Reply to Review Comments

We thank the anonymous referee and Ashley Ballantyne for their thoughtful comments and for their time and effort to review this manuscript. Original review comments are given in black, our answer in red, and new or revised text added to the manuscript in blue fonts.

Anonymous Referee #1

In the manuscript “20th – century changes in carbon isotopes and water-use efficiency: Tree-ring based evaluation of the CLM4.5 and LPX-Bern models” Keller et al. present the implementation of a carbon isotope scheme in two global models as well as their performance with respect to simulated spatial patterns and decadal trends. The model results are compared to two different datasets, tree-ring records and bulk leaf delta 13C data. This study is a valuable contribution to ongoing efforts on the implementation of carbon isotopes in global vegetation models. The overall approach as well as the results are presented in an adequate and clear manner.

Thank you.

My main criticism is on the conclusions that are drawn from these results. Some parts of the discussion will be subject to revision. More detailed comments are listed below.

We will address your points below.

Abstract The abstract is a nice summary and contains all important aspects of the paper. However, I disagree with the last sentence. Suggesting “fundamental problems associated with the prescribed relationship between conductance and assimilation” is rather provocative and not supported by the results of this study. This relationship is strongly supported by observations (see e.g. Wong 1979 and also De Kauwe et al. 2013, Global Change Biology; papers cited in this study) and consequently used in most global models. I.e. it would indeed be a fundamental problem in our understanding of plant physiology. If this thought is brought up at such a prominent position in the paper, it needs to be better discussed and corroborated later in the manuscript, see below.

The sentence will be revised to read: “The results suggest that the down-regulation of ci/ca and of photosynthesis by nitrogen limitation is possibly too strong in the standard setup of CLM4.5 or there may be problems associated with the implementation of conductance, assimilation, and related adjustment processes to long-term environmental changes.”

We agree that the relationship is experimentally well-established. Yet, it remains controversial whether and how the relationship between conductance and assimilation changes under changing environmental conditions, particularly those now addressed in the model experiments such as the monotonic increase in atmospheric CO2 over the past 170 years (Miner et al., 2016).

Introduction The introduction starts with a nice overview on the application of isotopes in the Earth System, which is a good motivation for this study. Following this part, the connection between carbon isotopes and plant physiological behavior is pointed out. The introduction closes with a very clear outline of the goals of this study.

Thank you.

Methods Page 4: the listing of all PFTs seems a bit unnecessary to me. It is enough to mention the classifiers (phenology, photosynthetic type, etc.). Alternatively, one could provide a table showing
the different PFTs and their main attributes in the appendix, but I don’t think this is necessary for this manuscript.

Done. PFT listing is deleted.

I would appreciate some more information on the carbon and nitrogen pools mentioned on the same page, in brief. How do they communicate? On what time scales?

Noted. Text will be added as requested.

Most text in the following explanations is taken directly from the technical description (Oleson et al., 2013). Separate state variables for C and N are tracked for leaf, live stem, dead stem, live coarse root, dead coarse root, and fine root pools. Each of these pools has two corresponding storage pools representing, respectively, short-term and long-term storage of non-structural carbohydrates and labile nitrogen. There are two additional carbon pools, one for the storage of growth respiration reserves, and another used to meet excess demand for maintenance respiration during periods with low photosynthesis. One additional nitrogen pool tracks retranslocated nitrogen, mobilized from leaf tissue prior to abscission and litterfall. Altogether there are 20 state variables for vegetation carbon, and 19 for vegetation nitrogen.

Decomposition of fresh litter material (including C and N) into progressively more recalcitrant forms of soil organic matter is represented as a cascade of transformations between decomposing coarse woody debris (CWD), litter, and soil organic matter (SOM) pools. The decomposition flux is the product of a decomposition rate and source pool size.

Depending on the C:N ratios of the upstream and downstream pools and the amount of carbon lost in the transformation due to respiration, the execution of this potential carbon flux can generate either a source or a sink of new mineral nitrogen. Steps that result in an uptake of mineral nitrogen (immobilization fluxes) are subject to rate limitation, depending on the availability of mineral nitrogen, the total immobilization demand, and the total demand for soil mineral nitrogen to support new plant growth.

If mineral N is less than the demand to support new plant growth and immobilization, both plant growth and immobilization are downregulated by the same fraction in order to match N supply.

In addition to the cycling of nitrogen within the plant – litter – soil organic matter system, CLM represents external sources, including atmospheric deposition and biological nitrogen fixation. CLM also represents sinks, including nitrification, denitrification, leaching, and losses in fire.

There is an abrupt jump from leaf level photosynthesis (the models by Farquhar and Collatz) to the canopy level (GPP). Please add a short sentence explaining how photosynthesis is scaled to the canopy level. However, I think it would make more sense to explain this (p.4, lines 14-19) after equation 6, and not in the general description of the model.

Noted. Text will be added as requested.

The maximum rate of carboxylation at 25 °C varies with foliage nitrogen concentration and specific leaf area (Thornton and Zimmermann (2007)) and is a PFT specific parameter. It is assumed that leaf nitrogen and sunlight decrease exponentially with cumulative leaf area index from the canopy top to
Accordingly, the carboxylation rate and other photosynthesis parameter decrease exponentially within the canopy. Leaf level photosynthesis is scaled to the canopy level by integration over all leaf area. This is done separately for sunlit and shaded leaves and by considering the exponential scaling.

