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Understanding the ¹⁷O excess glacial-interglacial variations in Vostok precipitation

Camille Risi,¹ Amaelle Landais,² Sandrine Bony,¹ Jean Jouzel,² Valérie Masson-Delmotte,² and Françoise Vimeux³

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[1] Combined measurements of δ^{18} O, δ^{17} O, and δ D in ice cores, leading to d excess and ¹⁷O excess, are expected to provide new constraints on the water cycle and past climates. We explore different processes, both in the source regions and during the poleward transport, that could explain the ¹⁷O excess increase by 20 per meg observed from the Last Glacial Maximum (LGM) to Early Holocene (EH) at the Vostok station. Using a single-column model over tropical and subtropical oceans, we show that the relative humidity at the surface is the main factor controlling ¹⁷O excess in source regions. Then, using a Rayleigh-type model, we show that the ¹⁷O excess signal from the source region is preserved in the polar snowfall, contrary to d excess. Evaporative recharge over mid and high latitudes and δ^{18} O seasonality in polar regions can also affect the Vostok ¹⁷O excess but cannot account for most of the 20 per meg deglacial increase from LGM to EH. On the other hand, a decrease of the relative humidity at the surface (rh_s) by 8 to 22% would explain the observed change in ¹⁷O excess. Such a change would not necessarily be incompatible with a nearly unchanged boundary layer relative humidity, if the surface thermodynamic disequilibrium decreased by 4°C. Such a change in rh_s would affect source and polar temperatures reconstructions from δ^{18} O and d excess measurements, strengthening the interest of ¹⁷O excess measurements to better constrain such changes.

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1. Introduction

[2] Stable isotopic measurements of water (δ^{18} O and δ D) have been performed for more than 50 years [*Dansgaard*, 1953; *Epstein and Mayeda*, 1953; *Craig*, 1961] with the aim to improve our understanding of the water cycle and its links to climate. At first order, variations of δ^{18} O and δ D are mainly due to the difference in saturation vapor pressures between the light and heavy isotopes, leading to an equilibrium fractionation. In addition, the larger diffusivity of light isotopes with respect to the heavy ones results in a kinetic fractionation, involved in evaporation and in snowflakes formation. Since these kinetic effects have a larger influence on δ^{18} O than on δ D [*Merlivat and Nief*, 1967; *Merlivat*, 1978], there is an added value in combining δ^{18} O and δ D measurements.

[3] This second-order effect is measured by d excess = $\delta D-8 \cdot \delta^{18}O$ [*Dansgaard*, 1964]. The d excess in oceanic water vapor is a tracer of evaporative conditions (sea surface

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temperature, surface relative humidity, wind speed) and airsea interaction [e.g., Merlivat and Jouzel, 1979; Gat, 1996]. In particular, d excess in water vapor over the ocean increases when humidity over the evaporative ocean decreases, which led Jouzel et al. [1982] to interpret isotopic records in polar ice as a tracer of past humidity changes. However, numerous fractionation processes take place along the transport of air masses from the evaporative to the polar regions, so that d excess in ice is also influenced by the moisture source and condensation temperatures, and by the isotopic composition of seawater [Vimeux, 1999; Petit et al., 1991; Stenni et al., 2001]. For instance, modeling studies have shown that the effect of relative humidity changes in evaporative regions on polar d excess could be overwhelmed by the effect of sea surface temperature changes at evaporation [Vimeux, 1999]. Therefore, d excess is now often interpreted as a tracer of the source temperature [Vimeux, 1999; Stenni et al., 2001].

[4] Recent experimental developments have made it possible to accurately measure $H_2^{17}O$ abundance and to define a new isotopic tracer in the water cycle, ¹⁷O excess [*Barkan and Luz*, 2007; *Landais et al.*, 2008], expressed in per meg (10⁻³‰):

$$^{17}\text{O}\,\text{excess} = 10^6 \cdot \left(\ln \left(\frac{\delta^{17}\text{O}}{1000} + 1 \right) - 0.528 \cdot \ln \left(\frac{\delta^{18}\text{O}}{1000} + 1 \right) \right)$$

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with the δ notation defined as

$$\delta = \left(\frac{R_{\text{sample}}}{R_{\text{SMOW}}} - 1\right) \cdot 1000$$

where R_{sample} and R_{SMOW} are the molar ratio of the heavy to light isotopes in the sample and in the SMOW standard [*Dansgaard*, 1964]. Note that ¹⁷O excess and d excess have slightly different definitions: d excess is defined in a linear scale and ¹⁷O excess in a logarithm scale (see *Luz and Barkan* [2005] for the advantage of the logarithm scale).

[5] As δD and $\delta^{18}O$, $\delta^{17}O$ and $\delta^{18}O$ show different sensitivities to equilibrium and kinetic fractionation processes so that ¹⁷O excess, as d excess, has the potential to bring additional information on climate conditions and the hydrological cycle.

[6] The ¹⁷O excess and d excess do not show similar spatial distributions and are thus expected to be complementary [*Landais et al.*, 2008]. While d excess in precipitation features a strong poleward gradient over Antarctica [*Dahe et al.*, 1994; *Masson-Delmotte et al.*, 2008], so far ¹⁷O excess has shown constant values in present-day Antarctic surface snow [*Landais et al.*, 2008]. Because of these different behaviors, it has been suggested using simple isotopic modeling that ice ¹⁷O excess was a more direct tracer of the evaporative regions than ice d excess [*Landais et al.*, 2008].

[7] A record of 17 O excess over the last 150 kyr has been obtained from the Vostok ice core (East Antarctica, 78°S, 106°E). The most prominent features are the significant increases of ¹⁷O excess by 20 per meg during the last two deglaciations [Landais et al., 2008]. Using a Rayleigh-type distillation model initialized by the traditional closure assumption (i.e., assuming that the vapor originates from surface evaporation only [Merlivat and Jouzel, 1979; Landais et al., 2008]) suggested that the large increase in ice ¹⁷O excess recorded in the Vostok ice core over the last deglaciation was due to a decrease of relative humidity at the surface (rh_s) by 10 to 20% over the source regions. However, general circulation models (GCMs) suggest very small changes of the near-surface air relative humidity (rh_a) over oceanic regions from LGM to present day [Bush and Philander, 1999] or in the context of future climate change [Bony et al., 2006]. If rh_s and rh_a vary in concert, which is assumed in studies interpreting water isotopes in polar ice cores using simple models [*Vimeux et al.*, 2001; *Stenni et al.*, 2001], GCMs simulations contradict the interpretation of 17 O excess as a tracer of rh_s.

[8] The goal of this article is thus to explore various processes that might explain the observed glacial-interglacial ¹⁷O excess shift, both at the source and during the poleward transport. To this aim, we use a single-column model for tropical or subtropical source regions and a Rayleigh distillation model for the effect along the air mass trajectory, as described and justified in section 2. We explore the effect of source conditions on ¹⁷O excess of the moisture source in section 3 and possible modifications of ¹⁷O excess between the source and Vostok in section 4. We discuss paleoclimatic implications in section 5.

2. Method

[9] Atmospheric general circulation models (GCMs) incorporating water stable isotopes seem the ideal tool to understand

the isotopic composition of worldwide water vapor and precipitation. However, GCMs still have difficulties to simulate surface temperature and snowfall amount over inland Antarctica [Masson-Delmotte et al., 2008]. Moreover, they have difficulties to simulate d excess glacial-interglacial variations of the right sign over high latitudes [e.g., Werner et al., 2001]. This difficulty could arise from either deficiencies in representing isotopic processes such as kinetic effects, or from a poor simulation of some LGM climatic properties influencing d excess, such as surface conditions at the evaporative source. In both cases, these deficiencies reduce the confidence in using GCMs to interpret d excess or ¹⁷O excess glacial-interglacial variations in polar ice cores. Consequently, quantitative interpretations of variations in ice d excess are still mainly based on simple Rayleigh distillation models [e.g., Johnsen et al., 1989; Ciais and Jouzel, 1994; Ciais et al., 1995; Kavanaugh and Cuffey, 2003; Stenni et al., 2001; Masson-Delmotte et al., 2005]. Finally, results from GCMs including $H_2^{1/O}$ have not, to our knowledge, been published yet.

[10] To explore processes at work during the transport of air masses from the source regions to the poles, we thus use a Rayleigh type distillation model: MCIM (Mixed Cloud Isotopic Model) [*Ciais and Jouzel*, 1994]. This model permits to quantify the water isotopic fractionation along the air mass trajectory as well as to explore the effect of evaporative recharge over midlatitude and high-latitude oceans (section 4.2).

