Responses to Referee 1

Review of "Diurnal, seasonal, and interannual variations in $\delta(180)$ of atmospheric O2 and its application to evaluate changes in oxygen, carbon, and water cycles" by Ishidoya et al., 2024

General:

The manuscript presents a study of highest relevance for the linkages among the carbon, oxygen and water cycles. For the first time, a research group has analysed long-term measurements of the atmospheric oxygen isotope ratio (¹⁸O/¹⁶O) intending to see the impact of natural (biospheric fluxes) and anthropogenic (fossil fuel combustion). It is a fascinating manuscript to read and I would like to congratulate them for their long-term measurement effort as well as their in-depth analyses.

The manuscript is nicely written and organized and can be considered for publication with minor changes outlined below.

Thank you very much for your significant and useful comments on the paper "Diurnal, seasonal, and interannual variations in δ (¹⁸O) of atmospheric O₂ and its application to evaluate natural/anthropogenic changes in oxygen, carbon, and water cycles" by Ishidoya et al. We have revised the manuscript, considering your comments and suggestions. Details of our revision are as follows. The line numbers denote those of the revised manuscript.

Major points:

(1) As already estimated in former studies, the expected signal in the atmospheric δ (¹⁸O) by the above mentioned isotope fluxes is minimal. This requires high precision measurements over an extended period in order to see such a small signal as the single uncertainty is about 10 times larger. The required precision and accuracy, especially important for trend analysis, can only be achieved by averaging over many measurements as shown in the manuscript.

I am not sure whether the mass spectrometer was dedicated only to these measurements and whether it was run day and night autonomously. Controlling the long-term stability only every month is quite rare because the mass spectrometers behavior can change suddenly due to maintenance (filament change, ion source tuning etc). Can you please comment on these points and maybe add an additional short section about it.

Lines 99-109: Following sentences have been added considering your comments.

"In general, mass spectrometers behavior can change suddenly due to maintenance, such as filament change, ion source tuning etc.. To minimize the uncertainties associated with the changes in the conditions of the mass spectrometer, we used the specific filaments for the measurements of air samples with the atmospheric level amount fraction of O₂ supplied by the Thermo Scientific co.. This enabled us to carry out the continuous measurements in the present study for 11 months without exchanging the filament (when we used the original filament supplied for the mass spectrometer, then we needed to exchange it every 3 months). After the exchange of the filament, several weeks are needed to stabilize the condition of the ion source of the mass spectrometer by flowing the sample and reference air, especially for the elemental ratios as O_2/N_2 , Ar/N_2 , and CO_2/N_2 . Once the condition is stabilized, we did not tune the ion source throughout the period using the same filament. Furthermore, the mass spectrometer was dedicated only to the measurements of δ_{atm} (¹⁸O) and related components including those for flask samples (e.g. Ishidoya et al., 2021, 2022) and it was run day and night autonomously to keep the condition of the ion source."

(2) Regarding the long-term estimated change in $\delta^{18}O(O_2)$, the authors assume a constant increase rate of the GPP per year. For me, this does not make sense as there are many studies out there discussing the CO₂ fertilization effect. Therefore, I would rather assume a scaling based on the excess CO₂ level (CO₂actual – CO₂ preindustrial). Even though this will most probably not affect their results significantly (Fig. 8), but it still better than use a constant increase rate. In particular as the authors have used a CO₂-dependent photorespiration rate (eq. 5).

Lines 461-469 and Fig. 8: We have assumed a scaling of GPP based on the CO₂ amount fractions from Scripps CO₂ Program, considering your suggestion. Specifically, following sentences have been added and the simulated $\delta_{\text{atm}}(^{18}\text{O})$ plotted in Fig. 8 has been revised.

"We first assumed that the global terrestrial GPP increases in proportion to the global average CO₂ amount fraction. As the global average CO₂ amount fraction, we used the data from Scripps CO₂ Program (https://scrippsco2.ucsd.edu/data/atmospheric_co2/icecore_merged_products.html) based on ice-core data and direct observations before and after 1959, respectively (Keeling et al., 2001; Rubino et al., 2019). We assume the initial terrestrial production of O₂ in 1871 as 16.7 Pmol a⁻¹ (Table 1), which corresponds to 107 Pg a⁻¹ (C equivalents) of global terrestrial GPP considering the r_{DR} of 0.59 and the OR_B of 1.1. Then, the GPP increased secularly with increasing CO₂ amount fraction, and it takes 141 Pg a⁻¹ (C equivalents) in 2006. Although this is somewhat larger than the average GPP of 125 Pg a⁻¹ (C equivalents) for the period 1992–2020 reported by Bi et al. (2022), it falls within a range of the global GPP estimates from various models summarized in Fig. 10 of Zheng et al. (2020)."

