**Anonymous Referee #2**

We are very grateful to the reviewer for reviewing this manuscript. We have carefully considered the suggestions and made revisions accordingly. Below we list detailed responses to the suggestions and comments. The suggestions and comments are in italics, followed by the response in normal font with changes highlighted in blue.

Cao et al. measure nitrate isotopes and concentrations in a 60-year firm core from South Pole and perform air-snow nitrate transfer simulations using the TRANSITS model to investigate whether nitrate isotopes at the site reflect changes in stratospheric ozone. The results are similar to previous Antarctic studies of ice core with similar snow accumulation rates that indicate d15N(NO3-) is insensitive to total column ozone. Decreases in the D17O(NO3-) record during the ozone hole are qualitatively attributed to atmospheric oxidation changes in the extratropical Southern Hemisphere nitrate source regions. The new dataset is a valuable contribution however, the manuscript could be improved by furthering our understanding of ice core nitrate isotopes in Antarctica which have a unique and not fully understood fingerprint. As such, I believe the authors have an opportunity to use the ice core dataset and the TRANSITS model to advance our understanding of ice core D17O(NO3-) to make a new and valuable contribution to the literature. I look forward to seeing the published.

**Suggestions for improvement**

A paper on nitrate isotopes in a snow pit (1960-2000) from the low-accumulation Dome A site was just published in June (Shi et al., 2022) and the authors conclude that nitrate isotopes (d18O, D17O, and d15N) record stratospheric ozone depletion and ultra-violet radiation at the Dome A site. The authors have discussed the modelled response of d15N(NO3-) to total column ozone at South Pole versus Dome A sites. Please update the manuscript in light of the newly published paper.


**Response:** Thanks for this and we have noted this paper in GRL. The Shi et al. (2022) paper observed changes in nitrate isotopes and concentrations in recent decades and concluded these are due to the effects of the ozone hole. However, carefully reviewing the figures and original data we found where the onsets of the isotope changes are not lined up with the onset of the ozone hole (neither the recovery of ozone hole and the corresponding changes). In addition, they also used the TRANSITs model to estimate changes in isotopes caused by the ozone hole, but the model parameters are not clear, e.g., snow e-folding depth, quantum yield of snow nitrate photolysis, using “similar parameters to Dome C as Erbland et al. (2015)” is impossible to get the reported model results in Shi et al. (2022). For example, “similar parameters to Dome C as Erbland et al. (2015)” as stated in Shi et al. 2022 will give a quantum yield of 0.026, applying this value to Dome A would lead to a modeled δ15N(NO3-) value of ~ 1150 ‰, more than 3 times of the observed value of ~ 300 ‰. What is more, our calculation indicated at Dome A the ozone hole can lead to changes in Δ17O(NO3-) by ~ 1 per mil the most, while the observed changes in Δ17O(NO3-) at Dome A is almost 5 per mil. In the revised manuscript we have
Now that there are a number of d15N(NO3-) measurements across Antarctica, a discussion on the sensitivity of d15N(NO3-) and D17O(NO3-) to total column ozone at various ice cores sites, including the new Dome A record, would be valuable addition for the community to make progress on the use of d15N(NO3-) and D17O(NO3-) as a UV or total column ozone proxy.  

**Response:** Thanks for this comment. Using TRANSITS model, we calculated the enrichments in δ15N(NO3-) and Δ17O(NO3-) caused by ozone hole alone (keeping other factors the same) in other East Antarctic sites including Dome A, Dome C, Vostok, Dome Fuji, and a west Antarctic site WAIS Divide in addition to the South Pole. In the revised manuscript, we added a new subsection (new subsection 4.3) to present this table with relevant discussion.  