[11] To represent the influence of source regions on water isotopic composition, most Rayleigh distillation models are initialized by the isotopic composition R_{BL} of an air parcel originating from the planetary boundary layer (BL), and the influences of changing climatic conditions on this isotopic composition are estimated through the so-called closure assumption [*Merlivat and Jouzel*, 1979] (auxiliary material Text S1, section 1)¹:

$$R_{\rm BL} = \frac{R_{\rm oce}}{\alpha_{\rm eq} \cdot (\alpha_K + rh_s \cdot (1 - \alpha_K))},\tag{1}$$

with α_{eq} and α_K the equilibrium and kinetic fractionation coefficients, R_{oce} the isotopic ratios of the ocean surface and rh_s the relative humidity at the surface, i.e., the relative humidity of near-surface air at the temperature of the ocean surface T_s :

$$\operatorname{rh}_{s} = \operatorname{rh}_{a} \cdot \frac{q_{\operatorname{sat}}(T_{a})}{q_{\operatorname{sat}}(T_{s})},$$
 (2)

where q_{sat} is the specific humidity at saturation and rh_a and T_a are the relative humidity and temperature of the near-surface air, respectively.

[12] The closure assumption relies on the hypothesis that the only source of vapor in the BL is from the surface evaporation (Figure 1a). However, progress in tropical meteorology, in our understanding of the physical processes that control the distribution of water stable isotopes in the atmosphere, and in isotopic modeling, makes the applicability of the closure assumption increasingly questionable [e.g., *Jouzel*

¹Auxiliary materials are available in the HTML. doi:10.1029/2008JD011535.

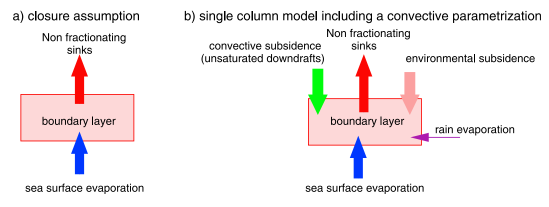


Figure 1. Scheme illustrating the moisture balance of the boundary layer (BL): (a) in the closure assumption [*Merlivat and Jouzel*, 1979] and (b) in the single-column model. Whereas the BL vapor in the closure assumption only originates from sea surface evaporation, the BL vapor in the single-column model receives water vapor from (1) sea surface evaporation (blue), (2) vapor just above the BL, entering the BL through subsiding motions (magenta), (3) unsaturated downdrafts, driven by the rain evaporation (green), and (4) rain reevaporation (purple). In both models the sinks of the BL vapor do not fractionate (e.g., updrafts).

and Koster, 1996; Delmotte et al., 2000]. Besides, in the tropics the BL air is not only influenced by sea surface evaporation but also by convective activity and the large-scale atmospheric circulation [e.g., *Raymond*, 1995]. Observations show that convective processes substantially affect the isotopic composition of the BL vapor [*Lawrence et al.*, 2004]. Large-scale subsidence and boundary layer mixing can also be important in dry regions [*Angert et al.*, 2008]. These reasons may explain why GCMs predict δ^{18} O (d excess) values that can differ from the closure assumption predictions by 3‰ (2‰) in the Tropics and by up to 6‰ (8‰) in midlatitudes [*Jouzel and Koster*, 1996].

[13] Therefore, to explore the impact on ¹⁷O excess of a broader range of processes than those from Landais et al. [2008], we use, instead of the closure assumption, a singlecolumn model (SCM) whose physics package incorporates the essential controls of water vapor and water stable isotopes in the tropics and subtropics [Bony and Emanuel, 2001]. In particular, the SCM includes the Emanuel convective parametrization [Emanuel, 1991], which represents in detail some convective processes such as rain reevaporation and allows for a detailed representation of the isotopic fractionation during these processes [Bony et al., 2008] (Appendix A). The representation by this SCM of cumulus convection and of isotopic processes has been carefully evaluated using tropical data [Bony and Emanuel, 2001; Bony et al., 2008]. As illustrated in Figure 1b, in contrast with the closure assumption, the SCM simulates the effect of both large-scale and convective subsidence on the isotopic composition of the BL.

[14] According to GCM studies [*Delaygue et al.*, 2000; *Werner et al.*, 2001], about 15% of the East Antarctic precipitation originates from moisture evaporated in the tropics, 30% from the subtropics (30°S–40°S) and 50% from midlatitude and high-latitude oceans. The SCM permits to explore the influence of processes (convective processes and large-scale motions in particular) occurring over the Tropics and the subtropics. The 50% of the Vostok precipitation that originates from evaporation in mid and high latitude can be considered as evaporative recharge of air masses during their poleward transport [*Hendricks et al.*, 2000; *Noone*, 2008]. In this paper, we take this recharge into account by adding it to the Rayleigh distillation model. Therefore, the combined analysis of the SCM and MCIM models allows us to consider the main processes that are likely to affect the isotopic composition of the polar snowfall: conditions at the evaporative source (surface conditions, convection), Rayleigh distillation and evaporative recharge over mid and high latitudes. Note that the condensation in frontal clouds is represented in a simple and implicit way by the Rayleigh distillation of MCIM. This latter does not allow to represent the impact, on the isotopic composition of the water transported poleward, of changes that might occur in the physical or microphysical properties of frontal cloud systems during climate change.

3. Single-Column Simulation of the Isotopic Composition of the Vapor Evaporated From the Tropics and Subtropics

[15] The single-column model and its isotopic implementation were extensively described by *Bony and Emanuel* [2001] and *Bony et al.* [2008], and are summarized in Appendix A. Each of our SCM simulation represents the steady state of the atmosphere in radiative-convective equilibrium, determined by the boundary conditions: surface temperature and wind speed, large-scale profile of vertical velocity, radiation, CO_2 . The model is ran over tropical and subtropical oceanic conditions.

[16] For example, for a wind speed of 5 m/s, sea surface temperature of 25°C and no large-scale ascent or descent (corresponding to average tropical conditions), the precipitation rate is about 3.6 mm/d and the simulated isotopic composition of the BL vapor is -12.9%, 14.7‰ and 20 per meg for δ^{18} O, d excess and ¹⁷O excess, respectively. These values compare well with measurements over tropical ocean: δ^{18} O $\simeq -10\%$ to -13% [Lawrence et al., 2004]. Very few data are available for d excess and ¹⁷O excess in tropical oceanic vapor. For comparison, unpublished results over the Amazon are in the ranges of 10–20‰ for d excess and 15– 20 per meg for ¹⁷O excess.

170¹⁷O ^{17}O $\delta^{18}O$ $\delta^{18}O$ $\delta^{18} {\rm O}$ if Excess d if Excess d in Excess Sensitivity SST р in BL in BL in BL Closure Closure Vif Closure rh. rh. in Ice Ice in Ice 61 (hPa/d) Test (m/s) $(^{\circ}C)$ (mm/d)(‰) (‰) (per meg) (‰) (‰) (per meg) (%) (%) (‰) (‰) per meg) 71.2 -54.2Control 5 25 15 1.4 -12.114.4 19.3 -10.512.9 16.2 78.8 15 42 5 21 15 2.3 -10.780.7 74.6 -51.98.3 34 SST -11.911.3 12.4 10.2 12.9 -11.6 SST 5 29 15 18.9 -10.172.1 40 2.615.2 13.6 16.0 79.9 -53.317.5 2 V_s 25 15 1.3 -13.320.9 24.8 -11.318.7 27.9 69.3 57.6 -53.420.2 49 V3 25 15 1.7 -12.9 18.2 23.7 -11.016.1 22.9 73.2 63.5 -53.5 19.1 48 V_{s} 6.5 25 15 2.6-11.612.5 153 -10.3112 12.7 80.7 75.1 -53.013.6 39 5 25 30 0.5 8.2 -10.211.31 31 -10.510.9 10.5 81.8 76.8 -52.98.3 $\omega_{\rm max}$ 5 25 0 -12.9147 198 -10.6134 17.270.6 157 44 36 777 -533 $\omega_{\rm max}$ 5 25 -306.9 -15.219.8 27.0 -10.915.5 21.5 73.8 65.0 -54.921.7 50 $\omega_{\rm max}$ 5 25 -60 9.8 -16.722.3 28.1 -11.016.4 23.4 71.9 62.8 -56.127.2 52 ω_{\max} 25 $\omega_{\rm max}$ 5 -9012.4-17.9249 28.9 -11.1174254 70.0 60.5 -56.831 5 53

 Table 1. Isotopic Composition of the Source Vapor and Vostok Ice Simulated by the SCM and MCIM Depending on Boundary at the Vapor Source^a

^aBoundary conditions imposed to the SCM: surface wind speed V_s , sea surface temperature SST, and large-scale circulation ω (the large-scale vertical velocity at 500 hPa; negative values indicates large-scale ascent). Also given are precipitation rate *P*, isotopic composition of the boundary layer (BL) (δ^{18} O, d excess *d*, and ¹⁷O excess), isotopic composition of the BL if it was predicted by the closure assumption [*Merlivat and Jouzel*, 1979], relative humidity of the air at the lowest-level rh_a and relative humidity at the surface rh_s , simulated at equilibrium by the single-column model. The humidity rh_s is calculated using equation (2). Finally, isotopic composition of the ice simulated by the MCIM (see section 5.1) for the different initial conditions given in the previous columns are shown in columns δ^{18} O, *d*, and ¹⁷O excess. Control entries are for the control simulation described in section 4.1.