In addition, rather than describing the stomatal model in words on page 6 and showing the Equation in the discussion (Eq. 14), I would show the equation at this point. Please make sure that its original source is cited and that the notation is consistent: here you use “ca” for atmospheric CO2, later in Equation 14 “CO2” is used.

Done. Equation 14 moved to method section as requested. Reference to (Ball et al., 1987) will be added.

Equation 6: Could this equation be double-checked? In my understanding the last term of this equation should be the overall resistance, i.e. \(1/(1.6*gs) + 1/(1.4*gb)\), which differs from the term here.

Done. Equation is correct and given as implemented in CLM.

Conductance, \(g\), and resistance \(r\), are inversely related. The boundary layer, \(gb\), and stomatal, \(gs\), conductance of H2O is related to the boundary layer, \(rb\), and stomatal, \(rs\), resistance of H2O by:

\[ g_s = 1/r_b \quad \text{and} \quad g_s = 1/r_s. \]

Total resistance is the sum of individual resistances. For H2O the total resistance is:

\[ r = r_b + r_s. \]

For CO2, diffusive transport is slower than for H2O as the molecular mass of CO2 is higher than of H2O. Thus, resistance is larger for CO2 than for H2O (by a factor of 1.6 for stomatal, molecular diffusion and by a factor 1.4 assumed for the boundary layer). It holds for the total resistance of CO2:

\[ r_{CO2} = 1.6 \times r_s + 1.4 \times r_b = 1.6/g_s + 1.4/g_b. \]

Rearranging yields for the conductance:

\[ g_{CO2} = 1/r_{CO2} = 1/(1.6/g_s + 1.4/g_b) = g_s + g_b / (1.6 \times g_b + 1.4 \times g_s). \]

Please also check the equation on page 6 l.13, including the unit for conductance.

Done. Equation and units correct. Conductance is usually related to the mixing ratio and given in units of mol m\(^{-2}\) s\(^{-1}\). Here, conductance is related to the partial pressure of the gas and given in units mol m\(^{-2}\) s\(^{-1}\) Pa\(^{-1}\). Typesetting has been corrected and “Pa-1” is now correctly typeset as “Pa\(^{-1}\)”.

The information on the LPX-Bern model is quite detailed and in some parts unnecessary. Again, the information on the PFTs can be shortened. E.g. for this paper it is not relevant what PFTs grow on peatland. Descriptive text elements such as “The CO2 flux from the atmosphere to the stomatal cavity is proportional to the CO2 difference between the atmosphere and the stomatal cavity (ca - ci)” are not needed and can be seen from the Equations (e.g. Eq.8) or are physical principles.
Done. The list of PFTs is deleted for natural non-peatland areas. The PFTs for peatland are still mentioned as discrimination is set constant for Spagnum Moss to 30 permil and is following the C3 path for C3 graminoids. This is of some relevance for the simulated δ13C signature of vegetation and GPP as shown in Figure 4 and the global terrestrial isotopic budget as given in Figure 1 and Table 2. The sentence on the CO2 flux has been deleted as requested.

The stomatal control as simulated in LPX is poorly described. It is stated that ci/ca is set to 0.8 for non-water stressed conditions. That reads as if ci/ca is constant whenever there is enough water, even under low light, high VPD etc. Further, it is not really clear how this optimization works. Is it an optimization in the sense of Cowan & Farquhar 1977? If yes, the original reference should be cited. If not, it would be good to either elaborate this aspect or cite another study at this point where it is explained in more detail.

Noted. Text will be clarified as requested, and in the below we provide an answer to the above question and additional information.

Yes, ci/ca is set to 0.8 whenever there is enough water. Note that the photosynthesis-conductance routines are solved on a daily time step in LPX.

The stomatal control, as many other model details are described by Sitch et al. (2003). The following description is taken from Keel et al. (2016).

Daily evapotranspiration is calculated for each PFT as the minimum of a plant- and soil limited supply function (E_supply) and the demand for transpiration (E_demand). E_supply is the product of root-weighted soil moisture availability and a maximum water supply rate that is equal for all PFTs (Sitch et al., 2003). E_demand is calculated following Monteith’s (1995) empirical relation between evaporation efficiency and surface conductance:

\[ E_{demand} = E_{eq} \alpha_m (1 - \exp\left(-\frac{g_m \phi}{g_m}\right)), \]  

(Eq. 1)

where \( E_{eq} \) is the equilibrium evaporation rate, \( g_m \) and \( \alpha_m \) are empirical parameters that are equal for all plant functional types, \( g_c \) the canopy conductance, and \( \phi \) the fraction of present foliage area to ground area (i.e., projected leaf area). Above equation is solved for \( E_{demand} \) using the non-waterstressed potential canopy conductance as calculated by the photosynthesis routine for a fixed ratio \( \lambda \) between the CO2 mole fraction in the stomatal cavity and the ambient air. \( \lambda \) is set equal to 0.8 following Sitch et al. (2003) to approximate non-water-stressed conditions and as a starting value for the iterative computation of carbon assimilation and transpiration. In case of water-stressed conditions when \( E_{demand} \) exceeds \( E_{supply} \), canopy conductance and photosynthesis are jointly and consistently down-regulated; \( E_{demand} \) is set to \( E_{supply} \) and Eq. (1) is solved for \( g_c \).