Minor points:

Title:

Diurnal, seasonal, and interannual variations in $\delta(^{18}O)$ of atmospheric O_2 and its application to evaluate changes in oxygen, carbon, and water cycles.

What kind of changes do you mean here? Changes in trend, seasonality, fossil fuel influence, natural changes?

Title: The title has been changed to "Diurnal, seasonal, and interannual variations in δ ⁽¹⁸O) of atmospheric O₂ and its application to evaluate natural/anthropogenic changes in oxygen, carbon, and water cycles".

L15-16

the amplitude is very small, how about its uncertainty? Because it is a mean of about 11 seasons, the seasonality could be smoothed.

Lines 387-390: The uncertainties for the seasonal amplitudes of $\delta_{\text{atm}}(^{18}\text{O})$ and $\delta(\text{O}_2/\text{N}_2)$ have been added as "The minimum of the seasonal $\delta_{\text{atm}}(^{18}\text{O})$ cycle appeared in late summer to early autumn, and the peak-to-peak amplitude was 2.1±0.6 per meg. The maximum of the seasonal $\delta(\text{O}_2/\text{N}_2)$ cycle occurred in summer, and its peak-to-peak amplitude was 113±10 per meg. The uncertainties for the amplitudes of $\delta_{\text{atm}}(^{18}\text{O})$ ($\delta(\text{O}_2/\text{N}_2)$) was evaluated as a standard deviation of the 10-year average monthly mean values from the best-fit curve shown in Fig. 7b.". In the abstract, we did not state the uncertainty since we use the phrase "about 2 per meg".

L18

The secular increase is even more delicate to determine, which requires an extreme stability of the instrument and the standard gas measurements. What about the influence of filament changes, power interruptions, ion source tunings, inlet system, gas flow regime. I am amazed about the stability that is required. Isotope ratio may be less prone to changes but elemental ratios as O_2/N_2 or Ar/N_2 are generally more dependent on such changes. Can you comment on these. Thank you.

As our reply to your "major points (1)", we have added related sentences to lines 99-109. We note that the gas introduction methods (Lines 80-86) are primarily important to obtain the stability (details of the system can be found in Ishidoya and Murayama (2014)).

L26-28

consider rewording to:

The ¹⁸O/¹⁶O ratio of atmospheric O₂, δ_{atm} (¹⁸O), is about 24 ‰ higher than that of ocean water (per definition 0 ‰ on the Vienna-Standard Mean Ocean Water (V-SMOW)) due to various processes in

the global oxygen and water cycle (e.g. Craig, 1961; Barkan and Luz, 2005) Lines 28-30: The sentence has been rewritten, as suggested.

L35-36

Please give corresponding references. There are many more than given here. Lines 39-40: Some corresponding references have been added, as suggested.

L37

.. of present air, ...

Line 41: The words "of air at present" have been changed to "of present air".

L37

....and that variations of the DME from the average are ± 0.2 ‰. This addition is not clear, please be more specific here.

Lines 41-44: The sentences have been rewritten as follows, considering your comments.

"B94 have reported that the DME is on average lower by 0.05 ‰ than that of present air during the past 130,000 years, and the standard deviation of the DME from the average was ± 0.2 ‰. They suggested that the DME was nearly unchanged between glacial maxima and interglacial periods, and the variability is small and may be due to variations of the relative rates of primary production on the land and in the ocean."

L45

Which value is now used? You may write...and have obtained a range of 22.4 to 23.3 for DME. Lines 50-51: The words "...and have reproduced the average DME of 22.4 or 23.3 ‰" have been changed to "...and have obtained the average DME of 22.4 to 23.3 ‰d", as suggested.

L63-65

This is a very interesting statement. Thank you very much for your evaluation.

L82-83

Switch sentence structure, 2nd part first and vice versa.

For the continuous measurements of stable isotopic ratios of O₂, N₂, and Ar ($\delta_{atm}(^{18}O)$, $\delta_{atm}(^{15}N)$, and $\delta_{atm}(^{40}Ar)$) as well as the O₂/N₂ ratio and amount fraction of CO₂, we repeatedly conducted alternate analyses of the sample and reference air.

Lines 88-89: The sentence has been modified following your suggestion as "We repeatedly conducted alternate analyses of the sample and reference air, for the continuous measurements of stable isotopic ratios of O₂, N₂, and Ar ($\delta_{\text{atm}}(^{18}\text{O})$, $\delta_{\text{atm}}(^{15}\text{N})$, and $\delta_{\text{atm}}(^{40}\text{Ar})$) as well as the O₂/N₂ ratio and amount fraction of CO₂".