[17] Compared to the closure assumption, the SCM yields δ^{18} O 2.3‰ lower, d excess 1.3‰ higher and ¹⁷O excess 3 per meg higher (Table 1), in agreement with GCM outputs [*Jouzel and Koster*, 1996]. In the SCM, this is due to the admixture into the BL of depleted and high d excess vapor by the unsaturated downdraft (Figures 1 and 2). Therefore, considering convective downdrafts in the SCM leads, in

regions of significant convection, to a significant modification of the BL composition compared to the closure assumption.

3.1. Sensitivity to Boundary Conditions

[18] To investigate what controls the BL composition in the SCM, we perform sensitivity tests to large-scale boundary

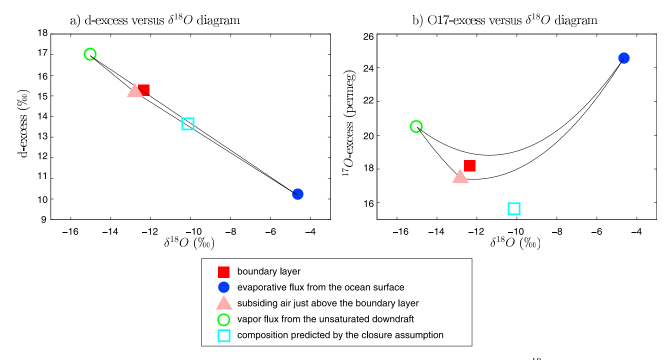


Figure 2. Isotopic composition of the BL simulated by the model (red solid square) in δ^{18} O versus (a) d excess and (b) ¹⁷O excess diagrams. The composition of the unsaturated downdraft vapor mass flux into the boundary layer, the evaporative flux from the ocean, and the subsiding environment just above the boundary layer are shown with green open circles, blue solid circles, and magenta solid triangles, respectively. Mixing lines (lines representing the ensemble of possible mixtures between two samples) are in black. The isotopic composition of the boundary layer predicted by closure assumption for the simulated surface conditions is shown with cyan open squares.

conditions: sea surface temperature (SST), surface wind speed and large-scale vertical velocity at 500 hPa (ω , controlling the convective intensity). Note that the surface relative humidity (rh_s) is not a boundary condition in the model, but rather a prognostic variable that depends mainly on the surface wind speed and on the large-scale convective activity.

[19] All sensitivity tests are performed around a control simulation corresponding to subtropical/tropical conditions (45% of the moisture source according to GCMs), with an SST of 25°C, surface wind speed of 5 m/s and a large-scale descent of 15 hPa/d at 500 hPa. We vary the boundary conditions of the SCM within a range for which the neglect of horizontal advections constitutes a reasonable approximation: SST from 21°C to 24°C, wind speed from 2 to 6.5 m/s and vertical velocity from –90 to +30 hPa/d.

[20] Note that we test independently the sensitivity to SST, surface wind and large-scale velocity to better isolate their effects, though in nature these variables vary in concert [Sobel and Bretherton, 2000; Bony et al., 2004].

3.1.1. Sensitivity to SST

[21] We test SST variations of up to 4°C, which is an upper bound of SST glacial-interglacial variations at low latitudes [*Harrison*, 2000; *Barrows and Juggins*, 2005; *Waelbroeck et al.*, 2009]. The sensitivity to SST simulated by the SCM is of 0.08 ‰/°C for δ^{18} O and 0.3‰/°C in d excess (Figure 3a, red solid line and Table 1). This sensitivity is predicted by the closure assumption (Figure 3a, dotted green line) and is due mainly to the variation of fractionation coefficients with temperature.

[22] The BL ¹⁷O excess is not expected to change with temperature [*Barkan and Luz*, 2005]. Indeed, $\frac{\ln(^{17}\alpha_{eq})}{\ln(^{18}\alpha_{eq})}$ remains equal to 0.529 for temperature variations between 10°C and 40°C (with ¹⁷ α_{eq} and ¹⁸ α_{eq} the fractionation coefficients at liquid-vapor equilibrium for H₂¹⁷O/H₂¹⁶O and H₂¹⁸O/H₂¹⁶O). The 7 per meg increase in ¹⁷O excess from 21°C to 25°C (Figure 3a) is only due to the 3% decrease in *rh_s* (driving a 3 per meg increase in ¹⁷O excess) and the 1 mm/d increase in precipitation (driving a 4 per meg increase in ¹⁷O excess). These sensitivities to *rh_s* and precipitation will be explained in sections 3.1.2 and 3.1.3.

3.1.2. Sensitivity to Surface Wind Speed

[23] We vary the surface wind speed V_s along a range of values typically found over tropical oceans from 2 to 6.5 m/s. We only test values in the smooth regime below 7 m/s (where the kinetic fractionation is constant), since 95% of the ocean surface is in this regime [*Eriksson and Bolin*, 1964]. Besides, V_s greater than 7 m/s would imply a large change in the kinetic fractionation coefficient at the transition between smooth and

rough regime [*Merlivat and Jouzel*, 1979], whose physical significance over the ocean remains unclear.

[24] As V_s increases, δ^{18} O increases (0.9 ‰/(m/s)) and both d excess and ¹⁷O excess decrease (-2.4‰/(m/s) and -4.4 per meg/(m/s), respectively: Figure 3b, solid red line). The sensitivity of the isotopic composition to V_s is mainly explained by coincident rh_s variations: when V_s increases, surface evaporation is enhanced, the water content in the BL increases and rh_s increases, by about 4.6%/(m/s) (Figure 4b). To quantify the effect of rh_s , we performed the same simulations but with rh_s artificially fixed constant at 70% in the calculation of the isotopic composition of the evaporation flux. Without the coincident variations in rh_s , the sensitivity of the isotopic composition to V_s becomes very small (Figure 4b, dotted green line), confirming that the underlying factor explaining this sensitivity is rh_s .

3.1.3. Sensitivity to Convective Intensity

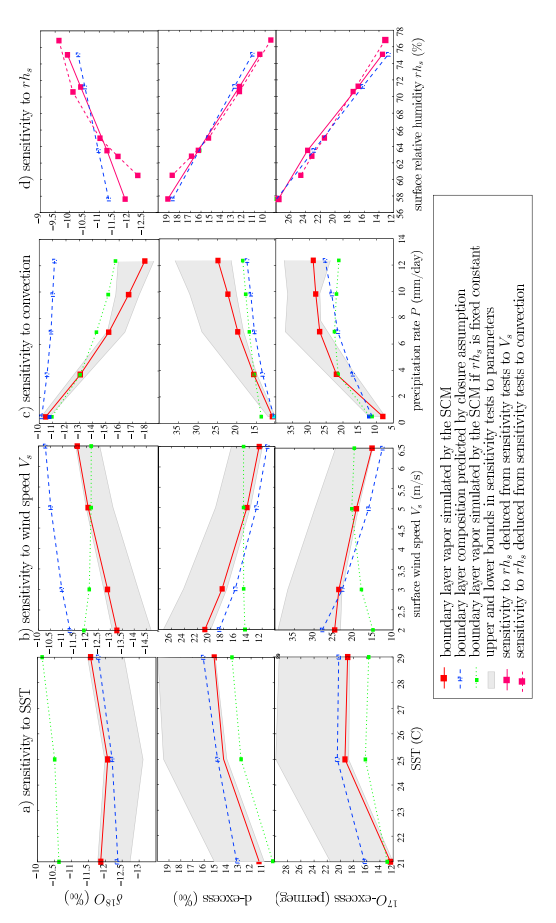
[25] Convective intensity is modulated in the SCM by prescribing a large-scale ascent or descent within the troposphere, and measured by the precipitation rate *P*.

[26] BL vapor δ^{18} O decreases by about 0.4 ‰/(mm/d) as convective intensity increases (Figure 3d), consistent with the well-known tropical amount effect observed in the precipitation [*Dansgaard*, 1964; *Rozanski et al.*, 1993]. This compares well with the measurements of *Lawrence et al.* [2004] over tropical ocean ranging from -10% in quiescent weather to below -20% after intense convective systems. Meanwhile, d excess increases by about 0.6‰/(mm/d) as convective intensity increases (Figure 3c, solid red line).

[27] Convective processes decrease δ^{18} O and increase d excess in the BL vapor mainly through convective mixing bringing down depleted and high d excess vapor [*Risi et al.*, 2008]. However, as for the sensitivity to V_s , part of the sensitivity of d excess to convection is explained by coincident rh_s variations: convection dries the BL in the model, owing to stronger upward transport of humidity by convective fluxes. Sensitivity tests to convective intensity in which rh_s is artificially held constant at 70% in the calculation of the composition of the surface evaporation (Figure 3c, dotted green line) show that the effect of rh_s contributes for 55% to the sensitivity to convective processes (Figure 3c, dashed blue line).

