Interactive comment on “Glacial–interglacial shifts in global and regional precipitation δ18O” by S. Jasechko et al.

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Re: C338: ‘interactive discussion’, Reviewer 3

We thank Reviewer 3 for their thorough review of our paper and for their pointing out the usefulness of model-data comparisons of late-glacial and late-Holocene precipitation isotope compositions.

The central idea behind the work of Jasechko et al. was the reconstruction of climatically-induced shift in the isotopic composition of global precipitation between the last Glacial and the Holocene, and to confront it with the predictions of state-of-the-art isotope-enabled general circulation models. The authors selected three different proxies of isotopic composition of past precipitation (groundwater, speleothem calcite, ice cores), conducted extensive literature search and came up with the reconstructed δ18O of precipitation for two time windows: (i) late Holocene (0 - 5000 calendar years), and (ii) Glacial (19500 - 50000 calendar years) at number of sites distributed globally. The resulting spatial distribution of the reconstructed δ18O(ice age) was then compared with the modeled δ18O(ice age) generated by five GCMs. This sort of global comparison was in fact long due and represents a valuable tool for assessing the performance of existing isotope-enabled GCMs. I highly appreciate the efforts and gigantic work done by the authors in compiling appropriate information. Still, when attempting this sort of comparison, a great care is needed in proper selection of the data in order to minimize the possibility of falling into "garbage in - garbage out" trap. Therefore, the proxy data selected for comparison should be carefully scrutinized. I do have a number of comments and suggestions which might assist the authors in improving the overall shape of the paper and which should be addressed in the revised version. They concern both the methodology used and the structure of the paper. Specific comments: 1. I would recommend adding a separate section in Chapter 2, where key characteristics of the selected proxies, of direct relevance to the presented model-data comparison, are discussed in depth. In particular, this discussion should highlight the following questions: (i) how well the given archive is preserving the (mean) isotopic composition of precipitation? (ii) what are the potential biases? (iii) can reliable chronology of the given archive be established?. Some information on those issues is dispersed throughout the main text and in the Supplement but the paper would definitely gain in clarity if all this information is gathered in one place. While the authors address to some extend the question of establishing age of groundwater (see comment no. 3), they refrain from any comment on the uncertainties of the chronology of speleothem and ice core samples used in the comparison, which can be significant (see e.g. Landais et al., (2015) for ice cores). We focus the majority of our revisions on this comment, pointing out potential
biases linked to seasonality (e.g., Werner et al., 2000; Jasechko et al., 2014; James et al., 2015) and uncertainties in the chronologies of each proxy record. We have added a new paragraph to our manuscript highlighting the potential for such biases.


2. It should be made clear that in view of significant uncertainties in establishing absolute chronologies of the archives selected for this work, particularly for the Glacial period, the boundaries of the selected time windows remain blurred. This is particularly true for the Glacial time window. Setting up a sharp lower boundary at 19500 calendar years does not make much sense in this context (should be rather 19000 or 20000 calendar years). We change the lower age limit to 20,000 years before present.

3. I have several objections with respect to the approach adopted by the authors to calculate groundwater ages, as described in the Supplement. Below we show how we revise our groundwater ages following the suggestion of the reviewer. Including the changes made below has resulted in little change in our groundwater-based measured \( \Delta^{18}O_{\text{late-glacial}} \) values. We agree with the reviewer that 14C-based ages have uncertainty, however, the plateauing of \( \delta^{18}O \) at each time interval and clear late-glacial to late-Holocene \( \delta^{18}O \) shifts supports our interpretation of such \( \delta^{18}O \) changes as records of late-glacial to late-Holocene climate change. We have added the following statement to the revised version of the manuscript: “Further, the chronologies of groundwaters and ice core records have uncertainties on the order of thousands of years, meaning that the time intervals used to calculate measured \( \Delta^{18}O_{\text{late-glacial}} \) values may be inaccurate. However, the plateauing of isotope content observed in most regional aquifers for 0-5,000 years before present and for >20,000 years before present supports our interpreting these data as records of late glacial to late Holocene isotopic shifts (see figures in the Supplement).”