L84

how come to determine a trend of 0.22 per meg or a seasonality of 2 per meg with a standard deviation of 20 per meg. This requires an well-defined long-term stability.

As described in the text (Lines 119-123), we evaluated the long-term stability from $\delta_{\text{atm}}(^{18}\text{O})$ of three secondary standards against the primary standard air. Variations of the annual average $\delta_{\text{atm}}(^{18}\text{O})$ of our three secondary standards were ± 0.9 per meg on average (Fig. 2), which corresponds to uncertainty of ± 0.13 per meg a⁻¹ for the 10-year-long secular trend. This enables us to detect the average trend of 0.22 per meg per meg a⁻¹ for the period 2013-2022. On the other hand, variations of each value of $\delta_{\text{atm}}(^{18}\text{O})$ of our three secondary standards were ± 2.3 per meg on average (Fig. 2). Therefore, it is difficult to detect a seasonality of 2 per meg of $\delta_{\text{atm}}(^{18}\text{O})$ from 1-year observation only, but we can evaluate it as 10-year average seasonal cycle ($\pm 2.3 / \text{sqrt}(10) = \pm 0.7$ per meg, which is consistent with the 2.1 ± 0.6 per meg evaluated as a standard deviation of the 10-year average monthly mean values from the best-fit curve shown in Fig. 7b).

L86

...calculated by ...Keeling...

Line 92: The words " $\delta_{atm}(^{18}O)$ by Keeling..." have been changed to " $\delta_{atm}(^{18}O)$ calculated by Keeling...", as suggested.

L88-89

For this purpose, the measured values of the $\delta_{\text{atm}}(^{18}\text{O})$ for the same air sample needed to not show any temporal drift, at least during the averaging period.

not clear what you want to say here, maybe you combine it with the previous sentence to

This averaging results theoretically in a standard error of the observed $\delta_{\text{atm}}(^{18}\text{O})$ of less than 0.6 per meg assuming no temporal drift during the averaging period.

Lines 93-94: The sentences have been combined and rewritten as you suggested.

L106

...an uncertainty of ± 0.13 per meg a⁻¹...how was this calculated?

Line 122: The uncertainty was calculated by $\pm \sqrt{(0.9)^2 + (0.9)^2}/10$ considering the long-term stability (1 standard deviation) of the annual average of $\delta_{\text{atm}}(^{18}\text{O})$ of three secondary standards (blue circles shown in Fig. 2).

L113-114

As seen in Figure 3a, $\delta_{\text{atm}}(^{18}\text{O})$ increased linearly with increasing amount fractions of CO₂.

Why, what are the reasons? There is no isobaric interference. Has it to do with isotope exchange between CO_2 and O_2 ? Have you done CO_2 additions with O_2 labelling?

Lines 132-135: Unfortunately, the mechanism has not been clarified yet, so that we have added following sentences.

"The mechanism of the positive correlation between the $\delta_{\text{atm}}(^{18}\text{O})$ and CO₂ was not clarified yet since there is no isobaric interference. In this regard, I found no significant influences of CO₂ amount fraction on $\delta_{\text{atm}}(^{18}\text{O})$ for a different mass spectrometer, Finnigan MAT-252 (Ishidoya, 2003). This suggest that the influences should be examined carefully for each mass spectrometer."

We agree with you that CO_2 additions with O_2 labelling will be a useful method to evaluate the possibility of isotope exchange between CO_2 and O_2 . We would like to leave it as a future task.

L120-121

reword toin our earlier flask studies in 2013.

Line 141: The words "in our earlier experiments in 2013 that involved use of flasks" have been changed to "in our earlier flask studies in 2013", as suggested.

L135-136

 R_{TS} and R_{ST} denote the ratios of the annual fluxes of O_2 between the troposphere and stratosphere, respectively.

to what? It is a ratio.

Lines 168-170: These are the ratios to the total amount of O_2 in the atmosphere. We have revised the sentence as "R_{Res}, R_{PS}, R_{OR}, and R_{OP} represent the relative ratios of the annual amounts of O_2 from terrestrial respiration, marine respiration, terrestrial production, and marine production, respectively, to the total amount of O_2 in the atmosphere (=3.706 x 10⁴ Pmol)."

L140

... the amount fraction of O_2 calculated by the box model was converted to $\partial (O_2/N_2)$.

how? Assuming a norm atmosphere or using the measurements to do it correctly. I ask this because of dilution effects.

Line 183: In the box model, the amount fraction of O2 was calculated assuming a norm atmosphere.

L152-153

Here, ε_{ST} was set to -4 per meg so that the diminution of δ_{atm} ⁽¹⁸O) at equilibrium was -0.4 ‰.

How come?

Lines 195-198, Table 1: In this revision, we have slightly changed the value of the stratospheric diminution, taking L&B2011 results into account. We have added Table 1 to show budgets (fluxes) and isotopic effects of atmospheric O₂ used in the box model. As shown in the table, stratosphere – troposphere exchange ($R_{ST} = 3.0 \times 10^3$ Pmol a⁻¹) is about 100 times larger than the total biospheric flux at the surface (16.7 + 9.8 Pmol a⁻¹). Therefore, ε_{ST} was set to 2.5 per meg to contribute to DME by about –0.3 per mil; –0.3 = $R_{ST} \times \varepsilon_{ST} / (R_{PS}+R_{OP})$.

L161-162

This uncertainty complicates the problem of inter-annual $\delta_{\text{atm}}(^{18}\text{O})$ change and suggests that gravitational separation may be involved in small fluctuations in the DME.

One needs to look into O_3 and ¹⁴C variations at high altitudes, ideally close to the tropopause.

We agree with you associated with the importance of S-T exchange through the tropopause for isotopically light O_2 in the stratosphere, but in this paragraph, we discuss gravitational separation which is not necessarily reflect STE alone, but is also influenced by the balance between molecular and eddy diffusion and/or strength of Brewer-Dobson circulation (e.g. Ishidoya et al., 2021). Therefore, we leave the sentence as it is.

L204-205

...and artificial inlet fractionation induced by radiative heating of an air intake (e.g., 205 Blaine et al., 2006).

You mentioned that thermal diffusion is not affecting the measurements due to the high flow rate. Line 268: The words "...diurnal $\delta(Ar/N_2)$ cycle, which is driven by" have been changed to "...diurnal $\delta(Ar/N_2)$ cycle, which is potentially driven by", to clarify both the night-time vertical temperature gradient and artificial inlet fractionation are just the possible causes.

L234

1.46in graph 1.45

Line 299: The number was corrected to 1.45. Thank you for pointing it out.

L239

why only to terrestrial and not to marine biosphere activities?

 O_2 and CO_2 fluxes between the terrestrial biosphere and the atmosphere are tightly correlated with each other (with ER ~ 1.1), while those between the ocean and the atmosphere are not due to carbonate dissociation effect. Therefore, we consider the summertime diurnal $\delta(O_2/N_2)$ with the ER of 1.08 could be attributed mainly to terrestrial biosphere activities.

L243-244

(https://www.enecho.meti.go.jp/statistics/energy_consumption/ec002/results.html#headline2, last access: 28 March 2024, in Japanese) (Ishidoya et al., 2020).

paper in 2020, reference in 2024?

We found the past URL in Ishidoya et al. (2020) was not convenience of the readers to find specific data, so that we have shown the updated URL accessed recently on 28 March 2024.

L244-246

The implication is therefore that the isotopic discrimination of O_2 during activities of the terrestrial biosphere was the main cause of the observed summertime diurnal $\delta_{\text{atm}}(^{18}\text{O})$ and $\delta(O_2/N_2)$ cycles, and the isotopic discrimination of O_2 during fossil fuel combustion was very small or negligible.

The same conclusion could be drawn by radiocarbon measurements. I guess ¹⁴C measurements are being done at your station. Why not use and show it?

We agree with your suggestion, but unfortunately, our institute has not observed Δ^{14} C of CO₂. I guess National Institute for Environmental Studies (NIES) observed Δ^{14} C at TKB, so that the results shown in the present study will be a useful tool in future to validate δ_{atm} (¹⁸O) and Δ^{14} C methods with each other.

L254-256

This method, hereafter referred to as the " $\delta_{\text{atm}}(^{18}\text{O})$ -method", enabled us to remove the impact on $\delta(O_2/N_2)$ of not only the activities of the terrestrial biosphere but also the contributions due to the air–sea O_2 flux, which is driven mainly by activities in the marine biosphere (e.g., Nevison et al., 2012; Eddebbar et al., 2017), from the estimated $\delta_{\text{FF}}(O_2/N_2)$.

Not clear as you first make the balance between observed and bio to obtain the FF. By doing this you cannot disentangle the air-sea O₂ flux from the terrestrial O₂ flux.