4.3 Estimated effects of the ozone hole on snow nitrate isotopes in other Antarctic sites  
In order to search for signals of the ozone hole, we used the TRANSITS model to further explore the maximum possible responses of ice-core preserved δ15N(NO3-) and Δ17O(NO3-) to the ozone hole at other Antarctic sites, including Dome A, Dome C, Vostok, Dome Fuji, and the West Antarctic Ice Sheet (WAIS) Divide in addition to the South Pole. The responses are defined as the differences between the isotopes before the ozone hole period and those in years with the most depletion. The results are listed in Table 1. As shown in the table, except WAIS Divide, other sites all display bigger responses to the ozone hole, especially the three East Antarctic Plateau sites. These patterns are mainly determined by the differences in snow accumulations rates at these sites (Frezzotti et al., 2013; Erbland et al., 2015; Shi et al., 2022), i.e., lower snow accumulation rates correspond to longer durations of nitrate in the photic zone, leading to larger effects of the ozone hole. In particular, at Vostok, Dome C and Dome A, the ozone hole alone can result in enrichments in δ15N(NO3-) by 31.2 %, 30.7 % and 26.5 %, respectively. These values are higher than that (~6.9 %) at the South Pole, and since the effects of the ozone hole were gradually increased given the enhanced level of depletion from ~ 1976 to the mid-1990s, gradual increase in δ15N(NO3-) might be possibly detected as long as snow accumulation rate at these sites stayed relatively constant before and in the period of the ozone hole. However, at east Antarctic Plateau sites (i.e., Vostok, Dome C and Dome A) where snow accumulation rates are extremely low, δ15N(NO3-) of preserved nitrate is above 300 %. It would be difficult to determine changes of ~ 30 % out of more than 300 %, especially considering the increasing pattern of snow accumulation rate in the East Antarctic Plateau since the ~1970s (Thomas et al., 2017) and the fact that ice-core δ15N(NO3-) is very sensitive to snow accumulation rate (Akers et al., 2022). Nevertheless, a recent study by Shi et al. (2022) reported firm core nitrate concentration and isotopes at Dome A, where nitrate concentrations in the 1990s and after decreased by close to one third compared to that in the 1970s (i.e., ~18 ng g⁻¹ to ~ 12 ng g⁻¹), with 20 to 30 % increases in δ15N(NO3-) and > 5 % decreases in Δ17O(NO3-). These appear to be qualitatively consistent with the effects of the ozone hole. However quick analyses indicate that changes in nitrate mass do not agree with the degrees of isotope changes resulted from the photo-driven post-depositional processing. For example, our preliminary calculations using the TRANSITs model suggest that at Dome A the ozone hole can only induce maximum 2.8 ng g⁻¹ decreases in nitrate concentration, and as little as 0.9 % decreases in Δ17O(NO3-). These results imply there are probably other factors regulating the observed nitrate concentration and isotopes at Dome A. Note Shi et al. (2022) also did TRANSITs modeling study, but the model parameters are not clear, e.g., snow e-folding depth, quantum yield of snow nitrate photolysis, and the modeled results can’t be reproduced given local Dome A conditions we complied. A comprehensive modeling effort in
combination with more thoughtful analyses on the observed data are necessary to investigate whether the signals can be detected at Dome A.”

Table 1. Model calculated maximum isotope changes resulted from the Antarctic ozone hole (i.e., the differences between isotopes in the pre-ozone hole period and that in years with the most depletion) at different Antarctic sites.

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<th>$\Delta(\delta^{17}\text{O}(\text{NO}_3^-))$</th>
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<td>106.84</td>
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</tr>
<tr>
<td>Dome Fuji</td>
<td>-77.32</td>
<td>39.7</td>
<td>28.8</td>
<td>16.3</td>
<td>-1.2</td>
</tr>
<tr>
<td>WAIS Divide</td>
<td>-79.48</td>
<td>-112.09</td>
<td>200</td>
<td>1.0</td>
<td>-0.3</td>
</tr>
</tbody>
</table>

Another recently published study (July 2022) on nitrate isotopes in relatively high accumulation rate sites (Summit Greenland) also highlights the importance of understanding post-depositional effects of ice core nitrate and it would be worth citing this paper.


Response: Thanks for this suggestion. We have added Jiang et al. (2022) as references in the Introduction in the revised manuscript.