[28] The ¹⁷O excess increases with increasing convective activity by about 0.8 per meg/(mm/d) as precipitation increases from 4 mm/d to 12 mm/d (Figure 3c). For these precipitation rates, the increase in ¹⁷O excess is mainly due to the decrease in rh_s : without coincident rh_s variations, the BL ¹⁷O excess varies by less than 0.1 per meg/(mm/d) (dotted green line).

Figure 3. Thick red line denotes sensitivity of boundary layer δ^{18} O, d excess, and 17 O excess to (a) sea surface temperature (SST), (b) surface wind speed, and (c) convective intensity, measured here as precipitation rate. Dashed blue line denotes sensitivity predicted by closure assumption for the same SST and the same relative humidity at the surface. Dotted green line denotes sensitivity simulated by the model when the relative humidity at the surface is held artificially constant at 70% in the calculation of the composition of surface evaporation, highlighting the direct effect of convective processes on the BL composition (sections 3.1.2 and 3.1.3). The gray shading represents the envelope of the simulated boundary layer compositions (δ^{18} O, d excess, and 17 O excess) when model parameters related to unsaturated downdraft are varied by $\pm 20\%$ and when parameter ϕ is varied from 0 to 0.9. (d) Sensitivity to rh_s , deduced from the sensitivity tests, to V_s (solid pink line) and convective activity (dashed pink line); the simulations with "constant" rh_s have been subtracted from the standard simulations to isolate the effect of rh_s . Also shown is the closure assumption (dashed blue line).





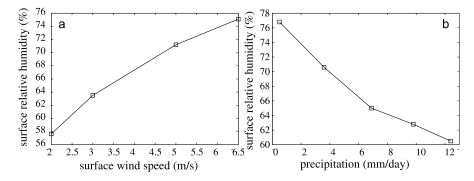


Figure 4. Sensitivity of surface relative humidity rh_s to (a) surface wind speed V_s and (b) precipitation in the SCM simulations. In Figure 4a, SST = 25°C, $\omega_{max} = 15hPa/d$, and V_s is varied from 2 to 6.5 m/s. In Figure 4b, SST = 25°C, $V_s = 5m/s$, and ω_{max} is varied from -90 to 30hPa/d.

The sensitivity of ¹⁷O excess to convective activity is higher for low precipitation rate: 4 per meg/(mm/d) for precipitation rates lower than 4 mm/d. Variations in rh_s account for 40% of this sensitivity for these precipitation rates.

3.2. Main Drivers of ¹⁷O Excess Changes in the Source Regions

[29] The previous experiments have revealed that convective processes and changes in rh_s were the two main processes underlying the sensitivities of ¹⁷O excess to boundary conditions.

[30] To calculate an upper bound for the impact of convective processes on ¹⁷O excess, we assume that the precipitation rate varies from 1 mm/d at LGM to 4 mm/d at EH in tropical and subtropical regions contributing to 50% of the Vostok precipitation. This is an extreme assumption since (1) GCMs rarely simulate larger precipitation changes, even regionally [*Braconnot et al.*, 2007], (2) tropical and subtropical regions are not expected to contribute more than 50% to the Vostok precipitation [*Delaygue et al.*, 2000] and (3) the sensitivity to convection is maximum at low precipitation rates. Excluding the effect of coincident rh_s variations, this would lead to a change of only 4 per meg in the Vostok precipitation, thus explaining 20% of the observed ¹⁷O excess change. Convection is therefore a minor influence on ice ¹⁷O excess.

[31] Sensitivity experiments to V_s and convection have revealed the large influence of rh_s on ¹⁷O excess. By comparing the sensitivity to V_s with and without fixed rh_s (Figure 3d, solid pink line), we estimate the sensitivity of ¹⁷O excess to rh_s to -1.0 per meg/%. Similarly, the δ^{18} O and d excess sensitivities to rh_s are 0.2‰/% and -0.5‰/%.

[32] These sensitivities to rh_s are roughly linear (Figure 3d). and consistent both with the closure assumption (dotted green line) and with the sensitivities deduced from the response to convection (dashed pink line), supporting the robustness of these sensitivities to rh_s . We thus suggest that these sensitivities to rh_s are robust and can be applied in extratropical conditions as well, as suggested also by d excess and ¹⁷O excess observations over midlatitude and high-latitude Southern Ocean [*Uemura et al.*, 2008, 2010].

[33] Taking into account uncertainties in the ¹⁷O excess sensitivity to rh_s (auxiliary material Text S1, section 2), a 12 to 22% decrease in rh_s from LGM to EH may explain

the 20 per meg change in BL 17 O excess, a conclusion similar to that of *Landais et al.* [2008].

4. Poleward Transport Processes

[34] To understand how 17 O excess variations in the source regions are transmitted to d excess and ¹⁷O excess in the polar regions, we initialize the MCIM with the BL isotopic composition simulated by the SCM (section 3) and simulate the isotopic composition of the snowfall over the Vostok station (-55°C, 3500 m above sea level [Petit et al., 1999]). Table 1 displays δ^{18} O, d excess and 17 O excess simulated in the ice at the Vostok location. The δ^{18} O, d excess and 17 O excess simulated by the model (-54.2%), 17.5% and 40 per meg, respectively) compare reasonably well with the measurements in Vostok for the present day (about -55‰, 16‰ and 40 per meg, respectively, for annual averages). Nevertheless, an accurate simulation of the Vostok snowfall composition is not expected given the idealized framework of both the SCM and MCIM. We thus focus here on the sensitivity to climatic conditions rather than the absolute value of the isotopic composition.

4.1. Effect of Pure Rayleigh Distillation

[35] The sensitivities of d excess and ¹⁷O excess in the source BL to source conditions is also visible in the polar snowfall isotopic composition (Table 1). For example, when increasing rh_s from 69 to 81%, d excess and ¹⁷O excess decrease both in the source BL and in the polar snowfall. However, distillation processes induce a dampening of the initial vapor d excess anomaly: while ¹⁷O excess decreases by 10 per meg both in the source vapor and in the snowfall, d excess decreases by 8.4‰ in the source vapor and only by 6.6‰ in the surface snow. Here are two reasons for these different behaviors of d excess and ¹⁷O excess along the distillation process.

[36] 1. A change in δ^{18} O of the source vapor influences d excess in the snowfall even if the source d excess remains constant [e.g., *Delaygue*, 2000; *Cuffey and Vimeux*, 2001; *Werner et al.*, 2001; *Vimeux et al.*, 2002; *Jouzel*, 2003]. On the contrary, ¹⁷O excess is not affected by such a change in δ^{18} O of the source region (Table 2, last two rows), due to the different definitions of d excess (linear) and ¹⁷O excess (logarithm) (auxiliary material Text S1, section 3).

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Table 2. Isotopic Composition of the Ice Simulated for Different Source and Site Temperatures and Different δ^{18} O of the Boundary Layer^a

T _{source} (°C)	T _{site} (Inversion) (°C)	Vapor δ^{18} O (‰)	$\begin{matrix} \text{Ice} \\ \delta^{18} \text{O} \\ (\%) \end{matrix}$	Ice d Excess (‰)	Ice ¹⁷ O Excess (‰)
26	-35	-13	-55.6	27.5	42
24	-35	-13	-54.5	24.5	42
28	-35	-13	-56.8	30.6	43
26	-38	-13	-59.6	35	43
26	-32	-13	-51.6	21	42
26	-35	-10	-52.7	20	42
26	-35	-16	-58.5	35	42

 $^{\mathrm{a}}\mathrm{In}$ the boundary layer, d excess is 15‰ and $^{17}\mathrm{O}$ excess is 20 per meg.

[37] 2. The d excess is influenced by air temperature while ¹⁷O excess is almost insensitive to temperature along the distillation (Table 2, first three rows), due to different evolutions of the equilibrium fractionation coefficients with temperature. Indeed, while $\frac{\ln(1^{17}\alpha_{eq})}{\ln(1^{18}\alpha_{eq})}$ remains equal to 0.529 for temperature increasing from 0°C to 20°C, $\frac{D\alpha_{eq}-1}{18\alpha_{eq}-1}$ varies between 9.6 and 8.7 ($^{D}\alpha_{eq}$ being the fractionation coefficient at liquid-vapor equilibrium for HD¹⁶O/H2⁶O). Note that a similar dependency on temperature is observed for $\delta D/\delta^{18}O$ if we consider the evolution of $\frac{\ln(D\alpha_{eq})}{\ln(1^{18}\alpha_{eq})}$ (9.1 at 0°C and 8.4 at 20°C) and the same behavior holds for vapor-solid fractionation. Therefore, the near-surface temperature in the moisture source regions influences the polar snowfall d excess [*Petit et al.*, 1991; *Vimeux et al.*, 2002] but not ¹⁷O excess.

[38] Consequently, the ¹⁷O excess variations of the BL vapor in the source region are perfectly preserved in the polar precipitation (Table 2). On the contrary, the polar d excess is sensitive to the initial and final temperatures of the distillation (Table 2), and thus only partially reflects the d excess signal from the source region.