We did not separate (disentangle) the air-sea O₂ flux from the terrestrial O₂ flux in this case. Instead, we separate the contribution of "the air-sea O₂ flux + the terrestrial O₂ flux" from that of fossil fuel combustion. This is based on (1) the simulated diurnal cycle of $\delta_{atm}(^{18}O)$ and the $\delta_{atm}(^{18}O) / \delta(O_2/N_2)$ ratio for the case considering terrestrial processes only were very similar to those for the case considering marine processes only, for both case the isotopic effects from B94 and L&B11 were incorporated into the box model (lines 312-315), (2) the contributions due to the air–sea O₂ flux is considered to be driven mainly by activities in the marine biosphere (e.g., Nevison et al., 2012; Eddebbar et al., 2017), and (3) the isotopic discrimination of O₂ during fossil fuel combustion was very small or negligible (Fig. 4). Then, we can estimate the variations of the observed $\delta(O_2/N_2)$ driven by the total activities of the terrestrial and marine biosphere (" $\delta_{BIO}(O_2/N_2)$ ") by dividing the observed variations in $\delta_{atm}(^{18}O)$ ($\delta_{atm}(^{18}O)$ is driven by the total activities of the terrestrial and marine biosphere).

L267-268

It is noteworthy that propane ($CH_3CH_2CH_3$), for which the OR_{FF} is 1.67 for complete combustion, should also be considered as the household gas consumed in the TKB area.

This is very interesting. Thank you very much for your interest.

L278-281

y. Similar separation has been carried out for CO₂ based on the simultaneous analysis of the Δ (¹⁴C) and amount fraction of CO₂ (e.g., Basu et al., 2020) or based on the simultaneous analysis of δ (O₂/N₂) and the amount fraction of CO₂ by assuming an average OR_{FF} based on a statistical assessment (Pickers et al., 2022).

There are more publications available that might be cited! Lines 362-364: We have added some references to be cited.

L291-292

Figure 7a therefore shows 116 and 120 δ_{atm} ⁽¹⁸O) and δ (O₂/N₂) data, respectively.

rewrite

Lines 374-375: The sentence has been rewritten as "Therefore, Fig. 7a shows 116 and 120 data of $\delta_{\text{atm}}(^{18}\text{O})$ and $\delta(\text{O}_2/\text{N}_2)$, respectively."

L306-309

Keeling (1995) expected $\delta_{\text{atm}}(^{18}\text{O})$ to be lower in summer than in winter by 2 per meg based on the assumption that the 100 per meg seasonal increase of $\delta(O_2/N_2)$ was driven by the input of photosynthetic O_2 , the $\delta(^{18}\text{O})$ of which is about 20 ‰ lower than $\delta_{\text{atm}}(^{18}\text{O})$.

Show how to calculate it!

Lines 393-398: The calculation method has been added, as suggested.

L316

We found that the box model could reproduce the observed seasonal δ_{atm} ⁽¹⁸O) cycles

You adjusted the corresponding values. Questions are remaining as to whether the used model values fall within known ranges.

Table 1: We have added Table 1 to clarify the specific values and references for the parameters incorporated into the box model, considering your comments.

L355-357

see major point 2

Fig. 2

The measurements are not equally distributed over time, this influences the uncertainty per year. Have you considered this?

We did not consider an effect of the non-uniform distribution you pointed out. I agree this could influence the uncertainty for secular trend, nevertheless the effect is not so serious since the measurements in 2012-2014 (the first period) and those after 2020 (the last period) is relatively denser than those in 2014-2020 as seen from Fig. 2. The uncertainty of the average secular trend throughout the period is determined mainly by the measurements in the first and last periods.

Responses to Referee 2

The authors present measurements of the ¹⁸O isotopic composition of atmospheric molecular oxygen from air taken at the roof of their building in the greater Tokyo region. They also measured O_2/N_2 ratios and CO_2 concentrations to assess the suitability of delta-¹⁸O in O_2 to probe the carbon cycle. They analyse the diurnal and seasonal variations as well at the interannual trend with a one-box model. The observations are pretty exciting. I am not aware of any group that measured the diurnal cycle of delta-¹⁸O in atmospheric O_2 before. Given the tremendous technical advances over the last years, it seems logical that they succeeded, eventually. They still have to average about 1000 individual measurements so that average diurnal cycles for each season are presented.

I am less convinced by the analysis with the one-box model. The method was not well explained, so that it is possible that I missed some things. I think that the manuscript needs some serious revisions before publication.

Thank you very much for your significant and useful comments on the paper "Diurnal, seasonal, and interannual variations in δ (¹⁸O) of atmospheric O₂ and its application to evaluate natural/anthropogenic changes in oxygen, carbon, and water cycles" by Ishidoya et al. We have revised the manuscript, considering your comments and suggestions. Details of our revision are as follows. The line numbers denote those of the revised manuscript.

Why are there only the values of Bender et al. (1994) but not the updated figures of Luz and Barkan (2011, doi: 10.1029/2010GB003883)?

Section 2.2 (lines 143-236) and Table 1: Considering your comment, we have carried out the simulations by using the isotope effects not only from Bender et al. (1994) but also from Luz and Barkan (2011) (referred to as "B94" and "L&B11", respectively, in the manuscript), and compared them especially for the long-term variations of δ_{atm} ⁽¹⁸O). Table 1 has also been added to summarize values of O₂ budgets and isotopic effects for both studies.