Photosynthesis is modeled following Collatz et al. (1991, 1992), which is based on the formulations by Farquhar et al. (1980) and Farquhar and von Caemmerer (1982) generalized for global modeling purposes. The N content and Rubisco activity of leaves are assumed to vary seasonally and with canopy position in a way to maximize net assimilation at the leaf level. For C3 plants assimilation is a function of the daily integral of absorbed photosynthetically active radiation. For a detailed description see Haxeltine and Prentice (1996a,b).

Canopy conductance, \( g_c \), is linked to daytime assimilation, \( A_{dt} \), through
\[ g_c = g_{\text{min}} + \frac{1.6A_t}{c_a(1 - \lambda)}, \quad \text{Eq. 2} \]

where \( g_{\text{min}} \) is a PFT specific minimum canopy conductance, \( c_a \) is the ambient mole fraction of CO\(_2\), and \( \lambda \) the ratio between the CO\(_2\) mole fraction in the stomatal cavity and the ambient air. The equations for water supply and demand, assimilation, and canopy conductance are solved simultaneously by varying \( \lambda \) to yield self-consistent values for \( \lambda, g_c, \) assimilation, and transpiration.

Are the two models forced with two different meteorological datasets? CRUNCEP and CRU TS3.23? Is there a reason for this? And could that affect the results in some way?

Yes, the two models are forced by two different, though closely related products for technical reasons. CLM4.5 is run with a sub-hourly time step, while a daily time step is used in LPX.

This does not affect our conclusions and differences in these data products cannot explain differences in simulated discrimination and iWUE, as we explain below.

CRU NCEP (https://www.earthsystemgrid.org/dataset/ucar.cgd.ccsm4.CRUNCEP.v4.html) is used to force CLM4.5. CRU NCEP is a combination of two existing datasets (ftp://nacp.ornl.gov/synthesis/2009/frescati/model_driver/cru_ncep/analysis/readme.htm): The CRU TS3.2 0.5°x0.5° monthly climatology covering the period 1901 to 2009 and the NCEP reanalysis 2.5°x2.5° 6 hours time step beginning in 1948.

The CRU TS3.2 climatology offers a good spatial resolution but only monthly mean fields are available which is a too low resolution for CLM4.5. On the other hand NCEP reanalysis has a temporal resolution of 6 hours and is compatible with the CLM4.5 time step. But the spatial resolution is low and precipitation of such reanalysis is known to be less reliable than CRU data based on station data.

As evident from Figure 9 and 10 of the manuscript, the influence on changes in climate on the change in discrimination and iWUE is small, except in semi-arid regions. But to more comprehensively address the reviewers’ question, we further investigated the potential impact of using two different data products. The CRU NCEP data used to force CLM4.5 are interpolated on the 1x1 degree LPX grid and integrated to monthly values. This re-gridded CRU NCEP data are then used in LPX. LPX results for both climate input data, CRU NCEP and CRU-TS3.22, are compared. We investigated two cases, the standard model setup where climate is changing transiently and the factorial setup with “constant climate”. Difference in simulated changes in discrimination and iWUE are small for both cases (Figure R1).
Figure R1: Century-scale changes in iWUE (%) of C3 trees as simulated by LPX forced by (a) the CRU-TS3.22 (top panels) and the CRU-NCEP climatology (bottom panel). The left panels show results obtained with the standard model setup including transient climate and CO₂ forcing. The right panels show results obtained by keeping climate constant. In both setups, the influence of the different climate input data on results is very small. Changes are based on decadal means (1990 minus 1900). Changes in iWUE as calculated from d13C tree-ring data are shown by colored circles. The upper left panel is shown as the bottom panel of Figure 11 in the originally submitted manuscript.

Just for clarification: when referring to delta 13C forcing (e.g. p.8, l.21) it would be clearer to write atmospheric delta 13C.

Done. The term forcing is not used anymore for prescribed atmospheric CO₂ and 13C in the manuscript.

The sentence “An empirical convective boundary layer parameterization (Monteith, 1995) couples the carbon and water cycle” does not make sense to me. Please explain why the convective boundary layer couples the water and carbon cycle. To me it would be more helpful to read how the leaf boundary layer is treated in the model, as it directly affects your calculations (see Eq. 6). In general, when describing the models I recommend putting more emphasis on the calculation of variables that are directly used for later calculations or referred to in the Results section (e.g. calculation of the leaf boundary layer, are there differences in how soil water stress affects gs or A?). This will certainly be of greater interest to the reader than a list of PFTs that occurs in every land surface model in a similar form.

Noted. Text will be clarified as requested.

Please see our reply to the comments above on stomatal controls.

Page 8: not everyone is familiar with the discrimination model by Lloyd & Farquhar 1994. Please mention the key differences between the two models here (Lloyd & Farquhar 1994 and Farquhar 1989). The fact that the two formulations give similar trends is an interesting aspect but it is a little
bit hidden in the Methods section. Lines 14 – 20 are better moved to the discussion and can be extended. I think it would be good to be more precise here: what processes are not considered in the discrimination formulation and what does that change or not change? For instance, why is the agreement with leaf delta 13C worse when the more complex model is used? Why does it not change the trend? Discussing such aspects may not be the focus of this study but it would be a valuable contribution to the discussion on how isotopes are (or should be) considered in global models.