4.2. Effect of Evaporative Recharge

[39] Classical Rayleigh distillation models assume that air parcels are isolated. However, in nature, air masses are partially recharged through surface evaporation: for example, *Trenberth* [1998] estimated a recycling ratio (proportion of the moisture originating from the local evaporation versus horizontal advection) from 10 to 20% at the 1000 km scale over subtropical and midlatitude oceans. The isotopic composition of Antarctic snowfall is affected by the evaporative recharge [*Kavanaugh and Cuffey*, 2003; *Lee et al.*, 2008] and the competition between mixing of surface evaporated water through the BL and poleward advection of moisture [*Noone*, 2008]. Evaporative recharge is expected to affect even more strongly ¹⁷O excess than δ^{18} O or d excess because mixing lines are curved in a logarithmic plot (Figure 5a), so that ¹⁷O excess is not preserved during mixing.

[40] To quantify the effect of evaporative recharge through the mixing of air masses with contrasted δ^{18} O, we perform an idealized sensitivity experiment. We start a distillation of the initial vapor (δ^{18} O = -12 ‰, d excess = 15‰, ¹⁷O excess = 18 per meg) until the vapor reaches a 2‰ δ^{18} O depletion. Then, we recharge 10% of the vapor by an evaporative source, corresponding to the composition of the evaporation flux simulated by the SCM in the control simulation: δ^{18} O = -5% (consistent with *Lee et al.* [2007]), d excess = 10‰, ¹⁷O excess = 25 per meg (Figure 2). The same distillationrecharge steps are repeated again until the precipitation δ^{18} O corresponds to the δ^{18} O measured at the coast of Antarctica. Then, the parcel is distilled without recharge until reaching the Vostok station. This experiment is roughly equivalent to a recycling ratio of 20% at the 1000 km scale (consistent with *Trenberth* [1998], until 70°S in latitude. Compared to an isolated parcel, ¹⁷O excess in Vostok snowfall is lower by 14 per meg, due to the curvature of the mixing lines (Figure 5). The Vostok snowfall d excess is just slightly reduced by 4‰ due to the admixture of lower d excess vapor by surface evaporation.

[41] Note that recycling in this test is artificial and does not accurately represent the complexity of recharge processes. Also, this test estimates an upper bound of the effect by mixing the distilled air mass with a very contrasted δ^{18} O. If the vapor was recharged with a less contrasted vapor (δ^{18} O = -12%) instead of -5%, for example, recharge by vertical mixing with the BL [*Noone*, 2008], the Vostok ¹⁷O excess would be lower by 10 per meg instead of 14 per meg. The recycling ratio used is also an upper bound, since air masses in mid and high latitudes are not transported poleward directly in contact with the ocean surface [*Noone*, 2008].

[42] Finally, we find a maximum sensitivity of ¹⁷O excess to the evaporative recycling ratio of -0.7 to -0.5 per meg/%. A decrease of the recycling ratio from LGM to EH by 30% to 40% (i.e., recycling ratio from 40–60% to 10–20%) would thus be required to explain by itself the 20 per meg ¹⁷O excess increase. For the LGM to present-day change, PMIP2 models [*Braconnot et al.*, 2007] simulate a slight decrease (smaller than 5%) of the recycling ratio between 20°S and 60°S from LGM to present day. This process is thus not likely to contribute to more than 20% to the observed ¹⁷O excess increase from LGM to EH.

4.3. Effect of δ^{18} O Seasonality in Antarctica

[43] Due to the curvature of the mixing lines in the loglog diagram, the annually averaged ¹⁷O excess in precipi-tation depends on the amplitude of the δ^{18} O seasonal cycle (Figure 5b): if the seasonal amplitude of δ^{18} O in snowfall increases, then the annually averaged ¹⁷O excess decreases. As an example, we assume that at present day, 40% of the Vostok snow accumulation occurs from October to March with a δ^{18} O of -50‰ and that the remaining 60% of the accumulation occurs from April to September with a δ^{18} O of -60% (values of the same magnitude as reported by *Ekaykin* [2003]). For simplicity, we assume a constant ¹⁷O excess in the snowfall all over the year. If the seasonal amplitude of δ^{18} O in Vostok snowfall increases from 10‰ today to 20% during the LGM, then the annual ¹⁷O excess in Antarctica snowfall decreases by 10 per meg (Figure 5b). Since mixing lines are linear when dealing with d excess, this last effect does not affect the d excess.

[44] No data describing δ^{18} O seasonality in Antarctica are available during the LGM. Some isotopic GCM show either no change [*Jouzel et al.*, 1994] or a small change (20% in simulations presented by *Jouzel et al.* [2007b]) in the seasonal cycle in Eastern Antarctica between the LGM and today. PMIP2 models simulate a decrease of the seasonal amplitude of temperature from LGM to present day, ranging from about 0°C to 6°C over Antarctica depending on

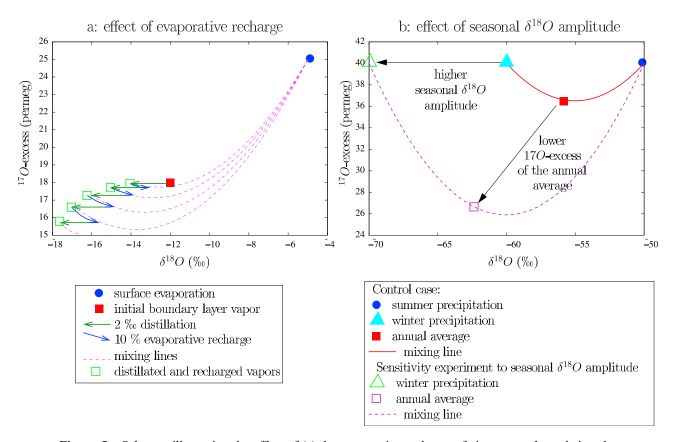


Figure 5. Schemes illustrating the effect of (a) the evaporative recharge of air masses along their poleward transport and (b) the seasonal amplitude of snow δ^{18} O over Antarctica. Both effects are due to the curvature of the mixing lines in the ¹⁷O excess versus δ^{18} O diagram, because of the logarithmic definition of ¹⁷O excess. In Figure 5a, in an idealized sensitivity experiment to evaporative recharge described in section 4.2, an initial boundary layer vapor (solid red square) undergoes successive distillation and recharge steps by a surface evaporation flux (blue circle). Both the initial vapor and surface evaporation have the composition illustrated on Figure 2. As the distillation depletes the vapor without significantly changing its ¹⁷O excess, evaporative recharge decreases ¹⁷O excess due to the curvature of the mixing lines. In Figure 5b, in an idealized sensitivity experiment, we assume that at the present day, 40% of the Vostok snow accumulation occurs from October to March with a δ^{18} O of -50‰ and the remaining 60% of the accumulation occurs from April to September with a δ^{18} O seasonal cycle in Antarctic snow is increased from 10‰ to 20‰, then the annual ¹⁷O excess of the snow decreases by about 10 per meg, due to the curvature of the mixing line.

models. We thus propose here a rough estimate based on temperature seasonality derived from GCM simulations of the LGM, and applying the modern seasonal slope of $\frac{d\delta^{18}O}{dT}$ 0 ~ 4‰/K [*Ekaykin and Lipenkov*, 2008]. As an upper bound for this effect, considering a decreased seasonal amplitude of 6°C from LGM to EH, the δ^{18} O seasonal amplitude in polar snowfall would decrease by 2.4‰ from LGM to EH. The consequence is a small increase of snowfall ¹⁷O excess of about 2.4 per meg from LGM to EH.

[45] Given these results, the variation in the seasonality of δ^{18} O is thus unlikely to contribute for more than 15% to the observed ¹⁷O excess shift.

5. Paleoclimatic Implications

[46] In sections 3 and 4, we have explored different processes affecting the source ¹⁷O excess, its evolution along air mass distillation and the impact of precipitation seasonality. While convective changes, recharge and seasonality may account for part of the ¹⁷O excess shift between the LGM and EH, these influences, even combined, are not likely account for more than 50% of the shift. On the other hand, a higher rh_s at LGM by 12 to 20%, would explain the observed shift (as in work by *Landais et al.* [2008]). This is an upper bound, neglecting the role of the aforementioned processes that have a secondary impact. We discuss here the realism of an extreme 20% change in rh_s and its implications for LGM temperature reconstructions using stable water isotopes in ice cores.

