars). c) Sulfur aerosol column of the lowest 3000 m of the LMS with standard errors, assuming particles of 75% sulfuric acid and 25% water, and AOD (right y axis) assuming stratospheric background aerosol particle size distribution.

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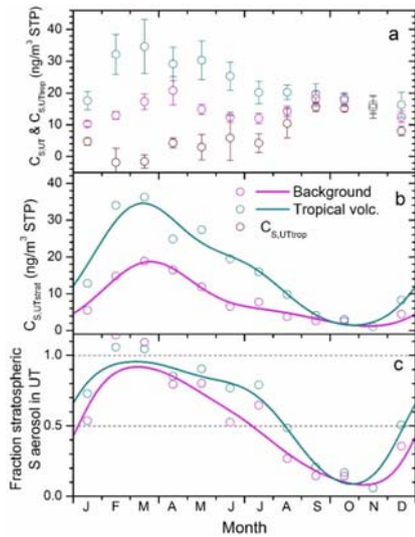


Figure 7. Seasonal variation of a) the UT particulate sulfur concentration during LMS background conditions and moderate volcanic influence, together with the estimated UT particulate sulfur concentration of tropospheric origin ( $C_{s,UTtrop}$ ), b) UT particulate sulfur concentration of stratospheric origin ( $C_{s,UTstrat}$ ) obtained by subtracting the  $C_{s,UTtrop}$  from the two UT concentration cases in Fig. 7a, and, c) the ratio  $C_{s,UTstrat}/C_{s,UT}$  for the two cases.

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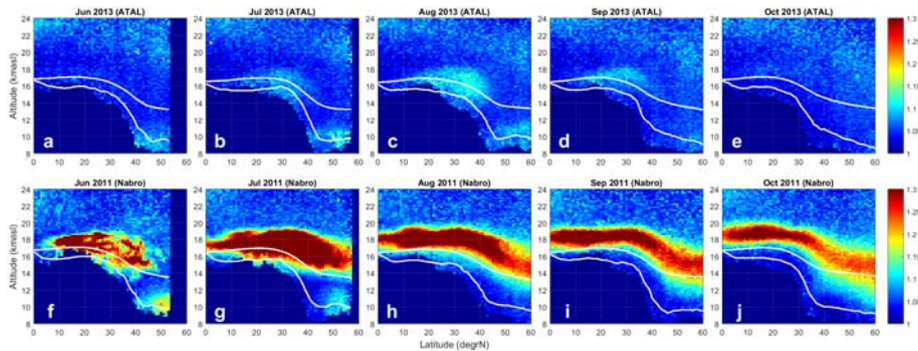


Figure 8. Latitude and altitude distribution of scattering from aerosol in the stratosphere and the UT. Two series, from 2013 (upper row, a - e) and 2011 (lower row, f - j), show the monthly mean 532 nm wavelength scattering ratio from June to October without respectively with fresh volcanic influence. The former series allows identification of the comparably weak ATAL in July to September. The lower series includes two volcanic eruptions: the high altitude cloud from the tropical volcano Nabro (eruption in June 12, 2011) and the low altitude and midlatitude faint volcanic cloud from Grimsvötn (May 21, 2011). The two white lines in every graph show the monthly averaged positions of 380 K potential temperature (upper line) and the 1.5 PVU dynamical tropopause.

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