Noted. Text will be added and clarified as requested and we will consider moving lines 14 to 20 to the discussion.

Results Overall, this section is nicely written and clearly structured.

Thank you.

Model results are compared to a study by Carvalhais et al. 2014. It would be good to provide a bit more detail here. Do you mean aboveground and belowground vegetation carbon? How was vegetation carbon estimated in the study by Carvalhais et al. 2014?

Noted. Text will be clarified as requested.

Estimates of total vegetation carbon, including below and aboveground biomass, were derived by Carvalhais et al. 2014 from a collection of estimates for pan-tropical regions and for northern and temperate forests based on radar remote-sensing retrievals. Above- and below-ground biomass uncertainty for the tropical regions was propagated from errors in measurements, allometric relations, sampling and predictions. In the Northern Hemisphere, estimates accounted for uncertainties in the BIOMASAR GSV data, wood density data and biomass compartment data. Herbaceous carbon is estimated from GPP data products. See the original publication for a more detailed description and references.

Section 3.3: Results and Discussion are mixed here. It would be better to focus on the Results and discuss uncertainties in section 4. Just a thought: Why not taking PFT-specific model output? One could only take the corresponding PFT of the simulations that matches the PFT of the measured species. Up to the authors.

We prefer to keep the text on the caveats and uncertainties related to the tree ring data-model comparison at the beginning of section 3.3. This is important information for the reader and we feel allows the reader to better put the results into perspective.

We prefer to use grid-cell average model output instead results for individual PFTs. As explained in section 3.3., differences in δ¹³C are reported for different species. In addition, differences within a species, even growing at the same site, can be as large or even larger as those between species. Differences in the rooting depth, water transport systems, root-to-leaf distance, leaf morphology, or in irradiance (sunlit versus shadow) affect discrimination. Further, the models as applied here use generic PFTs and not individual species.

The authors suggest that there is a stronger downregulation of stomatal conductance by water stress in LPX than in CLM4.5 in some regions. Here, it would be helpful to provide some possible explanations. Is it because water stress in LPX is stronger due to the climate forcing, the way soil moisture is simulated, or due to a stronger stomatal response to water stress?
Differences in the applied climate forcing are not responsible for the stronger downregulation of stomatal conductance in CLM compared to LPX (see Figure R1 and related discussion above).

Soil moisture and stomatal conductance are coupled and influence each other in both models in a non-linear way hampering somewhat a firm attribution of signals to these two processes. The available results suggest that the primary reason for the difference in modeled 20th century changes in discrimination and iWUE is rooted in the different parameterizations of the photosynthesis-conductance coupling in the two models.

**Figure R1:** Soil moisture (top) and 20th century changes (bottom) as simulated for CLM4.5 (left) and LPX (right) with the standard model setup. Changes are based on decadal means (1990 minus 1900). For CLM, soil liquid water content in kg m\(^{-2}\) and its change (1990 minus 1900) is shown. For LPX, the water-filled fraction of the available water holding capacity in the top soil layer and its change is displayed.

**Figure R2:** As figure R1, but with a model setup where climate is kept constant.
A primary input to soil moisture is precipitation which is similar in both models. Patterns of soil moisture are similar in both models (Figure R2, top).

20th century changes in soil moisture are small in the factorial simulations where climate is kept constant (Figure R3, bottom). For the standard model setup, changes in soil moisture simulated by CLM and LPX are also small in large parts of Europe and Asia, where most of our tree-ring data are located. Despite these small changes in soil moisture, large differences in discrimination and iWUE changes are found between the two models in these simulations and regions (see Figure 9, 10, 11).

This is in line with the conclusion stated in the manuscript at the end of section 3.4 (p15, l33-35): “This suggests that the simulated decrease in discrimination and in c_i/c_a in the CLM4.5 runs is mainly linked to increasing CO₂ and a corresponding downregulation of c_i/c_a (Eq. 6). The downregulation in CLM4.5 is larger than suggested by the tree-ring records.”

The following text is added to the manuscript at the end of section 3.4: “Soil moisture and stomatal conductance are coupled and influence each other and in turn 13C discrimination. 20th century changes in soil moisture are small in the factorial simulations where climate is kept constant. For the standard model setup, changes in soil moisture simulated by CLM and LPX are also small in large parts of Europe and Asia, where most of our tree-ring data are located. Despite these small changes in soil moisture, large differences in discrimination changes are found between the two models in these simulations and regions (Figure 9). This suggests that the primary reason for the model-model difference in simulated 20th century changes in discrimination is rooted in the different parameterizations of the photosynthesis-conductance coupling.”

Discussion This section contains many interesting thoughts, but its structure is not very clear. If it was divided in several subsections as it is the case for the results section, it would be easier to find certain aspects the reader is interested in.