(i) to calculate groundwater age the authors use eq.(S1) which numerical factor (-8267) contains more recent value for 14C half-life (5730 years). Then, they convert ages derived using eq.(S1) to calendar ages using corrections based on the calibration curve proposed by Fairbanks et al., (2005). However, by definition the calibration curve relates conventional radiocarbon ages to calendar ages (cf. Fig. 2 of Fairbanks et al., 2005). Conventional radiocarbon ages are calculated on the basis of Libby’s half-life (5568 years) which leads to numerical factor in eq.(S1) equal -8033, but not -8267. Besides, a more recent calibration curve (Reiner et al., 2013) synthesizing all available calibration data should be used rather than Fairbanks et al (2005) curve. We update our calculations using the updated calibration curve using data presented in Reimer et al. (2013). We also use the half-life proposed by Libby following the reviewer’s suggestion. This change led to the removal of a few samples that were previously used to calculate \( \Delta^{18}O_{\text{late-glacial}} \) values. Overall, groundwater-based \( \Delta^{18}O_{\text{late-glacial}} \) values changed little between our initial submission and this revised version.

Reimer, P. J. et al. (2013), IntCal13 and Marine13 radiocarbon age calibration curves 0-50,000 years cal BP, Radiocarbon, 55, 1869-1887.

(ii) Figures S4 - S62 have a horizontal axis labeled “Groundwater age (14C-years before present)”. Are those indeed radiocarbon ages calculated on the basis of eq.(S1), or perhaps radiocarbon ages converted already to calendar ages ?. In any case, they contain number of data points showing unrealistically high fi-
nite ages going up to 62000 years. The x-axis label for the supplemental figures is accurate (14C years). We agree with the reviewer that 14C ages exceeding 30 thousand years are highly uncertain, we modify our groundwater age calculations to convey the limitations associated with these old ages. However, some of the compiled works report high-precision 14C activities (e.g., 0.04 pmC in Larsen et al., 2002); we use the reported 14C activities as our best estimate of 14C content, but add an analytical uncertainty of ±1 pmC and propagate this added uncertainty through our age calculations.


(iii) In their uncertainty analysis the authors apparently forgot to include the analytical uncertainty associated with the measured radiocarbon content in the given sample (quantity A in eq.(S1)) Large majority of the reported radiocarbon data (my guess would be that this is around 80-90) was obtained by laboratories using conventional (i.e. decay-based) analytical techniques. Typical analytical uncertainty of radiocarbon analyses in such laboratories (one sigma level) usually varies between ca. 0.5 and 1.0 pmc (percent of modern carbon). In addition, sampling of groundwater in the field for radiocarbon analyses introduces additional source of uncertainty (possible contamination with modern radiocarbon from the atmosphere). Therefore, a realistic value for the Limit of Detection (LoD) can be set in this case around 1 pmc. Modern AMS laboratories can do a bit better but still the problem of contamination during sampling remains open. This LoD of approximately 1 pmc leads to conventional radiocarbon age of ca. 35000 years, not accounting for any geochemical correction - just taking into account radioactive decay only. This age limit transfers to approximately 40000 calendar years. Consequently, all groundwater ages higher than ca. 35000 conventional radiocarbon years or ca. 40000 calendar years should not be reported as finite ages but rather as ”> 35 ka BP or >40 cal. ka BP”. The additional uncertainty is now included in updated calculations. We report the best estimate of 14C age and show that 14C ages older than 35,000 years often have very large uncertainties (updated supplemental figures for each aquifer).

(iv) The authors use so-called Pearson correction model to account for the dissolution of carbonate phases in the aquifer (eq.(S2)). However, it is well-known since long time that this model does not describe correctly all possible interactions between the TDIC reservoir which is dated by the radiocarbon technique and the aquifer matrix. Often, more complex models which merge evolution of carbon isotopes with the evolution of water chemistry along the flowpaths need to be applied in order to obtain realistic groundwater ages. More advanced correction schemes may result in radiocarbon ages differing by several thousand years from the ages returned by application of the simple Pearson correction model. The reviewer is correct that 14C groundwater ages calculated in many studies are imperfect and susceptible to a variety of complexities (e.g., non-linear averaging of ages due to mixing; Torgersen et al., 2013). More complex age correction models are not appropriate for use here because of the wide variability of data available amongst the compiled publications. For example, some studies do not report major ion chemistry or $\delta^{13}$C values, whereas other publications report a more complete set of geochemical measurements.