5.1. Realism of a Higher rh_s Over Evaporative Regions During the LGM

[47] As mentioned in section 1, GCMs outputs do not produce significant changes of rh_a between the LGM and the present day [*Bush and Philander*, 1999] thus questioning the interpretation of the ¹⁷O excess shift. However, variations

in rh_s are not equal to variations in rh_a. The rh_s differs from rh_a owing to the difference between the air and surface temperatures (thermodynamic disequilibrium) [Angert et al., 2008; Pfahl and Wernli, 2008]. By differentiating equation (2) as a function of rh_a, T_s and T_s-T_a , and quantifying the different terms for rh_a ranging from 60 to 80%, T_s from 0°C to 15°C (conditions over midlatitude oceans, which are important sources of the Vostok precipitation) and T_s-T_a from -3°C to 3°C, we express the variations of rh_s as

$$\Delta rh_s = (1.0 \pm 0.2) \cdot \Delta rh_a - (2 \pm 11) \cdot 10^{-2} \cdot \Delta T_s - (4.7 \pm 0.7) \cdot \Delta (T_s - T_a).$$
(3)

[48] Variations in rh_s can thus arise either from a variation of rh_a of the same order of magnitude (first term on the right-hand side) or a variation in the thermodynamical disequilibrium between the sea surface and the near-surface air T_s-T_a (third term). The second term is negligible (ΔT_s of 10°C yields rh_s variations lower than 1%).

[49] A first possibility to explain the rh_s decrease from LGM to EH is a decrease of rh_a over the source region. This can be obtained through a general rh_a decrease over the Indian and Southern oceans or through a shift of the location of the moisture sources toward regions of lower relative humidity. On the one hand, PMIP2 simulations show variations lower than 3% between 20°S and 60°S. Moreover, GCMs simulate a poleward shift of the source regions from LGM to EH [*Delaygue et al.*, 2000; *Werner et al.*, 2001], which is not consistent with a decrease in rh_a from LGM to EH, since rh_a increases poleward [*Vimeux et al.*, 2001].

[50] On the other hand, Jouzel et al. [1982] argued that a higher rh_a during the LGM could be consistent with higher wind speeds over the ocean during this period, as suggested by highest aerosol and sea salt content measured in Antarctic ice [Petit et al., 1981, 1999]. In addition, global warming during the deglaciation are associated with a change in the intensity, frequency and latitudinal position of the storm tracks [Laîne et al., 2008; Toggweiler et al., 2006] and with a southward shift of the Intertropical Convergence Zone during the LGM [Chiang and Bitz, 2005; Kang et al., 2008]. Such large reorganizations could also affect the latitudinal distribution of relative humidity at low levels.

[51] A second possibility to explain the variations of rh_s is a change in the thermodynamical disequilibrium between the sea surface and the near-surface air in the source regions (equation (3)). For example, the 20% higher rh_s during the LGM could be explained by a 4°C lower $T_s - T_a$ (i.e., T_s increase from LGM to EH 4°C more strongly than T_a). However, LGM simulations with coupled ocean-atmosphere models conducted in PMIP2 [*Braconnot et al.*, 2007] do not show such a disequilibrium. They rather simulate a slightly lower rh_s (decrease of 3% in average between 30°S and 60°S) during the LGM.

[52] There is thus a mismatch between the 20% change in rh_s suggested by the ¹⁷O excess shift in polar ice and the small variations in rh_s simulated by GCMs.

5.2. Implications of Relative Humidity Changes on LGM Temperatures Reconstructions

[53] If true, a large change in surface relative humidity at the source may have strong consequences for the classical interpretation of ice δ^{18} O and d excess as indicators of site and source temperatures T_{site} and T_{source} [Vimeux et al., 2002; Jouzel et al., 2007a; Masson-Delmotte et al., 2005; Stenni et al., 2001]. Indeed, δ^{18} O and d excess in polar snowfall also depend on the variations in rh_s in the source regions. Until now, variations in rh_s were assumed to equal variations in rh_a, and variations in rh_a were either neglected or assumed to vary with T_{source} by -0.38%/°C [Vimeux et al., 2002]. This slope was obtained from spatial correlation between SST (between 7°C and 24°C) and rh_a from GCM outputs for present-day simulations [Vimeux et al., 2002]. However, these assumptions are questionable: (1) rh_s can vary even though rh_a is constant and (2) seasonal correlations between SST and rh_a in NCEP data [Kalnay et al., 1996] show weak values and variations in sign, questioning whether the slope given by Vimeux et al. [2002] is robust and holds in time (especially between glacial and interglacial periods). Therefore, changes in rh_s might have been underestimated in previous studies.

[54] In the light of the rh_s variations suggested by the ¹⁷O excess record, we thus reconsider the previous temperature reconstructions in Vostok over the deglaciation based on δ^{18} O and d excess records alone. To do so, we perform an inversion of the δ^{18} O and d excess ice core data at LGM to retrieve simultaneously T_{site} and T_{source} . The system of equations for δ^{18} O and d excess in ice are derived from MCIM simulations initialized by the SCM (Table 1), assuming linear relationships:

$$\Delta \delta^{18} \mathcal{O}_{\text{corr SW}} = -0.30 \cdot \Delta T_{\text{source}} + 1.0 \cdot \Delta T_{\text{site}} + 0.02 \cdot \Delta rh_s \quad (4)$$

$$\Delta d_{\rm corr\,SW} = 1.50 \cdot \Delta T_{\rm source} - 1.1 \cdot \Delta T_{\rm site} - 0.38 \cdot \Delta r h_s \qquad (5)$$

where $\Delta \delta^{18}O_{\text{corr SW}}$ and $\Delta d_{\text{corr SW}}$ are the variations of $\delta^{18}O$ and d excess in the Vostok ice from LGM to EH, corrected by the -1 ‰ change in seawater $\delta^{18}O$ from LGM to EH: $\Delta \delta^{18}O_{\text{corr SW}} = +7\%$ and $\Delta d_{\text{corr SW}} = +1\%$ [Jouzel, 2003].

[55] The coefficients for the sensitivity of Vostok $\Delta \delta^{18}$ O and Δd to ΔT_{source} and ΔT_{site} are given with a 20% uncertainty and are similar to those obtained by *Vimeux et al.* [2002] except for (1) $\frac{\partial d}{\partial T_{hs}}$: -0.15‰/% in work by *Vimeux et al.* [2001] and -0.38‰/% in this study, and (2) $\frac{\partial d}{\partial T_{site}}$: -0.5‰/K in work by *Vimeux et al.* [2002] and -1.1‰/K in this study, due to a different tuning of the MCIM (mainly supersaturation parametrization) based on the additional constraint from ¹⁷O excess [*Landais et al.*, 2008].

[56] Doing the extreme assumption that the increase in ¹⁷O excess from LGM to EH should be attributed to a decrease in rh_s only, we have $\Delta rh_s = -20\%$ and obtain $\Delta T_{site} = +7.8^{\circ}$ C and $\Delta T_{source} = +1.4^{\circ}$ C from LGM to EH. Such a variation of T_{site} over the deglaciation is in fair agreement with the results of *Jouzel et al.* [2003] and the ΔT_{source} is realistic compared to the available estimates of LGM oceanic temperature [*Barrows and Juggins*, 2005; *Sarnthein et al.*, 2003; *Waelbroeck et al.*, 2009]. Surprisingly, we find similar values as in work by *Vimeux et al.* [2002] despite different tunings of the MCIM, because the stronger dependency of d excess to T_{site} in our model is counterbalanced by the strong effect of rh_s .

[57] Taking into account the change in rh_s has a strong influence on the reconstruction of T_{source} (Table 3): for

Table 3. Variations of Site and Source Temperatures Deduced From the Inversion of δ^{18} O and d Excess Changes at Vostok Over the Last Deglaciation, Assuming Different rh_s Variations^a

Δrh_s (%)	$\Delta T_{\rm site}$ (°C)	ΔT_{source} (°C)
-20	+7.8	+1.3
-10	+8.5	+4.4
0	+9.2	+7.4

^a Δ refers to Early Holocene minus Last Glacial Maximum. T_{site} , site temperature; T_{source} , source temperature.

example, assuming a constant rh_s from LGM to EH yields T_{site} and T_{source} changes of +7.4°C and +9.2°C, respectively. The reconstruction of ΔT_{source} is twice more sensitive to the assumed Δrh_s than ΔT_{site} . Estimating the past rh_s variations through ¹⁷O excess measurements would thus have a strong impact on past temperature reconstructions, strengthening the added value of ¹⁷O excess measurements.

[58] This calculation contains many uncertainties: tuning of the MCIM, linearity assumption, uncertainties related to the SCM, uncertainties in the exact change of rh_s if part of the change should be attributed to convective changes, to the evaporative recharge, to the seasonal cycle or to other processes not considered here.