Noted. We will aim to clarify the structure of the discussion section.

p. 17, l.14-18: This paragraph can be expanded. As mentioned before, the differences between the discrimination model used here, and a more complex one (e.g. Lloyd & Farquhar, 1994), as well as possible implications for the simulated absolute values of discrimination and its trends can be discussed in more detail.

Noted. Text will be added as requested.

p. 18: The question that the reader will have is: why does CLM4.5 simulate such a strong trend in iWUE? The authors provide two possible explanations: 1) the downregulation of photosynthesis by nitrogen, and 2) an inadequate relationship between simulated stomatal conductance and assimilation. The first one is described well and is supported by other recent studies. In this context it would be helpful to know how the fdreg factor in Equation 6 changes over time, and whether it affects the relationship between An and gs.

The downregulation by nitrogen does not directly affect An and gs for carbon. It affects the amount of GPP that is allocated to the carbon pools. We will clarify this when revising the model description in the method section.
Figure R4: An index for nitrogen limitation, FPG top) and its 20th century change (bottom) as simulated by CLM4.5. FPG is indicative of nitrogen limitation and varies between 1 (no nitrogen limitation) and 0.

Nitrogen demand is the amount of nitrogen required to transfer all available carbon from photosynthesis into plant carbon. The fraction FPG describes by how much nitrogen demand to support plant growth is downregulated to match nitrogen availability. A value of 1 corresponds to non-nitrogen limited conditions. As shown in figure R3, 20th century changes in annual and grid-cell mean FPG are relatively small.

The fraction $f_{\text{dreg}}$ used to calculate the intercellular CO$_2$ partial pressure and thus $^{13}$C discrimination (Eq. 4 and 6 in the submitted manuscript) is closely related to FPG. We therefore also expect that annual mean values in $f_{\text{dreg}}$ changed little over the 20th century (Unfortunately, output for $f_{\text{dreg}}$ is not available).

It remains difficult to draw conclusions from annual or monthly values of FPG and $f_{\text{dreg}}$ as these variables are evaluated on the model time step and influence simulated fractionation and iWUE in a non-linear way (see e.g. Eq 4, 6, and 9 in the submitted manuscript).

Concerning the second explanation, I don’t understand what the key message should be. Is it the general form of the Ball-Berry model and the prescribed relationship between gs and An? In this case it should be mentioned that this model or similar models are used in most land surface models (see e.g. Sato et al. 2015, JGR Biogeosciences). If the reason for the strong iWUE trend is due to an inadequate relationship between gs and An, we should see a similar behavior in other land surface models. A comparison with other models is missing here.
CLM4.5 employs the Ball-Berry equation using a constant time invariant slope value, m (Eq. 14). Thus, any potential adjustment of m to changes in environmental conditions, including the century-scale increase in atmospheric CO₂ or changes in water stress are not considered. It is currently unclear whether such adjustment processes occur and our current understanding of the underlying physiological mechanisms of stomatal responses is less than complete (e.g. (Miner et al., 2016)).

Sato et al. (2015) investigate the influence of the use of vapor pressure deficit (VPD) or relative humidity (RH) in Ball-Berry-type stomatal conductance formulations. About half of the investigated models apply RH as a driving variable (Eq. 14), despite that VPD is considered the more relevant controlling factor. The global warming simulations reveal an increase in VPD and little change in RH. Their results suggest that the increase in VPD under global warming leads to a stronger downregulation of stomatal conductance (gₛ) and ci/ca for the VPD formulations compared to the RH formulation. This implies that replacing RH by VPD in the Ball-Berry equation may, without further adjustments, even increase the disagreement between modeled and reconstructed changes in discrimination and in iWUE.

We will clarify the text in the revised manuscript according to the above two paragraphs.

It is then argued that the trend may partly be attributed to changes in relative humidity, but no data are shown that would support this statement. What does the CRUNCEP climate forcing dataset suggest? Is there a trend in relative humidity that could explain the strong trend to some extent? Do areas that show a decrease in discrimination also show a decrease in relative humidity? The role of relative humidity (and possibly other climate variables) is an interesting aspect to discuss at this point, but it should be supported by data and discussed in context of the factorial simulations that were made.

This was not our intention. We do not argue that the changes in discrimination are due to long-term changes in relative humidity. We rather suggest “that assimilation is shifted towards times with lower relatively humidity (page 18, line 28)”.

This point is rather subtle and follows from Eq. 14. We will delete the text from lines 25 to 31 on page 18 of the submitted manuscript to avoid misunderstanding and for brevity.

It also mentioned that the value of the stomatal slope parameter m might be too high. It would be good to provide some more information, here or in the method section. What is the value of m? Is it constant across PFTs? I agree that m is probably too high for coniferous forests, but not necessarily for other vegetation types. If the value of m is to be discussed here, the authors should at least cite Lin et al. 2015, Nature Climate Change, who looked more generally at patterns of m across PFTs. They used a slightly different model, but that shouldn’t affect the patterns of m, see also Miner et al. 2016 Plant, Cell & Environment.

Text will be adjusted as requested and references added. The value of m is now mentioned in the methods section. Parameter values are m=9 for C3 plants and for m=4 C4 plants. These parameters are time invariant.