Summarizing, the groundwater data need a major overhaul here. 4. In their compilation of groundwater data the authors do not refer to the IAEA database. This is by far the largest collection of isotope data for groundwater systems worldwide. Therefore, I would strongly recommend that three IAEA Atlases are consulted (Atlas of Isotope Hydrology - Africa, IAEA 2007; Atlas of Isotope Hydrology
ogy - Asia and the Pacific, IAEA 2008; Atlas of Isotope Hydrology - the Americas, IAEA 2009) and, if appropriate, additional data obtained from this source included in the global picture of reconstructed Delta18O(ice age) presented in the paper. The Atlases provided by the International Atomic Energy Agency are a useful facility, which highlight the many aquifers investigated by IAEA coordinated research projects. However, these Atlases do not explicitly point out occurrences of paleowaters, instead providing ranges of isotope compositions observed for different samples (e.g., springs, groundwaters). We agree that these data could be used in future studies to further investigate spatial patterns of paleowater isotope compositions.

5. In Chapter 3.3 the discussion of each region should be accompanied by appropriate regional maps showing the locations of relevant sites, each labeled by two numbers: reconstructed and simulated Delta18O(ice age). Colors should be avoided because they hide to some extent the real differences. Such regional maps would guide the discussion and would help to identify regions which are most problematic with respect to the model-data comparison pursued in the paper. Figure S3 should be then removed from the Supplement. Regional maps are added to the main text. We follow the reviewer’s suggestion and labelled each measured Δ18Olate-glacial value on the figure. Although we cannot match a single model simulated Δ18Olate-glacial value to each because we report five different general circulation models. However, we display the multi-model ensemble median as a grid to aid the discussion of model-versus-measured Δ18Olate-glacial values.

6. Figure S1 should be moved to the main text. Perhaps the authors may consider adding on the map presented in Fig. S1 the Holocene-Glacial noble gas recharge temperature differences reconstructed for number of aquifers which are included in the model-data comparison discussed in the text. Noble gas temperatures are considered excellent proxy of ground level air temperatures and it would be instructive to confront Annan and Hargreaves (2013) reconstructions with those derived from noble gas data. From this perspective, the 15 degree Celsius of temperature suppression for Hungary at the last glacial maximum, reported on page 848, line 16, is clearly an exaggeration (noble gas data indicate only 8-9 degree Celsius - see e.g. Corcho Alvarado et al., 2011). Global maps of surface temperature changes developed by Annan and Hargreaves (2013) will be added to the main text. We agree that future studies could compile and analyse the noble gas records, providing a useful set of paleo-temperature records that could be used to better understand the spatial patterns of late-glacial to late-Holocene temperature change. We have added reference to the temperature change for Hungary suggested by Deák et al. (1987) of 5-7°C, a lower value than proposed on the basis of geomorphic data (Fábián et al., 2014): “Geomorphic evidence suggests permafrost covered portions of Hungary at the last glacial maximum, suggesting that land temperatures may have been up to 15°C cooler than present day (Fábián et al., 2014), a larger late-glacial to late-Holocene temperature shift than earlier, noble gas based reconstructions (5-7°C; Deák et al., 1987).”


7. The conclusions should stress the fact that the compilation of reconstructed Delta18O(ice age) presented in the paper constitute a strong challenge for isotope enabled GCMs. Figure 3 makes it clear that the selected crop of isotope-enabled GCMs is not performing particularly well. The frustrating thing is that apparently not much progress has been made in this respect in the past twenty years or so. Perhaps the necessary first step towards improving the situation would be a comprehensive model-data comparison with the present-day spatial
distribution of mean delta18O(delta2H) in global precipitation. We agree with the reviewer that the models and observations do not agree in all locations. The compiled data presented here can be used to continue to improve isotope enabled general circulation models.

Interactive comment on Clim. Past Discuss., 11, 831, 2015.