We thank the reviewers for their constructive comments, suggestions, and corrections. This version has been largely improved owing to their helpful review. In our revision, we have addressed the reviewer’s concerns. The following is a one-to-one response to their questions.

Response to Comments by Anonymous Referee #1

General Comments:
The authors provide a detailed analysis for the long-range transport of tropospheric O$_3$ from Africa to Asia. They indicated that African O$_3$ have important influences on free tropospheric O$_3$ over Asia, and the imported African O$_3$ peaks in winter because of the shifts of transport and emission patterns. I recommend the paper for publication after consideration of the points below.

1) The paper isn’t concise enough for me. For example, Section 5 provides a summary for the transport and emission processes, which is actually a repeat of Section 4.2. In addition, considering the small contribution from SHAF (shown by Figure 4), it may not be necessary to have an individual section (Section 4.3) to discuss its influence.

Thanks for the comments and suggestions. We have reconstructed the paper to make it more concise. Section 5 in the last version has been removed. The presentation is polished throughout the paper and more in-depth discussion is added. We have reconstructed the section on the interhemispheric transport of ozone from SNAF and added more analyses in this section (now section 4.2). Therefore, we think it is better to keep this section.

2) The discussion should be improved. The authors should explain why the seasonal variability of biogenic isoprene is so weak (Figure 6); and revise the discussion about the contributions from various sources (i.e. biogenic, biomass burning and lightning, Section 4.2).

Thanks for the points. In the last version, the color scale for the seasonal variation of biogenic isoprene in Figure 6 was not appropriate so that the seasonal variation was not shown apparently. We have edited the color scale so to better show the magnitude of the seasonal variation of biogenic isoprene (now Figure 8). The regional means of
the seasonal variation of biogenic isoprene over Africa is shown in Figure 9, which shows large seasonality of biogenic emissions. The discussion about the contributions from various sources has been revised and discussed in more depth (now in section 3.3).

Specific Comments:
1: Line 147-149 Are the O₃ production and loss rates generated using the full-chemistry simulation with the current model settings or from other studies (Wang et al. 1998; Zhang et al. 2008)?

In this study, we generated the ozone production and loss rates from the full-chemistry simulation using the current model settings of GEOS-Chem v9-02. We have clarified this in this revision (see section 2.1).

2: Line 149-153 It would be better to show these regions as boxes in the map (e.g. Figure 5). It is difficult to imagine the regions just based on these lat/lon numbers.

Thanks for your suggestion. We have added Fig. 1 to show the definitions of the regions. The sites used in the GEOS-Chem validation and trajectory analysis are also shown in Fig. 1.

3: Line 155-157 Did the authors evaluate the possible influences from interannual variations of meteorology on chemistry?

Thanks for the question. In this study, we focus on the impact of meteorology on the transport. Therefore, we keep the ozone production and loss rate fixed in one year and allow the meteorology to vary from year to year.

Yes, meteorology also affects chemistry. If keeping ozone production term constant, the meteorology influence on chemistry is ignored. We have pointed this out in this revision (section 2.1, the last 2nd paragraph). We have not directly evaluated this impact. Instead, we have tested if this impact will significantly alter our results. Therefore, we conduct a sensitivity test. First, we run GEOS-Chem in full chemistry mode to generate ozone production rate and lose frequency in other two different years. Years
2001 and 2004 are selected because these are years when the extreme anomalies of imported African ozone appear in Asia. In the two full chemistry simulations, we used the same anthropogenic and biomass burning emissions in 2005 but with the different meteorology in 2001 and 2004 respectively. Therefore, the differences in the ozone production rate and loss frequency data can be regarded as the meteorology influences on chemistry as the emissions are fixed. Then we use these two sets of daily ozone production and loss frequency to run the tagged ozone simulations for 20 years. First, we compare the 20-year mean of imported African ozone from these two runs with the default run in Fig. 1. The differences of imported African ozone over Asia between three datasets are small, varying from -1 ppbv to 0.2 ppbv for most layers and months, although the differences are large in NH winter over the upper troposphere, reaching 1-3 ppbv.