Where is fossil fuel in the box model? It looks like the model comes from Bender et al. (1994) who analysed the last 130,000 years and did not need fossil fuel.

Equations (3) and (4) and related sentences, lines 172-178, and lines 322-329: In the revised manuscript, contributions of fossil fuel combustion has been included in the equations for the box model. However, we assume that atmospheric oxygen is consumed without isotope effects in fossil fuel combustion considering high temperature during the industrial combustion processes. We have added some sentences to note future needs to examine the combustion processes as follows.

"(lines 172-178) We assumed that atmospheric oxygen is consumed without isotope effects in fossil

fuel combustion ($\mathcal{E}_{FF}=0$), taking into account that the industrial combustion processes usually occur at high temperature. Therefore, we consider no contribution to DME from fossil fuel combustion in this study. In this regard, it is known that large oxygen isotope fractionation occurs in the combustion processes such as biomass burning due to complex combustion processes (Schumacher et al., 2011). In such cases, it will be necessary to consider isotopic fractionation in the consumption of atmospheric oxygen associated with combustion. However, at present, little is known about the impact of this on DME."

"(lines 322-329) It would be generally reasonable since the combustion occurs at high temperature, which minimizes isotopic discriminations. However, Schumacher et al. (2011) reported isotopic discriminations on the order of up to 26 ‰ for stable oxygen isotopic ratio of atmospheric CO₂ ($\delta_{CO2}(^{18}O)$) derived from combustion of different kinds of material. They suggested that natural combustion processes on the long term might enrich $\delta_{atm}(^{18}O)$ and contribute to the DME. Therefore, isotopic discriminations of $\delta_{atm}(^{18}O)$ due to combustion processes should be examined carefully in future, based on precise observations of $\delta_{atm}(^{18}O)$."

The method of ignoring terrestrial or marine fluxes was not well explained. Only the terrestrial fluxes had a sinusoidal cycle in the methods. So how does the model calculate a diurnal cycle if terrestrial O₂ fluxes are ignored?

Lines 231-233: The sentence "We also carried out simulations of diurnal changes considering marine O_2 consumption and production approximated by the similar simple function, to examine sensitivities of $\delta_{atm}(^{18}O) / \delta(O_2/N_2)$ ratio to the terrestrial and marine signals" has been added to make the method clearer.

The rationale behind their calculations of the delta-¹⁸O_atm-method were not given. Why does the ratio give you delta-¹⁸O bio? Is it because fossil fuel is missing in the box model?

As you expected, it is because we ignored the isotopic effect of fossil fuel combustion. Regarding this assumption, we have added some sentences to note future needs to examine the processes (please check our reply to your comments "Where is fossil fuel in the box model?..." above).

The box model has a shifted diurnal cycle by about two hours. You would get a strange signal when dividing two sinusoidal signals shifted by some delta. This does not seem the case here and I was wondering why? Is it possible that you get a false diurnal cycle of the biospheric fluxes because of the wrong timing of the box model?

In the simulations, diurnal cycles of respiration and photosynthesis fluxes were approximated by simple sinusoidal signals. We could not know true phases and amplitudes of the diurnal cycles, so that we adjusted the amplitude of the respiration and photosynthesis fluxes arbitrarily under the constraint

that both fluxes became the largest around noon. Therefore, if we shift the peaks of the fluxes by two hours earlier, then we can reduce the phase shifts of the simulated $\delta_{\text{atm}}(^{18}\text{O})$ and $\delta(\text{O}_2/\text{N}_2)$ found in Fig. 4a. However, we consider it would be reasonable that the largest respiration and photosynthesis fluxes are found around noon, so that we leave the simulated diurnal cycles in Fig. 4a as they are.

The ER method is not explained. Where are the numbers 1.1, 1.4-1.7 coming from? I guess nowadays 1.05 is more accepted for photosynthesis.

Lines 335-338: The sentence has been modified considering your comments as "...we assumed the OR for activities in the terrestrial biosphere (OR_B) to be 1.1 (Severinghaus, 1995), which has been widely used in past studies (e.g. Manning and Keeling, 2006; Tohjima et al.), and the OR_{FF} to be 1.4, 1.5, 1.6, or 1.7 considering mixed combustion of solid fuel, liquid fuel, and natural gas. It is noted some recent studies have used the OR_B of 1.05 rather than 1.1 (e.g. Morgan et al., 2021)".

How is it possible that in O₂ leaf water isotopes are increasing by 9 per meg (Figure 8) when leaf water isotopes of H₂O are increasing by nothing until 2000 and then only about 0.2 permil? Lines 512-516: Following sentences have been added to show the reason clearer.

"It is noted that the contributions of the changes of $\delta_{LW}(^{18}O)$ to the simulated $\delta_{atm}(^{18}O)$ increased with time monotonously while clear increase of $\delta_{LW}(^{18}O)$ was found after the 1980s (Figs. 8b–c). This is due to the choice of the initial $\delta_{LW}(^{18}O)$ in 1871; we set it to be 4.4 or 6.5 ‰ (the values for steady state by B94 or L&B11). As seen from Fig. 8c, the average $\delta_{LW}(^{18}O)$ during 1872–1980 was higher than the initial values, which made the monotonous increase of the $\delta_{atm}(^{18}O)$ driven by the $\delta_{LW}(^{18}O)$ changes..

What are the leaf water scenarios in Figure 7? I could not find any explanations. I would be curious how you get a time shift of up to two months from different formulations of leaf water.

Lines 421-435: The sentences have been rewritten to explain the scenarios. The scenario for the thick dashed blue line in Fig. 7 was determined based on some past studies reported seasonal variations of $\delta_{LW}(^{18}O)$, and other two scenarios represented by two-dot chain and dotted lines were carried out as sensitivity tests to the phase difference in seasonal $\delta_{LW}(^{18}O)$ cycle.

You get an increasing or decreasing secular trend if the right-hand side of Eq. (3) is non-zero. This can have many reasons and you do not have to have increasing GPP or anything. The authors have tweaked so many fluxes in their model that I do not think that we can say anything about the secular trend. Section 3.3 (lines 441-558) and Fig. 8: We have revised the analysis and discussion for the secular trend including the simulations by the box model incorporated not only B94 but also L&B11. As you expected, the secular trends of the simulated $\delta_{\text{atm}}(^{18}\text{O})$ are highly sensitive to the isotopic effects associated with the DME. Therefore, further studies are needed to determine the isotopic effects precisely, in order to evaluate long-term changes in GPP and photorespiration based on δ_{atm} ⁽¹⁸O).

This reminds me of the literature of delta-¹⁸O in atmospheric CO₂. There were the same issues: a secular trend due to unbalanced fluxes, a one- to two-month time shift in the seasonal cycle, etc. The authors could learn at lot from that literature but not a single paper is referenced in the manuscript. Lines 408-413 and 498-502: Following sentences have been added to note the characteristics of seasonal and interannual variations in stable oxygen isotopic ratio of atmospheric CO₂ (δ_{CO2} (¹⁸O)) for a comparison with those of δ_{atm} (¹⁸O).

"(lines 408-413) In this context, seasonal cycles of $\delta_{CO2}(^{18}O)$ have been reported by some past studies (e.g. Peylin et al., 1999; Cunz et al., 2003; Murayama et al., 2010). Peylin et al. (1999) and Cunz et al. (2003) used 3-D atmospheric transport models to reproduce the observations, and they found the main contributors are respiration and production for the respective seasonal cycles of $\delta_{CO2}(^{18}O)$ and CO_2 amount fraction. These characteristics are different from the seasonal cycles of $\delta_{atm}(^{18}O)$ and $\delta(O_2/N_2)$ observed in this study, both of which are driven mainly by production (Fig. 7c)."

"(lines 498-502) It should be noted that Welp et al. (2011) suggested that $\delta_{CO2}(^{18}\text{O})$ increases with increasing $\delta_{\text{precip}}(^{18}\text{O})$ and $\delta_{LW}(^{18}\text{O})$ through the redistribution of moisture and rainfall in the tropics during an El Niño, which leads to substantial interannual variations in $\delta_{CO2}(^{18}\text{O})$ during 1977–2009 obtained from the Scripps Institution of Oceanography global flask network. Therefore, it will be important in future studies to examine not only secular trend discussed in this study but also interannual variations in $\delta_{LW}(^{18}\text{O})$ and $\delta_{atm}(^{18}\text{O})$."

Nobody thinks that GPP increase over the last century comes solely from a decrease of photorespiration. The discussion from page 14 line 397 up to page 15 line 417 is weird regarding the carbon cycle and anything we know about photosynthesis.

The discussion you pointed out in the previous manuscript has been removed from the revised manuscript.

The leaf water from MIROC5-iso looks like source water. Most global models of water isotopes do not calculate leaf water enrichment because they assume steady state so that the transpired water is the same as source water. If not, I would have loved to know how delta-¹⁸O of leaf water is calculated in MIROC5-iso.

Lines 247-254: MIROC5-iso does calculate the isotopic fractionation effect when transpiration occurs. We added the equation to make the point clearer.

More minor comments are:

Why is the ¹⁸O in parenthesis in δ ⁽¹⁸O)? This is a weird notation.