There are extremely scarce measurements of e-folding depth in Antarctica. A much shallower e-folding depth of 2-5 cm was observed at DML. This was also shallower than estimated by Zatko et al. (2013). What is the uncertainty on your estimated e-folding depth of 20 cm? How appropriate is that estimate in the context of measurements and modelled estimates? Given that recent studies have shown the importance of e-folding depth on nitrate recycling, a discussion and sensitivity analysis of a range of possible e-folding depths for South Pole site is highly encouraged.

Response: Thanks for these suggestions. The snow e-folding depth is determined by snow chemical and physical properties. At the South Pole, snow black carbon (BC) concentration measured by Casey et al. (2017) at South Pole is $(0.26 \pm 0.13) \text{ ng g}^{-1} (1\sigma)$, the uncertainty on model estimated e-folding depth is about $\pm 5 \text{ cm} (1\sigma)$. Due to the lack of measurement of specific surface area (SSA) and impurities of snow grains at the South Pole, we assumed total LAIs is $\sim 10$ times of BC and applied the average vertical SSA profile measured at Dome C by Gallet et al. (2011) when calculating e-folding depth at the South Pole.

In addition, we agree that e-folding depth are important to the modeled isotopes. However, here we are focusing on the long-term trends of the isotopes, over the studied period we don't expect e-folding depth have decreased or increased significantly (but of course with annual variabilities). As discussed,
e-folding depth is determined by snow LAIs, and snow physical properties (e.g., density, grain size). Over the studied period, snow LAIs are relatively constant as indicated by South Pole ice core records (as mentioned in the original manuscript). Regarding snow physical properties, the grain size is inversely proportional to specific surface area (SSA) (Zatko et al., 2013), and both snow density and SSA are affected by wind speed and temperature (Kaspers et al., 2004; Domine et al., 2009). Although surface temperature at the South Pole were not changed significantly after the 1950s (below Figure S2), surface wind speed (below Figure S3) displays a decreasing trend since ~1970. In Polar Regions the wind action will increase the SSA of surface snow, however, it is the wind storm (>55km/h) that increase the SSA in Antarctica (Domine et al., 2009). Using the empirical relationship between surface snow density with temperature, wind speed and snow accumulation rate in Kaspers et al. (2004) and the parameters in Sugiyama et al. (2012) as follows:

\[ \rho = 305 + 0.629T + 0.150A + 13.5W \]

Where, \( \rho \) is surface snow density in kg/m\(^3\), \( T \) is the annual average surface temperature in °C, \( A \) is the accumulation rate in m w.e. a\(^1\) and \( W \) is the annual wind speed in m s\(^{-1}\) at 10 m above the surface. The calculated surface snow density from 1957 to 2005 at the South Pole is plotted in Figure S4. As shown in the figure, snow density after the 1970s (i.e. the ozone hole period) is ~20 kg/m\(^3\) lower than before that. This would lead to an increase in e-folding depth by only ~1cm, corresponding to ~1‰ changes in the preserved \( \delta^{15}\text{N(NO}_3\text{)} \). Thus, the effects of the e-folding depth can be ignored.

In the revised manuscript, we added a quick discussion on this (i.e., sensitivity evaluation) as follows:

Page 6, line 168: “... Here, we assumed these factors are constant from 1944 to 2005 and the e-folding depth was the same throughout the record for simplicity, though in this period surface wind speed at the South Pole has a decreasing trend (Fig. S3) which may have affected snow density. However, the caused effects on snow e-folding depth due to changing wind speed is only ~1 cm and the resulted difference in \( \delta^{15}\text{N(NO}_3\text{)} \) is only ~1‰ and thus can be ignored (SI)...”. Where in SI, we added the sensitivity discussion on the effects of the possible long-term changes in e-folding depth.
Figure S2 The annual mean atmospheric temperature from 1957 to 2005 at the South Pole (Surface station data of annual atmospheric temperature and wind speed in the South Pole (https://ramadda.data.bas.ac.uk/repository/entry/show/?entryid=569d53fb-9b90-47a6-b3ca-26306e696706).