Second, we compare the three datasets on the interannual variations of imported African ozone over Asia in Fig. 2. Indeed, there are differences between the three datasets. However, the interannual variations of the three simulations are similar and all positive correlated to the ITCZ. This sensitivity test suggests that our treatment is robust in capturing the variation of ozone transport from Africa to Asia from year to year.

4: Line 166-167 Is there any other station available? Why are these three stations selected?

GEOS-Chem simulations have been validated with ozonesonde data extensively, for example, in North America (Zhang et al., 2008; Zhu et al., 2017b), Europe (Kim et al., 2015), and East Asia (Wang et al., 2011; Zhu et al., 2017a; Zhu et al., 2017b). However, few studies have validated the simulations in Africa. We specifically validate the performance of GEOS-Chem over Africa for an enhanced confidence on our analysis. These three stations are selected for their representative locations and relative long records in Africa. In this version, the ozone data from three ozonesonde stations in India are added to compare with the GEOS-Chem simulations. In addition, the Tropospheric Emission Spectrometer (TES) satellite observations are compared with
the GEOS-Chem simulation in the middle troposphere shown in the supplementary file (Fig. S6).

Figure S1. The differences of imported African ozone over Asia between the 2005 and the 2001 datasets (1st col.) and between the 2005 and the 2004 dataset (2nd col.). The values are averaged over 60-145°E from 1987 to 2006. Associated with the three datasets, the ozone production and loss data are generated with the same emissions but different meteorology in 2001, 2004, and 2005.
Figure 2. The interannual variations of the anomaly of imported African ozone over Asia simulated using the three sets of ozone production and loss data at (a) 200 hPa, (b) 600 hPa, and (c) 975 hPa in January. Associated with the three datasets, the ozone production and loss data are generated with the same emissions but different meteorology in 2001, 2004, and 2005.
5: Section 2.3 Is the meteorological data the same as used by the HYSPLIT model? If they are the same, it would be better to combine Section 2.2 with Section 2.3.

The meteorological data are the same as what are used by the HYSPLIT model. The two parts have been combined into Section 2.3.

6: Line 237-239 The influence of African O₃ to south America across Atlantic is discussed, but isn’t shown in the Figure. It could be better to remove the discussion about the transatlantic transport here.

The discussion about the transatlantic transport here has been removed.

7: Line 262-269 Although may not be necessary to explain, I am just curious about the reason for the discrepancy between western and eastern Africa.

In general, the latitudinal position of ITCZ follows the rotation of the sun. In eastern Africa, the seasonal migration of ITCZ with latitude is more symmetrical around the equator, while in western Africa, the migration is limited (Collier and Hughes, 2011). The seasonal migration of the ITCZ in western Africa is complicated. Generally, in NH summer, the convergence zone is formed by the flows from the Atlantic cold tongue and the Saharan heat low, locating around 20°N (Nicholson, 2009, 2013). In NH winter, the anticyclonic wind from northern Africa converges with the southerly wind from Atlantic. The ITCZ over western Africa still stays in the continent (Nicholson, 2013). Therefore, the seasonal migration of the ITCZ in western Africa is within a narrower range of latitudes than in eastern Africa.

8: Line 276-277 Figure 6 shows significant seasonal variation for biomass burning CO. Surprisingly, the seasonal variation of biogenic isoprene is ignorable, which seems inconsistent with other study (e.g. Marais et al. 2014). Is it associated with the color scale? On the other hand, the normalized magnitudes of seasonal variability (Figure 7) are comparable between CO and isoprene. Is it due to the usage of standard deviation in the calculation? The approach for normalization is confusing.

Marais, E. A., Jacob, D. J., Guenther, A., Chance, K., Kurosu, T. P., Murphy, J. G.,

Thanks for the points. Yes, the narrow seasonal variation of biogenic isoprene shown in the old Fig. 6 indeed is due to the use of the color scale. We have edited the color scale in the figure (now Fig. 8) so that the seasonal variation of biogenic isoprene in Africa is better presented. Fig. 9 (old Fig. 7) shows the seasonal variability of isoprene and CO. The biogenic emission peaks in spring and autumn. The magnitude of biogenic isoprene is comparable to the results in Marais et al. (2014).