We understand your comment, however, I have used the phrase following the Editor's instruction.

Given the current notation. It is never clear which molecule is looked at. Sometimes $\delta_{LW}(^{18}O)$ is O₂ and sometimes H₂O. Perhaps making it clearer, e.g. adding the molecule behind such as $\delta^{18}O$ -O₂ or $\delta^{18}O(O_2)$?

In the revised manuscript, we have used $\delta_{LW}(^{18}O)$ for H₂O only. The words "...and the respective contributions of GPP (solid green line), photorespiration (solid red line), and $\delta_{LW}(^{18}O)$ (solid blue line) to the simulated $\delta_{atm}(^{18}O)$..." in caption of Fig. 8 has been changed to "...and the respective contributions of the changes of GPP (green line), photorespiration (red line), and $\delta_{LW}(^{18}O)$ (blue line) to the simulated $\delta_{atm}(^{18}O)$..." to make the meaning clearer.

Lots of references are missing like all the references for the emission ratio method. Or ER = 1.67 for propane?

References for the DME, emission ratio methods, and $\delta_{CO2}(^{18}O)$ have been added in the revised manuscript (e.g. lines 39-40, 154, 176, 336-338, 362-364). As for the OF_{FF} for propane, the words "...the OR_{FF} is 1.67 for complete combustion" have been changed to "...the OR_{FF} is 1.67 assuming complete combustion" (line 350). If propane completely burned, $C_3H_8 + 5O_2 \rightarrow 3CO_2 + 4H_2O$, then the OR is 5/3 = 1.67.

I was wondering if MIROC5-iso has no carbon cycle? Most models have nowadays. So why not using these fluxes, or at least its dirunal and seasonal variations, instead of simple sinusoidal fluxes? Thank you for the valuable suggestion. Unfortunately, the carbon cycle is not included in MIROC5-iso, although it is included in another version of MIROC; MIROC-ESM (Watanabe et al., 2011). We would like to leave it as a future task.

Responses to Referee 3

Jeff Severinghaus,

Ishidoya et al. have produced a stunning and groundbreaking extension of the well-known millennialscale variations in atmospheric oxygen isotopes (namely ¹⁸O of O₂), sometimes known as the Morita-Dole Effect, that are recorded in ice cores. Their extension brings to the table totally new and fascinating information - namely the first high-quality observations of diurnal and seasonal cycles in ¹⁸O of O₂. Their tour-de-force treatment of extremely difficult analytical techniques makes it possible now to ask totally new questions about the role of the terrestrial biosphere in the last 5 decades of (unplanned) anthropogenic CO₂ fertilization due to fossil fuel burning, as just one example among many.

The quality of their measurements is superb, and unparalleled. Their deep attention to details, and exploration of potential pitfalls, makes their conclusions robust and convincing.

One very minor comment I would make is that their box model estimate of ~1500 years for the turnover time of atmospheric O_2 may be a little too long. My ice core work shows that ¹⁸O of atmospheric O_2 relaxes with a characteristic asymptotic decay curve after abrupt climate change events on a timescale of about ~1000 years, implying that the turnover time of O_2 in the atmosphere is about ~1000 years.

The authors are to be congratulated for a true breakthrough that will no doubt open many doors for future study of the interlinked carbon, oxygen, and argon cycles in Earth's atmosphere. These studies will no doubt shed light on the ongoing anthropogenic perturbations to the cycles of these gases.

Thank you very much for your significant and useful comments on the paper "Diurnal, seasonal, and interannual variations in δ ⁽¹⁸O) of atmospheric O₂ and its application to evaluate natural/anthropogenic changes in oxygen, carbon, and water cycles" by Ishidoya et al. We are so happy to hear that you are pleased with the work that was done. We have revised the manuscript, considering your comments and suggestions, and those from other two reviewers. We have added following sentences, considering your comments regarding the turnover time.

"(lines 214-221) The biospheric turnover time of O_2 in the steady state was 1398 years, which is longer than the 1200 years estimated by B94. This may be a little too long, since the $\delta_{\text{atm}}(^{18}\text{O})$ variations reported by Severinghaus et al. (2009) based on the ice core measurements that showed a characteristic asymptotic decay curve after abrupt climate change events on a timescale of about ~1000 years, implying that the turnover time of O_2 in the atmosphere is about 1000 years. The biospheric turnover time is inversely proportional to the sum of the terrestrial and oceanic productions of O_2 incorporated into the box model, which is 26.5 (16.7 + 9.8) Pmol a⁻¹ in this study (Table 1). This implies that total production of O_2 for the initial value in our model is underestimated. In this regard, turnover time decreases to about 1000 years when we simulate a case in which the GPP is increased, as will be discussed later."