Figure S3 The annual mean wind speed from 1957 to 2005 at the South Pole
Figure S4 The calculated surface snow density from 1957 to 2005 at the South Pole

Please add a section of assessing the validity of the TRANSITS model especially in regards to D17O(NO3-). The model doesn’t simulate the observed decreasing D17O(NO3-) trend from ~1976 to 2000. Why is this? How much can you take away from the simulated D17O(NO3-) results? How can you improve the model? How does the model help you understand D17O(NO3-) at South Pole. TRANSITS simulations of D17O(NO3-) would be an area where the authors can contribute new understanding to the literature.

Response: Thanks for this suggestion. But the TRANSITS model is an air-snow exchange model, it only includes snow chemistry and chemistry in the overlying atmosphere. This is saying, the model can only be used, or the best used, to investigate changes related to what occurs locally (in snow or the above). It is from the model results that we can conclude that the local processes (i.e., largely the post-depositional processing) cannot result in any long-term changes in Δ17O(NO3-), so that the observed decreasing Δ17O(NO3-) should be from other factors which are most likely related to changes in primary nitrate (the starting values of Δ17O(NO3-)). From here, we further discussed effects of source regions oxidation environment and varying transport on Δ17O(NO3-) of primary nitrate. To thoughtfully investigate and/or discern the reasons why Δ17O(NO3-) of primary nitrate has decreased since the 1970s, a chemical transport model with detailed NOx source and chemistry changes in the past are necessary. This is however out of the scope of this study but in our to-do-list.

Introducing the South Pole site in terms of the snow accumulation and also atmospheric nitrate isotopes (Walters et al., 2019) in the introduction would be helpful to put the site into context of other records given that the nitrate isotopes are sensitive to accumulation rate.

Response: Thanks for this suggestion. In the end of the introduction part, we have added the following statements: “…In this study, we have … from a south Pole ice core. At the South Pole, snow accumulation rate is relatively low (0.073 w.e. m yr⁻¹) and the effects of post-depositional processing
are well observed as reflected by the large differences between atmospheric and snow nitrate isotopes (Walters et al. 2019). …”

*It is not always clear in the discussion if the authors are talking about the results from TRANSITS or observations.*

**Response:** Thanks for your comment. In the revised manuscript we have explicitly distinguish the simulated and observed results by adding describing terms of ’observed’ or ’modeled’ when related content is discussed or mentioned.

**Specific comments**

L1 The title is misleading as nitrate isotopes at South Pole do not reflect changes stratospheric ozone changes.

**Response:** Thanks for this suggestion. We have changed the title as “On the potential fingerprint of the Antarctica ozone hole in ice core nitrate isotopes: a case study based on a South Pole ice core”.

L26 HCl and ClONO2

**Response:** This seems to be in L36, and we have added a word ‘produce’ before “HCl and ClNO2’ if this is the review meant.

L65-67 The photic zone at DML is 15 cm which is less than Dome C (Winton et al., 2020).

**Response:** Thanks for your comment. We have revised it as follows:

“… This nitrate recycling process at the air-snow interface can occur multiple times before NO3- is permanently buried below the snow photic zone which is usually 15 to 60 cm deep and below this depth more than 95 % of the radiation is attenuated (Erlbrand et al., 2015; Zatko et al., 2013; Winton et al., 2020).”

L71-73 This sentence focusses on fractionation constants on the EAP. Relevant to this study are fractionation constants in the “transition zone” characterized by snow accumulation rates typical of sites located between the EAP and coast (5–20 cm yr−1 w.e.; Erbland et al. 2015).

**Response:** No, here we meant fractionation constant associated with snow nitrate photolysis, which is only related to wavelength. Erbland et al. (2015) derived different fractionation constants at different sites, but which are “apparent fractionation constant” and basically are mixed effects from the whole post-depositional processing (photolysis is the trigger step).

L86-102 Recent studies have shown the importance of e-folding depth on nitrate recycling. This is important to mention here.

**Response:** We thank the reviewer for pointing this, but e-folding depth is essentially determined by snow chemical and physical properties which are more direct when discussing post-depositional processing.