The review of Miner et al. (2016) yields mean values for m of 9.8, 8.7, and 6.8 for angiosperm evergreen, angiosperm deciduous and for gymnosperm trees, respectively (Their figure 1).
Changing m would certainly affect the absolute values of iWUE and discrimination, but would it make a difference to the simulated percentage trend in iWUE as shown in Figure 7? If the value of m is taken as a reason for the overestimated trend in iWUE by CLM4.5 this needs to be shown somehow. In my opinion, a change in m would primarily change the spatial patterns of the simulated discrimination.

Duarte et al. (2016) present in their figure A1 results from sensitivity simulations where only the parameter m was changed in the model setup. The change in leaf δ13C is larger and the change in discrimination is smaller for the lower value of m (m equal 6 instead of 9).

The formulation implemented in the LPX is better able to capture the observed iWUE trend. But is that really because of the optimization? I would argue that also the Ball-Berry model (Eq. 14) predicts a constant ci/ca and thus a trend in iWUE that is proportional to ca, provided that rH and m do not change over time. In my eyes this is indicative of changes in rH, or more likely, problems with the nitrogen downregulation, as discussed earlier. Why not testing this? The CLM4.5 model could be run with a version that does not include the nitrogen downregulation. The comparison of this alternative version with the version used in this study could be used to answer the question whether the problem lies in the nitrogen downregulation or in the stomatal conductance scheme (Eq. 14). If the alternative model version still shows a stronger iWUE trend than expected, this would be a stronger indication that an optimization based approach indeed works better. Maybe new global runs are not necessary, and a simple analysis based on Eq. 6 would suffice. However, without testing this, the statement (p.20, l.27f) remains speculative and should not be mentioned in the conclusion of the paper. In general, this part of the discussion needs to be revised according to the comments above.

We will delete the statement (“and may be interpreted as supportive of a plant strategy towards optimizing assimilation and minimizing water loss under changing environmental conditions”) on p20, l27 as requested. The remaining text will read: “The results suggest constant ci/ca over the 20th century. The results suggest that it is desirable to adjust the implementation of photosynthesis and conductance in CLM4.5 towards a better agreement with observation-derived century-scale trends in δ13C discrimination and intrinsic water-use efficiency.”

p. 19: The behavior of iWUE and ci/ca as reconstructed from the tree-ring measurements and modeled by LPX-Bern is compared to other studies. The nice thing on this paragraph is that it is very comprehensive. But it could be clearer with respect to the method used in the cited studies. Rather than just listing the studies you could sort them by method, i.e. mention other isotope-based studies first, then other methods. At the moment studies using the same methods (e.g. FACE) are mentioned in different parts of this section (Ainsworth & Long, 2005 and De Kauwe et al. 2013) which seems a bit fuzzy. I think this aspect is important as different methods are associated with different uncertainties (which, however, do not have to be discussed here).

Noted, we will consider this option during the revision of the discussion section.

With respect to the eddy covariance records it may be interesting to mention that a recent study (Knauer et al. 2016, New Phytologist) found that large-scale carbon and water fluxes are not in agreement with a constant ci, but rather with a constant ci/ca.

Noted. We will complement the text and add the citation.
p. 20: The effects of land use change and representation issues between the datasets/model simulations are adequately addressed. It may be helpful for the reader to mention the Figures again where the described aspects can be seen.

Noted. Figures will be mentioned where appropriate.

Figures In some figures (e.g. Fig. 2), the color code and the associated numbers are very small and hard to read. It would be ok to have fewer color classes as they are hard to distinguish.

Noted. We will reduce the number of labels and increase their size in Figures 2 to 4. We will also change units from gC m\(^{-2}\) to kgC m\(^{-2}\) to reduce the number of digits.

Fig. 2: For the difference maps, please state what is subtracted from what, at least in the legend.

The differences are based on decadal means (1996-2005 minus 1896-1905). The caption will be adjusted.

Fig. 5: representing the differences in mean delta13C as barplots is not appropriate here. I recommend to remove the bars and show the error bars only, also in Fig. 6.

We prefer to keep the bars for a clear distinction between the difference and the error estimates.

From Fig. 5 onwards: Some of the points on the map are hard to see. It would be helpful if their representation could be changed.

Our choice represents a trade-off between the visibility of individual dots and the visibility of all dots.
Review 2: A. Ballantyne

Here the authors compare a compilation of tree ring and leaf isotope data from around the world with isotopic simulations from two common land surface models. While several studies have compared isotopic estimates of iWUE with model simulations of iWUE, especially at regional scales. This study is novel in that it is one of the few to actually investigate isotopic tracer simulations within models as a critical diagnostic for how accurate models are at simulating the global C cycle. In principle, this approach allows us to evaluate to what extent the terrestrial biosphere is being fertilized by increased atmospheric CO2; however, I think that the authors could further partition the response of iWUE into its component processes of assimilation and transpiration (at least in the models). This may also help reconcile why the models appear to show differing degrees of iWUE response.

General Comments:

I suspect that the two models investigated here differ considerably in how stomatal conductance is simulated and this is having a big impact ultimately on the isotopic tracers. While these models may be responding similarly to increases in atmospheric CO2 they may be responding to different metrics of atmospheric water vapor. As the authors point out, assimilation in CLM is modeled as a function of RH and CO2, while it is my understanding that in LPJ stomatal conductance is modeled as a function of VPD. While RH and VPD may be inversely related in some environments, this is not always the case and their relationship might vary over the 20th century. It would be nice to see how assimilation and transpiration have responded over the 20th century independently in the two models. This may also help explain why LPX and CLM show different responses of iWUE over the 20th century.