[59] To estimate the uncertainties related to the linearity assumption and the neglect of convection in our simplified equations (e.g., equations (4) and (5)), we performed experiments using rh_s , T_{source} and T_{site} variations given in Table 3. To do so, for each Δrh_s scenario, we perform simulations of the isotopic composition of the Vostok ice for the LGM and EH by the SCM and MCIM (Table 4). The inputs of the SCM and MCIM are such that the change of rh_s , T_{source} and T_{site} between EH and LGM (columns 9, 10 and 12 of Table 4) are almost identical to those in Table 3. Changes in rh_s are obtained in the SCM either by varying V_s or ω (column 3) of Table 4), or both simultaneously when ω variations only are not sufficient to explain rh_s variations. We take EH conditions from the control simulation (except when the change in rh_s was impossible to simulate through reasonable V_s or ω variations), but we focus on the EH-LGM differences rather than on the absolute values. Simulations show that whatever the method to vary rh_s in the SCM, be it through an extreme variation of V_s or of convective activity, the simulated EH-LGM change in ¹⁷O excess (in per meg) corresponds to the change in rh_s (in %), with an error of 2 per meg maximum (last column of Table 4). Taking a slope of 1 per meg/% and neglecting the effect of convective processes on ¹⁷O excess are thus very robust assumptions. Simulated changes in δ^{18} O and d excess in ice are similar to observations with maximum errors of 1.5 and 1.1‰, respectively (columns 16 and 17 of Table 4). This translates into uncertainties of 2.2 and 2.4°C on the reconstruction of ΔT_{site} and ΔT_{source} , respectively.

[60] To estimate the uncertainties related to the models, we perform sensitivity tests with the SCM and the MCIM. We vary tunable parameters in the SCM (auxiliary material Text S1, section 2) and in the MCIM (within ranges of values for which the modeled evolutions of ¹⁷O excess and d excess in Antarctica still agree with the data). In the highest deviation from the inversion presented above, obtained by an extreme tuning of the SCM, the reconstruction of T_{site} is

virtually unchanged (-0.2° C) whereas the reconstruction of T_{source} is more affected (-1.9° C).

[61] Therefore, when considering the uncertainties mentioned above, the maximum uncertainty ranges for T_{site} and T_{source} are on the order of 2°C and 4°C, respectively. However, the major source of uncertainty in this reconstruction remains the estimated change in rh_s, if other factors than rh_s contribute to the observed ¹⁷O excess shift (Table 3): 0.4°C for ΔT_{site} and 9°C for ΔT_{source} .

6. Conclusion

6.1. Summary

[62] We have explored various processes, both at the evaporative source and during the poleward transport, that could explain the +20 per meg increase over the last deglaciation.

[63] Using a single-column model (SCM) over tropical and subtropical oceans, we show that the ¹⁷O excess of the low-level vapor is affected mainly by the relative humidity at the surface, rh_s , with a sensitivity of -1.0 per meg/%. This sensitivity is robust and similar to that predicted by the closure assumption, so that it can likely be extended to all latitudes. Given this sensitivity, a 12 to 22% increase in rh_s would be necessary to explain the +20 per meg increase over the last deglaciation. Changes in rain rates over tropical or subtropical sources, conversely, are not likely to contribute to more than a few per meg to the shift.

[64] Using a Rayleigh type distillation model, we show that the amplitude of ¹⁷O excess variations in the moisture source regions is well recorded in the ¹⁷O excess of the polar precipitation, while they are damped in d excess. The ¹⁷O excess in polar snowfall can also be influenced by evaporative recharge and by the amplitude of the δ^{18} O seasonal cycle at the precipitation site, but the contribution of these effects to the observed ¹⁷O excess shift is expected to remain secondary (at most 35% of the shift).

[65] Among the different processes considered in this study (changes in convective activity, SST or relative humidity at source regions, in evaporative discharge over midlatitude to high-latitude oceans, in the seasonal cycle of Antarctica precipitation), only one can explain the large magnitude (+20 per meg) of the ¹⁷O excess shift observed in Antarctica over the deglaciation: the decrease of the surface relative humidity (rh_s) by 8–22% from LGM to present (in agreement with *Landais et al.* [2008]). This might arise either through a decrease of the surface air relative humidity (rh_a), or through an increase of the thermodynamical disequilibrium between the surface and the near-surface air as the global mean temperature increases.

[66] The fact that current GCMs do not simulate any large change in rh_s during the LGM raises questions. This mismatch could have a link with the inability of current isotopic GCMs to simulate the observed increase of d excess from LGM to EH over polar regions [*Werner et al.*, 2001]: if GCMs simulated a higher rh_s during the LGM, they would more likely simulate a lower d excess, closer to observations.

[67] The possibility of such a large change in rh_s strengthens the interest of ¹⁷O excess to provide more accurate reconstructions of source and site temperatures than from the combination of δ^{18} O and d excess only. The assumption

	Δrh_{z}			T_{zz}	V_{z}	(11)	rh.	Δrh_{e}	ΔT_{zz}	T_{a}	ΔT_{zi}	$\delta^{18}\Omega_i$	d_i	$^{17}O_{c}$	$\Delta \delta^{18}$,	Δd_i	$\Delta^{17}\Omega_i$
Simulation	Scenario	Method	Age	(_C)	(m/s)	(hPa/d)	(%)	(%)	(°C)	(0 ^C)	(°C)	(0%)	(%0)	(per meg)	(0%)	(00)	(per meg)
1	-20	V_{s}	EH	29	2	0	56.0			-55		-55.96	23.1	57			
2	-20	V_s	LGM	27.7	6.9	0	76.3	-20.3	1.3	-62.8	7.8	-61.57	23.2	37	5.61	-0.1	20
Э	-20	ω and V_s	EH	29	4.5	06	61.2			-55		-56.01	22.9	52			
4	-20	ω and V_s	LGM	27.7	6.5	30	80.5	-19.3	1.3	-62.8	7.8	-61.85	21.2	32	5.84	1.7	20
5	-10	V_s	EH	29	3.6	15	65.6			-55		-55.64	18.0	47			
9	-10	V_s	LGM	24.6	6.9	15	75.9	-10.3	4.4	-63.5	8.5	-61.60	16.3	37	5.96	1.7	10
7	-10	3	EH	29	5	-45	66.2			-55		-54.88	21.0	47			
8	-10	Э	LGM	24.6	5	30	76.5	-10.3	4.4	-63.5	8.5	-61.48	19.2	37	6.60	1.8	10
Control	0	V_s	EH	29	5	15	72.5			-55		-54.2	15	42			
6	0	V_s	LGM	21.6	4.5	15	72.3	0.2	7.4	-64.2	9.2	-61.86	14.3	40	7.66	0.7	2
Control	0	3	EH	29	5	15	72.5			-55		-54.2	15	42			
10	0	З	LGM	21.6	5	13	72.1	0.4	7.4	-64.2	9.2	-62.15	14.2	40	7.95	0.8	7

by be in the SCM by two methods (Method): wind speed (V_s) or large-scale vertical velocity at 500 hPa (ω), or both simultaneously when ω variations alone are not For each scenario and method of rhs variation, we perform two simulations, one for EH and one for LGM (Age column), so that the Δrhs is as close as possible to the between EH and on the absolute values. o are given in columns 5, 6, and 7. The rh_s simulated by the SCM as well as the Δrh_s between EH at the polar temperature (T_{si}). The Vostok ice composition (δ^{18} O, d excess, and 17 O excess) simulated t and method of rh_s variation are given in columns 16, 17, and 18. These simulated variations are to the spectively. Note that we focus here on the EH-LGM differences, rather than on the absolute value excess, respectively. Note that we focus $V_{
m s}$, and ω) used to achieve the Δ rh_s scenario are given in columns 5, 6, i for each scenario and scenario. The inputs of the SCM (sea surface temperature T_{ss} , V_s , and ω) used to achieve the Δrh_s scenario are listed in columns 8 and 9. The outputs of the SCM are then used as input for the MCIM, together with tCIM is given in columns 13, 14, and 15, and the corresponding EH-LGM variations for each scenario an 0 and for Vostok ice δ^{18} O, d excess, per meg rh_s +1‰, and +20 temperature (ΔT_{si}). For each scenario, we vary sufficient to explain Δrh_s (simulations 3 and 4). observed +7%o. the compared with Δrh_s scenario. the MCIM LGM 8

that rh_e changes are either negligible or linearly related to temperature [Vimeux et al., 2001; Stenni et al., 2001] can be relaxed. Using ¹⁷O excess measurements to constrain the change in rh_s yields a LGM source temperature about 1.4°C lower than at EH, and a Vostok temperature 7.8°C lower. Both these estimates are consistent with previous studies [Vimeux et al., 2001], but owing to compensating effects. These estimates are particularly sensitive to the reconstructed change in rh_s.

6.2. Perspectives

[68] While we explore the influence of different climate conditions both at the evaporative source and during the poleward transport, our approach is still incomplete. First, we considered only one evaporative source and one trajectory. whereas the polar snowfall originates from different sources through various trajectories [e.g., Helsen et al., 2006]. Given the sensitivity of ¹⁷O excess of mixing of contrasted δ^{18} O moisture, the heterogeneity of the sources could have an impact on the polar ¹⁷O excess. Second, the SCM was run for subtropical conditions, whereas middle and high latitudes are also important sources of vapor for Antarctica snowfall [Delaygue et al., 2000; Werner et al., 2001]. Whereas the sensitivity of the isotopic composition of low-level vapor to surface relative humidity can be applied to all latitudes [Jouzel et al., 1982; Landais et al., 2008], the sensitivity to tropical convective processes is more difficult to generalize to extratropical latitudes. In addition, some effects controlling the ice ¹⁷O excess in middle and high latitudes might have been ignored. For example, 3D large-scale advections or changes in cloud dynamics and microphysics in frontal systems [Gedzelman and Arnold, 1994] might play an important role. The type of water transport (diffusive or advective [Hendricks et al., 2000; Kavanaugh and Cuffey, 2003] might also affect the ¹⁷O excess, since diffusive transport would involve some mixing, which decreases the ¹⁷O excess due to the nonlinearity of the mixing curves in the logarithmic diagram (Figure 2). GCM simulations with isotopic capabilities including $H_2^{17}O$ would be necessary to explore these processes and thus to better understand what information is recorded in snowfall ¹⁷O excess.

Appendix A: Description of the Single-Column Model

Physical Package A1.

[69] The SCM includes bulk formulas for sea surface evaporation, a radiation parametrization [Fouquart and Bonnel, 1980; Morcrette, 1991], the Emanuel convective parametrization [Emanuel, 1991; Emanuel and Zivkovic-Rothman, 1999] and a statistical cloud scheme coupled to this convective scheme [Bony and Emanuel, 2001]. The only dimension is altitude, discretized with 40 pressure levels. A complete description of the isotopic and nonisotopic aspects of this model is given by Bony et al. [2008] and Bony and Emanuel [2001], respectively.

[70] The convective parametrization represents the net effect on the large-scale environment of an ensemble of convective systems. Air parcels from the BL are adiabatically lifted to different levels until they precipitate. The falling precipitation partially or totally reevaporates as it falls through unsaturated atmospheric layers, driving an unsaturated downdraft.

[71] The boundary conditions of this model are surface conditions (surface wind, sea surface temperature and albedo), insolation, CO_2 and large-scale atmospheric forcing. Large-scale vertical motions control the convective activity: for example, a large-scale ascent is associated with large-scale moisture convergence and thus strong convection. In the model, the large-scale circulation is represented by a large-scale ascent or descent, prescribed as a vertical profile of vertical velocity of cubic shape.

[72] Horizontal advections of temperature and humidity are computed from the large-scale velocity. Horizontal gradients of temperature and humidity are neglected, as justified by *Bony et al.* [2008] over tropical oceans.

A2. Representation of Isotopic Processes

[73] The representation of isotopic processes is described in detail by *Bony et al.* [2008]. Water isotopic species ($H_2^{16}O$, $H_2^{17}O$, $H_2^{18}O$ and HDO) are passively transported by the large-scale and convective mass fluxes. As for humidity, horizontal isotopic gradients are neglected. Isotopic fractionation is introduced at each phase change, with fractionation coefficients for $H_2^{18}O$ and HDO given by *Bony et al.* [2008]. For $H_2^{17}O$, we took the experimental values determined by *Barkan and Luz* [2005, 2007] for liquid-vapor equilibrium fractionation coefficients, in the absence of experimental determination, we took the theoretical determination by *Van Hook* [1968].

[74] Isotopic fractionation during surface evaporation is represented by the *Craig and Gordon* [1965] equation (equation (1) of the auxiliary material Text S1). The kinetic fractionation coefficient α_K depends on surface wind speed V_s according to *Merlivat and Jouzel* [1979]. For V_s below 7 m/s, which represents 95% of the ocean surface [*Eriksson and Bolin*, 1964], the kinetic fractionation coefficient is constant.

[75] We assume isotopic equilibrium with the vapor for liquid condensation (above 0°C), and a Rayleigh distillation for ice condensation (below -15° C). Between 0°C and -15° C, the composition of the condensate is assumed to be a linear combination of the compositions of the liquid and solid phases.

[76] Following *Jouzel and Merlivat* [1984], we take into account kinetic effects due to supersaturation with respect to ice S_i , assuming that S_i varies linearly as a function of temperature T,

$$S_i = \mu - \lambda \cdot T \tag{A1}$$

where μ and λ are tunable parameters [Jouzel and Merlivat, 1984]; μ is set to 1 [e.g., Hoffmann et al., 1998; Noone and Simmonds, 2002]. λ has typical values of 0.002 [e.g., Landais et al., 2008], 0.003 [e.g., Hoffmann et al., 1998; Noone and Simmonds, 2002] or 0.004 [e.g., Schmidt et al., 2007; Vimeux et al., 2001; Stenni et al., 2001]. Here we take $\lambda = 0.002$ to be consistent with Landais et al. [2008], who used both d excess and ¹⁷O excess data to constrain λ .

[77] In the unsaturated downdraft, sublimation of ice is assumed not to fractionate due to low isotopic diffusivities in ice.

[78] Fractionation during rain reevaporation and diffusive exchanges between rain and vapor are represented following

Stewart's model [*Stewart*, 1975], as described by *Bony et al.* [2008]. In this model, the evolution of the isotopic composition of the rain drops is calculated using mass conservation equations and assuming that at each instant, the isotopic composition of the rain evaporation R_e is given by *Craig and Gordon* [1965]:

$$R_e = \left(\frac{D'}{D}\right)^n \cdot \frac{\frac{R_l}{\alpha_{eq}} - h_{eff} \cdot R_b}{1 - h_{eff}}$$
(A2)

where R_l is the isotopic composition of the rain, R_b the isotopic composition of the vapor in the unsaturated downdraft (i.e., surrounding the rain shaft) and h_{eff} is the relative humidity at the droplets surface. *D* and *D'* are the diffusivities of water and isotopic species, respectively, taken from *Merlivat* [1978], and *n* is an exponent set to 0.58 [*Stewart*, 1975]. The h_{eff} is parameterized as a function of the relative humidity in the unsaturated downdraft h_{dd} as

$$h_{\rm eff} = \phi + (1 - \phi) \cdot h_{\rm dd},\tag{A3}$$

with ϕ a tunable parameter between 0 and 1. This parameter represents the humidification around the rain drops and controls how much the rain drops reequilibrate with the unsaturated downdraft vapor by diffusive exchange: if $h_{\text{eff}} = 0$, no diffusion occurs. If $h_{\text{eff}} = 1$, only diffusive exchanges occur and the vapor and the droplets tend toward isotopic equilibrium. The ϕ is set to 0.9 to optimize the simulation of the isotopic composition of the tropical rain [*Bony et al.*, 2008]. The isotopic composition of the rain R_l and the downdraft vapor R_b are then calculated using mass balance equations for both water and isotopic species.

Appendix B: Distillation Model MCIM

[79] To model the evolution of the isotopic composition of air masses from the source region to the poles as well as the composition of the Vostok snowfall, we use the Mixed Cloud Isotopic Model (MCIM) [*Ciais and Jouzel*, 1994] adapted for the calculation of ice ¹⁷O excess [*Landais et al.*, 2008], with the same fractionation coefficients as in the SCM.

[80] The MCIM is an extension of Lagrangian models based on a Rayleigh distillation [Merlivat and Jouzel, 1979; Jouzel and Merlivat, 1984]. It describes the isotopic processes at each phase transition and thus the isotopic composition of (1) the condensed phase (liquid water or solid ice) and (2) the water vapor at each step from the oceanic source region to the precipitation site on the ice sheet. During liquid precipitation, only equilibrium fractionation occurs. Then, kinetic fractionation is taken into account for snowflakes formation. Depending on the temperature, the MCIM allows for a zone of mixed clouds, where liquid droplets and ice crystal can coexist. In this zone, the Bergeron-Findeisen process associated with significant kinetic fractionation effects is considered (details given by Ciais and Jouzel [1994]). As for the SCM, the relative proportion of equilibrium versus kinetic fractionation is controlled by the supersaturation.

[81] The transport path of the water mass is described in terms of temperature and pressure. The air parcel is transported in saturated (or supersaturated in polar regions) conditions from the source region to the precipitation site, hence implying continuous fractionation. In nature, air parcels are transported

mainly in unsaturated conditions and most of the saturation occurs during the last day of transport [Helsen et al., 2006]. However, since the depletion associated with the distillation depends mainly on the initial and final temperatures, considering transport in saturated conditions has little impact on the results (less than 2‰ for δ^{18} O, and negligible for d excess [Helsen et al., 2006]).

[82] The model receives as main inputs (1) the temperature and pressure of the source region as well as the isotopic composition of the initial water vapor and (2) the temperature and pressure at the precipitation site.

[83] The MCIM includes several tunable parameters [*Ciais* and Jouzel, 1994] such as the dependence of supersaturation on temperature, the fraction of condensate remaining in clouds. We performed numerous sensitivity experiments to tune these parameters and kept only those enabling a reproduction of the δ^{18} O, d excess and 17 O excess on the Antarctic transect [Landais et al., 2008]. Note that the same dependency of supersaturation with temperature has been taken for the MCIM and the SCM. For the other tunable parameters, we choose values very similar to those used in previous studies [Ciais and Jouzel, 1994; Vimeux et al., 2001; Stenni et al., 2001]. Other parameters have however been tested (section 5).

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