L108-111 See the recently published paper by Shi et al. (2022)

**Response:** Thanks for this suggestion. We add the following statement in our revised text:

“… annual variations in snow accumulation rate as suggested by Ming et al. (2020). A recently study by Shi et al. (2022) measured firm core nitrate concentrations and isotopes at Dome A, Antarctica, and
there appears to be responses of nitrate concentrations and isotopes to the Antarctic ozone hole. However, the onsets of the observed changes (i.e., isotopes and column ozone) are not lined-up, the model efforts in the study is ambiguous and it is unclear whether the effects of the ozone hole can quantitatively (or even at the qualitative level) explain the observed changes. In this study, we measured and examined …”

*L131* Did you decontaminate the samples?
**Response:** Yes. As stated in lines 137 to 139, we cleaned samples surface with a bandsaw and melted samples in a clean beaker at room temperature, and the concentrated process using ion-exchange resin can also filter meltwater and further decontaminate the samples.

*L134* Suggest moving reference to Geng et al. further up in the methods section.
**Response:** Thanks for this suggestion. Revised it accordingly

*L119-136* Please add protocols for minimising contamination. Please state the sample resolution in terms of depth and age here.
**Response:** Thanks for your comment. In the original manuscript we have described the protocols for minimizing contamination. “… After cutting, the surface of each sample was cleaned with a bandsaw and the cleaned sample was melted in a clean beaker at room temperature. The nitrate in the meltwater was then concentrated using ion-exchange resin…” We add the following statement in the Sect.2.1 in our revised text for the sample resolution in terms of depth and age: “…As a result, a total of 62 samples were cut from the top 8.4 meter of the SP04C6 core covering the years from 1944 to 2005. The depth resolution of these samples varies from 11 to 38 cm and each sample covering 1 year. Among these samples, …”

*L133 UW*
**Response:** Corrected as suggested

*L138-140* This sentence seems out of place.
**Response:** Thanks for this suggestion. The purpose of this sentence is to show that the ozone hole does cause a significant increase in surface UV radiation. This would further enhance the photo-driven post-depositional processing of snow nitrate and can assess through TRANSITs model. We add the following statement in revised text:
“…As shown in Fig. 1, compared to years without an ozone hole (represented by the case in 1976), in years with an ozone hole (represented by the year of 1993), surface actinic flux was significantly enhanced in the summer half year especially in spring when the ozone hole was developed. The stronger surface actinic flux in the ozone hole period presumably would enhance the photo-driven post-depositional processing…”

*L164 How did you calculate the e-folding depth?*
**Response:** Thanks for your comment. We used the two-stream Analytical Radiative Transfer in Snow (TARTES) model (Libois et al., 2013) to calculate the depth profile of actinic flux at different wavelength. We have described this in revised manuscript. We have revised it as follows:
“As a result, the e-folding depth of actinic flux at 305 nm was calculated to be 20 cm at the South Pole
using the Two-stream Analytical Radiative TransfEr in Snow (TARTES) model (Libois et al., 2013), shallower than…”

L223-226 Seems out of place.
Response: Thank you for raising this issue. The observed total column ozone and surface ozone should not be put in Sect.3.1 describing ice-core observations. We have moved these two sentences describing spring TCO and surface ozone trends to Sect.2.2 in revised manuscript where ozone data were first mentioned.

L234 Add the dates of the pit
Response: Thanks for pointing out this. We have added the snowpit date in revised manuscript:
“…All of these ice core results are however lower than \(\omega(\text{NO}_3^-)\) of (\(\sim100 \text{ – 200}\)) \(\text{ng g}^{-1}\) in a 6-m snowpit (1977-2003) at the same site reported by McCabe et al. (2007).”