Figure R5: Simulated transpiration (in mm/s) for the decade around year 2000 (top) and 20th century changes (bottom; 1996-2005 minus 1896-1905).

The panels shown in Figure R5 will be added to figure 2, next to the panels showing GPP and GPP changes (Layout and labels will be adjusted).
Simulated 20th century changes in annual-mean transpiration are generally small in both models, except in Australia for LPX and in parts of Latin America for CLM4.5. Generally an increase in transpiration is found in boreal and temperate forest regions in both models. Transpiration is slightly decreasing in LPX and slightly increasing in CLM4.5 in most tropical forest regions. The overall increase in water use efficiency (not shown), the ratio of assimilated carbon to transpired water, is primarily driven by the increase in GPP in both models.

**Specific Comments:**

P1L12 ‘water loss by transpiration.’

Done. Text changed as requested.

P2L4 ‘and water transpiration’

Done. Text changed as requested.

P2L22 Graven article is on 14C not 13C as cited. Check reference as they may have also included 13C in their simulations.

Done. Reference removed.

P2L27 conductance can be of CO2 or H2O, could be more specific here and say ‘transpiration’ as the process and H2O as the mass.

Stomatal conductance is used here as a general term for H2O and CO2. Transport of 12CO2, 13CO2, and H2O through the stomata are by molecular diffusion. This process depends on molecule mass and leads to the “fractionation” between the 13CO2 and 12CO2 fluxes. Text is not changed.

P3L7 While the authors mention many 13C tree ring records, they fail to mention the pioneering work by Tans et al. which is found in the references.

Done. Reference added.

P3L22 ‘to complement recent advances in simulating marine carbon isotopes’

Done. Sentence changed to read “This is a step towards fully coupled isotope-enabled CESM applications and complements recent advances in simulating marine carbon isotopes”.

P4L24 ‘reactant to product’

Done. Text changed as requested.

P5L10 I believe that diffusion is only relevant for fractionation in non vascular plants such as bryophytes as well.

Done. Text clarified by adding reference to Farquhar et al., 1989 at the beginning of the paragraph. Non-vascular plants are not included in the standard setup of CLM4.5.

P6L6 more realistically related to the gradient between internal water pressure and atmospheric water pressure (approximated as vapor pressure deficit or VPD).
Done. Text clarified. The description of the relationship between conductance and assimilation in CLM4.5 is clarified and Eq. 14 (submitted MS) is moved here to page 6.

P7L7 del 13C signature of what? Atmosphere? Please clarify

Done. Text clarified to read: “Note that no $\delta^{13}$C observational data, e.g., from tree-rings or atmospheric samples, was used as a constraint in the assimilation.”

Eqn 8 Isn’t this the same as Eqn ?? But not quite sure $c^*$ is specified in Eqn 7.

Noted. Model description will be revised in reply to the comments by referee 1 and Eq. 8 may be deleted.

P9L9 del 13 C is estimated as the ‘weighted flux’ of component GPP fluxes from PFTs from within grid cell. Omit ‘GPP is used as a weight’.

Done. Sentence deleted as requested.

Eqn 11. I don’t think that this equation is necessary (especially given the number of equations already included) and this can simply be explained.

Done. Equation 11 deleted and text adjusted.

P10L14 While this approximation of 36.16 holds well within 0.1% isotopic ratio differences are per 1000, so is this enough significant figures?

Done. Text clarified. The approximation is for Eq. 12 (in the submitted MS) and not for 36.16. It holds exactly: $q=1.6 (a-b) = 1.6 (27-4.4) = 36.16$. Additionally, four significant digits are enough as units here correspond to “permil units” and are not “per 1000”. Text clarified to read: “The approximation given in Eq. 12 holds well within 0.1%.”

P11L1 Not sure that you need to correct for the offset if you are only focusing on the trends and normalizing them across sites, regardless this should not affect your analysis.

We agree with the reviewer that this correction hardly influences our analyses and does not alter conclusions. Nevertheless, the relative change in iWUE depends weakly on the magnitude of the discrimination ($\Delta(t_1)$) as evident from equation 13 and as clearly stated in the manuscript.

P12L18 Were the CLM and LPX simulations conducted with or without land use change and does this have any impact on the global isotopic budget.

Yes, anthropogenic land use is considered in the CLM and LPX simulations as stated on p6, line 26 for CLM and on page 8, line 22). Land use maps from Hurtt et al., 2006 are used in both models. Implications for the global isotopic budget are discussed on page 20, line 14 to 21 in the submitted manuscript.

P13L13 ‘changes in the atmospheric del13C source’

Done. Text adjusted as requested.

P13L15 ‘globally-averaged’ what? Soil, atmosphere?
Done. Text clarified to read: “There is a substantial offset between the models for globally-averaged soil and vegetation pools, with CLM4.5 being around 5 per mil more negative.”

P13L34 2.42 and 3.22 per mil these should have units
Done. Per mil units added.