Normalization is not taken in this revision to avoid confusing.

9: Line 315-362 The discussion in this section is superficial. The authors discuss the contributions from various sources without detailed calculations. For example, the authors indicated: 1) “In boreal spring, a region with high ozone concentrations (>40 ppbv) appears in higher altitudes and ... mainly due to the highest biogenic emissions in the NHAF” 2) “In boreal autumn, the locations of the ITCZ and the Hadley cell are similar to these in boreal spring. Ozone in the African middle troposphere ... attributed to stronger lightning NOx emission” However, there is no evidence to demonstrate that the contributions from biogenic and lightning activities are evaluated carefully.

The discussion is simply based on the spatial distribution of Figure 6. The biogenic and lightning activities are highly similar between spring and fall, and it is hard to explain why the spring-time O3 is biogenic dominant, whereas autumn-time O3 is lightning dominant.

Thanks for the comments. Aghedo et al. (2007) has suggested that the biogenic and lightning emissions are the two important sources influencing African middle and upper tropospheric ozone and affecting global tropospheric ozone burden. To further explore the differences between the situations in NH spring and in NH autumn, we have conducted 3 sensitivity experiments by switching off the biogenic, lightning, and biomass burning emissions, respectively. The separate contribution of the three
sources to tropospheric ozone over Africa is shown in the supplementary file (Fig. S7). In NH spring and autumn, the influence of ozone from SHAF on Asia is small and similar (Fig. 7) so we can mainly focus on ozone in NHA. It appears that in NH spring, elevated ozone abundances from biogenic emissions are higher than that from lightning NOx emissions while in NH autumn, elevated ozone abundances from biogenic emissions are lower than that from lightning NOx emissions (Fig. S7, and also see Aghedo et al. (2007)). We have revised our paper accordingly. We also have made the discussion in more depth in section 3.3.4.

10: Section 4.2 It seems that Figure 8 and Figure 9 are already sufficient for the discussion. I suggest to remove Figure 10 to make the paper more concise. Thanks for the points. Fig. 10 (in the last version) has been removed.
Response to Comments by Anonymous Referee #2

This is a relatively straightforward analysis of the interplay of meteorological processes and atmospheric chemistry in venting out ozone and ozone precursors from Africa and reaching Africa. The manuscript is well written, the figures appropriate.

As the authors state- there is relatively little literature discussing Africa-to-Asia transport, so this is a welcome addition, despite it doesn’t make use of the recommendation in the HTAP2 exercise to harmonize region definitions to allow comparability of results.

We thank the reviewer for the encouragement and for the valuable and thoughtful comments. In Fig. 1, we show the regional definitions in this study and in HTAP2, so the reader can have an idea for the similarity and difference between the two definitions.

A minor remark is that I don’t see terribly much added value of the trajectory analysis in figure 12.

The trajectory analysis is improved in this revision as follows. (1) Trajectories in two more seasons, spring and autumn, are added for comparison between the four seasons instead of just between two seasons in the last version, (2) trajectories from four more stations in Africa are added for a wider coverage and representation, (3) the mean transport paths for the trajectories that arrive Asia are illustrated with lapse times indicated, and (4) more discussions are added in this revision to supplement the discussion on the mechanisms that control the transport of African ozone to Asia throughout section 3.3 and in section 4.2 and in the conclusions (section 5).

Although some attempt has been made to demonstrate the model’s ability to model ozone over Africa, I think this could be done more convincingly- there is meanwhile a host of other observations (surface, aircraft, satellite tropospheric ozone columns) that could be explored.

Thanks for the point. In this revision, we add more validations between the
ozonesonde observations and GEOS-Chem simulations from perspectives of variceal profiles and the seasonal and interannual variability of tropospheric ozone over Africa in the surface layer, lower, middle, and upper troposphere. The comparisons at three more ozonesonde stations in India are added. Furthermore, the ozone data from the Tropospheric Emission Spectrometer (TES) satellite instrument are used to evaluate the GEOS-Chem simulation in the middle troposphere (464 hPa) globally in the supplementary file (Fig. S6). The detailed comparisons are shown in Figs. 2-4 and Tables1-2 and are discussed in section 2.2.

Are signals from African ozone visible in soundings over India?

The transport of airmass from African in summer is reflected in the ozonesonde data at Poona and Thiruvananthapuram in western India (in an added figure, Fig. 4). The impact of the Somali jet on the ozone in western India is obvious as low ozone concentrations appear in the lower troposphere in NH summer.

The organization and discussion of methods could be somewhat more systematic.

Thanks for the suggestions. The methods are reorganized and the presentation is polished in section 2.

I suggest that the authors explore somewhat further these aspects, and recommend the manuscript to be accepted after taking these major and minor comments below into account.

Thanks. We have followed these suggestions.

Minor comments.

1. 11-30 the abstract could be somewhat more explicit in describing the regions and attribution methodology.

Thanks. The abstract is rewritten to explicitly state the regional definitions and attribution methods.
l. 16 Replace boreal by NH winter. Or find better way of describing which months are discussed. Are the > and < really meant to express minima and maxima?

Thanks. We have replaced boreal winter with northern hemisphere (NH) winter in this revision. We have removed the > and <, and used certain numbers to express minima and maxima.

l. 30 I miss some statement on the relevance of this analysis. How much of the Asian ozone was produced in Africa or from African precursor emissions- where is it most important (not only vertical but also geographically.

Vertically, the influence of African ozone on Asia is mainly in the middle and upper troposphere. Geographically, the imported African ozone mainly distributes over latitudes south to 40°N in Asia. We have added more detail about this in the abstract.

l. 35 give reference time to which this RF estimate pertains.

Thanks. The reference time is given: “It also acts as a greenhouse gas, whose global mean radiative forcing is about 0.4 ± 0.2 W/m² for the period 1750-2011 (Myhre et al., 2013)”.

l. 46 add: as well as a range of papers in the HTAP2 (Galmarini 2016) special issue.

Thanks. Galmarini et al. (2017) and several other papers in the HTAP2 special issue are now added in the citations.

l. 53 The issue is also very connected to legislative issues related to the control of ozone and ozone exceedance in the western states of the USA, e.g. as discussed in Huang et al. (2017; already cited).

Thanks. This point is now included in Introduction.

l. 54 One reference on LRT transport between South Asia and East Chakraborty et al. Science of the Total Environment, 523, 2015

Thanks for this recommendation. The reference is helpful to the study and it is added
in the reference.

1. 73 … makes a contribution … how is contribution defined? Zero out of emissions? This is important because later you present a different method.

We have clarified this term in Introduction. In this study, we used the tagged ozone simulation to track ozone from source regions to a receptor region. We did not use the perturbation simulations that turn the emissions off to see the contribution of source regions to a receptor region.

The contribution of the source regions to the receptor region can be presented as absolute and fractional contributions. The former refers to the concentrations of the imported ozone in a unit of ppbv, while the latter is the ratio of the imported ozone to the total ozone in a grid, a layer, or a region. We have also described the term in Introduction and rephrased this sentence to make it clear.

1. 117 which resolution is used for GEOS-CHEM; what was the underlying resolution of the assimilation product. Importantly for this paper, how is convection parameterized, is there any evaluation over Africa of these process. Interhemispheric mixing and similar: refer to any relevant application of the model that demonstrates it is fit-for-purpose for this study. I realize that these are discussed later, but I would have expected these descriptions here.

Thanks for these comments. The detailed descriptions of GEOS-Chem have been moved to an earlier part in section 2.1. In this study, the simulations are driven by GEOS-4 meteorology at a 4° latitude by 5° longitude horizontal resolution, degraded from their native resolution of 1° latitude × 1.25° longitude. There are 30 vertical layers including 17 levels in the troposphere (see the 2nd paragraph in section 2.1). GEOS-4 uses the schemes developed by Zhang and McFarlane (1995) for deep convection and by Hack (1994) for shallow convection. GEOS-4 meteorology is found to be characterized with stronger deep convection in tropics than GEOS-5 (Liu et al., 2010; Zhang et al., 2011). Liu et al. (2010) and Zhang et al. (2011) have shown good agreement of GEOS-Chem simulations driven by GEOS-4 with satellite observations.
in the tropical troposphere. Choi et al. (2017) compared the simulations of the Global Modeling Initiative (GMI) chemistry and transport model (CTM) driven by three meteorological data sets (fvGCM for 1995, GEOS-4 for 2005, MERRA for 2005) with ozonesonde and TES observations. They found that ozone simulated by GEOS-4 has the highest correlation with the observations. These previous studies and the good validation results over Africa from this study provide us confidence on the model performance. We have provided more details on the model descriptions in this revision (see section 2.1).

1. 125- If I understand correctly the authors merge the EDGAR3.2 global inventory with regional ones. Which period? How do these inventories compare with e.g. the HTAP2 inventory for 2008/2010 in this special issue, or EDGAR4.2 products (for time series).

We conducted the full chemistry simulation in GEOS-Chem to generate ozone production and loss data in 2005. The merged emission data are from the global EDGAR 3.2 inventory in the base year 2000. The regional emission inventories include the INTEX-B Asia emissions inventory in 2005 with base year 2006, the NEI05 inventory in North America in the base year 2005, the EMEP inventory in Europe in the base year 2005, the BRAVO inventory in Mexico in 2005 with base year 1999, and the CAC inventory in Canada in the base year 2005. This part has been described in more details in section 2.1.

We compared the anthropogenic emissions of CO and NOx from the GEOS-Chem for 2000 inventories with those in the HTAP2 inventories for 2008 (http://edgar.jrc.ec.europa.eu/htap_v2/) and showed the comparison in the supplementary file (Figs. S3 and S4). Compared to the HTAP2 inventory for 2008, the CO and NOx emissions in GEOS-Chem are lower in Africa throughout the year. The annual anthropogenic emissions of CO in Africa in GEOS-Chem and from the HTAP2 inventory are about 12.2 Tg yr\textsuperscript{-1} and 62.5 Tg yr\textsuperscript{-1}, respectively. The anthropogenic emissions for NOx in Africa in GEOS-Chem and HTAP are about 2.27 Tg yr\textsuperscript{-1} and 4.53 Tg yr\textsuperscript{-1}, respectively.
The contribution of anthropogenic emissions to generation of African ozone is considered to be smaller than that of other emissions. Aghedo et al. (2007) estimated that anthropogenic emissions emitted in Africa account for approximately 11% (4.7Tg/42.8Tg) of the total African emissions that impact the global tropospheric ozone burden. In this study, the annual biomass emissions of CO in Africa in GEOS-Chem are 182.8 Tg yr\(^{-1}\), which is much larger than anthropogenic emissions of CO (4.7Tg yr\(^{-1}\)). Nevertheless, we should consider the impact of this issue and we have stated in this revision that “Although the anthropogenic emissions contribute less significantly to the ozone generation in Africa, these differences in emission inventories imply that African ozone simulated by GEOS-Chem is with some uncertainties.” (Line 188-190).

1. 135 What is the global lightning source strength and specific for Africa. How does this compare to other studies.

Lightning NO\(_x\) emission used in the study is shown by annual mean and in each season in the supplementary file (Fig. S1). The annual global lightning NO\(_x\) source amount is 5.97 Tg N yr\(^{-1}\), comparable to 6±2 Tg N yr\(^{-1}\) in Martin et al. (2007) and 6.3 Tg N yr\(^{-1}\) in Miyazaki et al. (2014). Miyazaki et al. (2014) estimated the annual global lightning NO\(_x\) emission by assimilating observations of NO\(_2\), HNO\(_3\), and CO measured by OMI, MLS, TES, and MOPITT into the global chemical transport model CHASER. The annual lightning emission is 1.72 Tg N month\(^{-1}\) in Africa, 0.80 Tg N month\(^{-1}\) in NHAF, and 0.79 Tg N month\(^{-1}\) in SHAF, shown in the supplementary file. We have added the information in section 2.1 in the 3\(^{rd}\) paragraph.

1. 137 briefly describe what is the ‘standard’ tagged ozone method. Pro’s and con’s-limitations. Comes now later

The tagged ozone method tracks ozone that is generated in different regions. The method was first proposed by Wang et al. (1998) and then further developed and used by a number of studies (Fiore et al., 2002; Sudo and Akimoto, 2007; Zhang et al., 2008;
Liu et al., 2011; Sekiya and Sudo, 2012, 2014; Zhu et al., 2017b). This is one of the standard modes in GEOS-Chem. It is done by the following two steps. First, the daily production rates and loss frequencies of odd oxygen ($O_x = O_3 + NO_2 + 2NO_3 + 3N_2O_5 + HNO_3 + HNO_4 + PAN + PMN + PPN$) were generated and archived from a full chemistry simulation before the tagged ozone simulation. Since ozone accounts for most of $O_x$, ozone instead of $O_x$ is used for clarity. Second, GEOS-Chem is run again in the tagged ozone mode using the archived ozone production rate and loss frequency, with separate tracers for the ozone produced from each of the specific source regions. Therefore, the tagged ozone tracer method can assess the contributions of African ozone to Asia. The advantages of and issues with the tagged ozone simulation are discussed in section 2.1 (see the last paragraph in section 2.1) with an additional figure in the supplementary file (Fig. S5).

1. I expected this description earlier.

Thanks. The description has been moved into an earlier part of the section. Please see the 2nd paragraph in section 2.1.

1. 140-143: better include with the GEOSCHEM description.

Thanks. This part has been included with the GEOS-Chem description. Please see the 2nd paragraph in section 2.1.

1. 150- what is the reason for not simply taking the ‘african mask’- instead two blocks. I suggest adding a simple figure, showing these masks on top of a map (perhaps along with the HTAP2 definition of Africa).

Thanks for the suggestion. Fig. 1 is added to show the definitions of the source and receptor regions in this study. The definition of Africa in HTAP2 is also presented to show the similarity and difference between the two definitions. The reason for using the blocks to define regions is because this is the default way that GEOS-Chem uses for the tagged ozone simulation.
I think most papers that I know keep (anthropogenic) emissions constant— but that is not necessarily the same as keeping the production terms constant. What could be the impact?

*Thanks for this comment.* Meteorology affects both chemistry and transport, a physical process. If keeping ozone production term constant, the meteorology influence on chemistry is ignored. If keeping emissions constant, the impacts of meteorology on both chemistry and transport are considered. We have pointed this out in this revision (section 2.1, the last 2nd paragraph).

what about sfc observations, tropospheric residual from satellite. The comparison is fairly superficial

*Thanks.* In this revision, we have included more comparisons between the GEOS-Chem simulations and ozonesonde observations by adding 2 figures and two tables. The seasonal and interannual variation GEOS-Chem simulations have been further compared with ozonesonde data in the surface layer, lower, middle, and upper troposphere at the African sites. Three ozonesonde stations at India are added for the comparison. Figs. 2 and 3 and Tables 1 and 2 are added in section 2.2. The correlation coefficient (r), bias in percentage, and root-mean-square error (RMSE) between the simulations and the ozonesonde data are presented. The comparison in global distribution with the TES satellite ozone data are shown in the supplementary file (Fig. S6).

Imported ozone=>try to describe more exactly what it is. Region-average abundance of imported ozone (imported ozone could be also the flux through the western border, for instance).

*Thanks for the point.* We define ozone that is generated over Africa under the tropopause as African ozone. Following Holloway et al. (2008), “Imported ozone” is used to refer ozone that is distributed over the receptor region. In the paper, when we discuss African ozone over Asia, we use “imported African ozone” to differentiate it from the overall ozone concentrations in Asia. We have clarified this term in
Introduction (the last paragraph in section 1).

p. 295 /figure 6: I guess if the units are molec/cm²/s this pertains to the integrated amount over a model layer; otherwise it should rather be per cm³?

Thanks for the point. Yes, in the last version, the unit pertains to the integrated NOₓ amount over a model layer. We have converted the unit into NOₓ emissions per cubic meter per second. Therefore, the lightning NOₓ emissions are expressed as molec/m³/s in this revision (see Figs. 8 and 9).