L282 Can you use the approach of Weller et al. (2004) to calculate nitrate loss? And then compare to the TRANSITS estimate of nitrate loss?
Response: Thanks for your comment. Weller et al. (2004) quantified the total loss of NO3- by comparing snow nitrate concentration of the first year to the 100-year mean concentrations retrieved from the firn core. They used the first-year snow concentration to represent the surface snow concentration. Unfortunately, our ice core was drilled in the winter season of 2004/2005, the first few center meter sample represents the first couple of months in 2005. This means we don't really have the “first year” data/ We want to note, this is not a good approach as even the first year data is available it is already affected by post-depositional processing and not equal to primary nitrate. Weighted monthly average surface snow concentration could be better as the starting point.

L295 Heading should reflect that this section is about TRANSITS modelling
Response: Thanks for your suggestion. We changed the heading into: “4.2 Modeled effects of the ozone hole on the \(\delta^{15}\text{N(NO}_3^-)\) and \(\Delta^{17}\text{O(NO}_3^-)\) records”

L334 This assumption ignores other factors that influence e-folding depth. While we don’t know how e-folding depth changes over time, based on changes in grain size, snow density and impurity content it is fair to assume e-folding depth at any site is not constant through time. Sensitivity studies show that nitrate isotopes are sensitive to changes in e-folding depth.
Response: Thanks for this comment. Ice-core records shows that in the past 50 to 100 years at the South Pole impurity (i.e., LAI) are relative constant (no decreasing or increasing trend). Regarding the physical peripiteries, we have assessed them in earlier response and their long-term effects on e-folding and the consequences on \(\delta^{15}\text{N(NO}_3^-)\) are negligible. In the revised manuscript, we have explicitly discussed this in supplemental materials.

L346-357 Update in light of the published work by Shi et al. (2022).
Response: Thanks for your suggestion. We have added a subsection to discuss other sites including Dome A.
“4.3 Estimated effects of the ozone hole on snow nitrate isotopes in other Antarctic sites
In order to search for signals of the ozone hole, we used the TRANSITS model to further explore the
maximum possible responses of ice-core preserved $\delta^{15}$N(NO$_3^-$) and $\Delta^{17}$O(NO$_3^-$) to the ozone hole at other Antarctic sites, including Dome A, Dome C, Vostok, Dome Fuji, and the West Antarctic Ice Sheet (WAIS) Divide in addition to the South Pole. The responses are defined as the differences between the isotopes before the ozone hole period and those in years with the most depletion. The results are listed in Table 1. As shown in the table, except WAIS Divide, other sites all display bigger responses to the ozone hole, especially the three East Antarctic Plateau sites. These patterns are mainly determined by the differences in snow accumulations rates at these sites (Frezzotti et al., 2013; Erbland et al., 2015; Shi et al., 2022), i.e., lower snow accumulation rates correspond to longer durations of nitrate in the photic zone, leading to larger effects of the ozone hole. In particular, at Vostok, Dome C and Dome A, the ozone hole alone can result in enrichments in $\delta^{15}$N(NO$_3^-$) by 31.2 ‰, 30.7 ‰ and 26.5 ‰, respectively. These values are higher than that (~6.9 ‰) at the South Pole, and since the effects of the ozone hole were gradually increased given the enhanced level of depletion from ~ 1976 to the mid-1990s, gradual increase in $\delta^{15}$N(NO$_3^-$) might be possibly detected as long as snow accumulation rate at these sites stayed relatively constant before and in the period of the ozone hole. However, at east Antarctic Plateau sites (i.e., Vostok, Dome C and Dome A) where snow accumulation rates are extremely low, $\delta^{15}$N(NO$_3^-$) of preserved nitrate is above 300 ‰. It would be difficult to determine changes of ~ 30 ‰ out of more than 300 ‰, especially considering the increasing pattern of snow accumulation rate in the East Antarctic Plateau since the ~1970s (Thomas et al., 2017) and the fact that ice-core $\delta^{15}$N(NO$_3^-$) is very sensitive to snow accumulation rate (Akers et al., 2022). Nevertheless, a recent study by Shi et al. (2022) reported firm core nitrate concentration and isotopes at Dome A, where nitrate concentrations in the 1990s and after decreased by close to one third compared to that in the 1970s (i.e., ~18 ng g$^{-1}$ to ~12 ng g$^{-1}$), with 20 to 30 ‰ increases in $\delta^{15}$N(NO$_3^-$) and > 5 ‰ decreases in $\Delta^{17}$O(NO$_3^-$). These appear to be qualitatively consistent with the effects of the ozone hole. However, quick analyses indicate that changes in nitrate mass do not agree with the degrees of isotope changes resulted from the photo-driven post-depositional processing. For example, our preliminary calculations using the TRANSIT model suggest that at Dome A the ozone hole can only induce maximum 2.8 ng g$^{-1}$ decreases in nitrate concentration, and as little as 0.9 ‰ decreases in $\Delta^{17}$O(NO$_3^-$). These results imply there are probably other factors regulating the observed nitrate concentration and isotopes at Dome A. Note Shi et al. (2022) also did TRANSITs modeling study, but the model parameters are not clear, e.g., snow e-folding depth, quantum yield of snow nitrate photolysis, and the modeled results can’t be reproduced given local Dome A conditions we complied. A comprehensive modeling effort in combination with more thoughtful analyses on the observed data are necessary to investigate whether the signals can be detected at Dome A.”

Table 1. Model calculated maximum isotope changes resulted from the Antarctic ozone hole (i.e., the differences between isotopes in the pre-ozone hole period and that in years with the most depletion) at different Antarctic sites.

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</table>
The ice core data in the figures suggest interannual variability.

Response: Thanks for your suggestion. In this section we discussed the long terms. The revised text is shown as follows:

“…No apparent long-term trends in ice-core $\omega$(NO$_3$) and $\delta^{15}$N(NO$_3$) likely reflect that main nitrate sources to the South Pole and post depositional effects have not changed in the studied period…”

A concluding sentence about oxidation for this paragraph would be helpful here.

Response: Thanks for your suggestion. According to the suggestions of Referee # 1, we have changed the orders of the subsection in the revised manuscript, and in this part, we have added a conclusion sentence as follows:

“…Therefore, the observed $\Delta^{17}$O(NO$_3$) decrease after the 1970s is more likely due to the potential decreases in O$_3$/HO$_x$ ratio in the extratropical Southern Hemisphere. This remains to be explored and confirmed with future studies”

The EAST ANTARCTIC PLATEAU snow sourced

Response: Corrected as suggested.

Figures: It would be very helpful for the reader to visualise the TCO and nitrate isotope trends on the same figure.

Response: Thanks for your suggestion. In the revised manuscript, we have merged Figure 3 and Figure 4 and plotted a new Figure 3. Grey shading area correspond to the first stage of the ozone hole period (from 1976 to 1996).
Figure 3. Left panels: time series of annual snow accumulation rate (a), spring (average from September 22 to October 13) TCO (total column ozone) (b), and summer half year surface $O_3$ concentrations (c) at the South Pole over the period of the ice core record. Red curves are the 5-year moving averages. Right panels: ice core nitrate concentration and isotopic compositions at the South Pole in 1944-2005 (black: observations; red: modeled). The thin lines represent the observed and modeled annual (d) $\omega(\text{NO}_3^-)$, (e) $\delta^{15}N(\text{NO}_3^-)$ and (f) $\Delta^{17}O(\text{NO}_3^-)$ from 1944-2005. The thick lines represent the 5-year moving averages. Yellow shading area represents the period with changes in nitrate concentrations and isotopes from surface snow to below the photic zone. Grey shading area represents the ozone hole period.

Fig. 5: please add in the nitrate isotope observations.

Response: Thanks for your suggestion. We have added the ice-core observed nitrate isotopes in new Figure 4.
Figure 4. Sensitivity results of the modeled isotopes, i.e., $\delta^{15}$N(NO$_3^-$) (a) and $\Delta^{17}$O(NO$_3^-$) (b), to TCO and snow accumulation rate. Grey curve: ice core observed record; Red curve: modeled results with observed accumulation rate and TCO; Green curve: modeled results with observed TCO but mean accumulation rate throughout the record; Blue curve: modeled results with observed accumulation rate but TCO were kept the same before and after 1976. Grey shading area represents the ozone hole period.
Reference