P14L8 Maybe these global mean estimates should be reported first before noting all the regional differences and more nuanced results.
Done. Global mean estimates are now reported before the regional differences.

P14L12 ‘bias of the models’
Done. Typo corrected.

P14L28 Were any trend statistics (e.g. Mann-Kendall) conducted on the observations or models?
As suggested, we performed a Mann-Kendall analysis on the discrimination time series. The trend in discrimination of C3 trees is most monotonic in CLM-4.5 ($\tau = -0.9$), less monotonic in the average tree-ring record $\tau = -0.6$ for annual data and -0.7 for decadal mean data) and almost negligible in the LPX-Bern output ($\tau = -0.2$).

The Mann-Kendall test provides information about the extent to which a trend is monotonic, but does not provide information about the magnitude of the trend. We do not plan to add the results from this test to the manuscript. It is clear from Figure 7 that the trends in discrimination and iWUE are larger in CLM4.5 than in LPX and the tree-ring data (see also Figure R6 below).

P15L10 These aren’t really ‘spatial’ correlations
Done. Word spatial deleted.

P15L15to21 This paragraph seems to fit better in the methods
Done. Paragraph moved to method section. This implied a slight rearrangement of the subsections in the method section. Spin-up and transient simulations for both CLM4.5 and LPX are now described in the same subsection.

P16L24 Model simulations with increased CO2 and constant climate change could be compared at least quantitatively to FACE data.
This is beyond the scope of this study. For a comparison between carbon-nitrogen land models and FACE data we refer to the literature, e.g., Zaehle et al. (2014)

P16L34 ‘Recall, however, that ...represent annual or multi-annual averages that have been weighted by C assimilation or alloclation’
Done. Text adjusted as proposed.

P17L28 This paragraph is rather short think about combining.
We prefer to keep the paragraph as is to give the same weight to the conclusions drawn at the end of this and the previous paragraph.

Eqn. 14 Would also be interesting to compare how conductance is simulated in LPX. While assimilation in both models is clearly responding to increasing atmospheric CO2, I suspect that transpiration may be responding differentially in the models due to different stomatal response to atmospheric water demand.

Noted. The methods section will be adjusted to better explain how conductance is modelled in LPX. See response to a similar comment by reviewer 1.

P18L27 For the CLM response you should look at the relative changes in CO2 and relative humidity over time (this should be a prognostic variable in the model). Also see work by Isaac Held on the response of the hydrologic cycle to atmospheric warming. Essentially, at the global scale RH does not change in response to warming; however, this might not be true over land. So it would be interesting to see in CLM how RH has changed at the tree ring sites.

This point is rather subtle and follows from Eq. 14. We will delete the text from line 25 to 31 on page 18 of the submitted manuscript to avoid misunderstanding and for brevity. Please see also reply to reviewer 1, concerning this point.

P20L5 Similar work by Penuelas et al (2011) has shown an increase in water use efficiency but not necessarily an increase in annual ring width. However, a true test would be the relationship between WUE and Biomass - not sure if Klein looked at biomass in this study.

Noted. We will mention the work by (Peñuelas et al., 2011).

P20L9 It seems that both of these FACE studies report a consistent increase in WUE, but of slightly different magnitudes. It is interesting that the responses are so different between European forests and the N. American forests. Unfortunately, most of the FACE studies have been conducted in the Eastern US, where there are no tree ring isotope records.

Thank you. We do not plan to extend the discussion on these FACE studies.

Figure 3. not so sure that the Carvalhais estimates are ‘observations’, maybe ‘derived from’ or ‘constrained by’ observations.

Done. Text modified to read “derived from observations”.

Figures 5 and 6. I am not sure that you need both of these figures as they illustrate the same data. Perhaps move one to supplemental.

We will keep both figures as part of the manuscript. This decision is also in response to the comment of reviewer 1 on the visibility of the tree-ring data dots. Showing only regional maps as in Figure 6 would provide incomplete information of the global picture, whereas showing only the global map as in Figure 5 would affect clarity and visibility of regional results.

Figure 7. Can you include all of the tree ring records as thin grey traces in this figure? Would be nice to see some distribution of the observations to see if all the obs are bound by the model simulations.
Figure R6: Changes in discrimination and iWUE. Similar as figure 7 in the original manuscript, but with individual tree ring series added (thin gray lines). All model and tree-ring series represent decadally-smoothed values.

Figure R7: Same as Figure R6, but tree-ring series represent annual values.

We will replace Figure 7 with Figure R6 as requested.

Figure 8. Not sure that you need the discrimination equation, which should be defined in the text.

Done. Equation removed from figure caption. Equation 5, showing this definition, is now mentioned.

Figure 9. The right hand panels where certain variables have been kept constant is not explained in the caption.

Done. Explanation added to the caption.
Figures 11 and 12. Once again these figures are both great but they illustrate redundant information maybe move one to the supplemental information.

We will keep both figures as part of the manuscript for the same reason as discussed above for figures 5 and 6.

In summary, with tree ring isotope data we are only able to approximate iWUE and cannot partition this response between assimilation and transpiration. However, in the models you can partition these processes, so it would be interesting to see how transpiration and assimilation are responding in the models, which may help identify processes that can reconcile these model simulations.

Figure on transpiration will be added as requested. See response to main comment above.

References:


