Modelling water isotopologues (\(^1\)H\(^2\)H\(^16\)O, \(^1\)H\(^2\)\(^17\)O) in the coupled numerical climate model iLOVECLIM (version 1.1.5)

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Abstract.
Stable water isotopes are used to infer changes in the hydrological cycle for different climate periods and various climatic archives. Following previous developments of \(\Delta^{18}\)O in the coupled climate model of intermediate complexity iLOVECLIM, we present here the implementation of the \(\Delta^{2}\)H and \(\Delta^{17}\)O water isotopes in the different components of this model, and calculate the d-excess. We also present results of modelled \(^17\)O-excess in the atmosphere and ocean, that was currently only available in the LMDZ4 model. Results of a 5,000 years equilibrium simulation under preindustrial conditions are analysed and compared to observations and several isotopes-enabled models for the atmosphere and the ocean components.

In the atmospheric component, the model correctly reproduces the first order global distribution of the \(\Delta^{2}\)H and d-excess as observed in the data (\(R=0.56\) for \(\Delta^{2}\)H and 0.36 for d-excess), even if local differences are observed. The model-data correlation is within the range of other water isotopes-enabled General Circulation Models. The main isotopic effects are properly modelled, with respect to changes in precipitation or temperature, similarly to LMDZ4 model. The latitudinal gradient is also correctly reproduced in our model and is close to previous water isotopes-enabled General Circulation Models simulations despite a simplified atmospheric component in iLOVECLIM. One exception is observed in Antarctica where the model does not correctly estimate the water isotope composition, consequence of the non-conservative behaviour of the advection scheme at very low moisture content. The modelled \(^17\)O-excess presents a too important dispersion of the values in comparison to the observations and is not correctly reproduced in the model mainly because of the complex processes involved in the \(^17\)O-excess.

For the ocean, the model simulates adequate isotopic composition in comparison to the observations, except for local areas such as in the surface Arabian Sea, a part of the Arctic and West equatorial Indian ocean. Data-model evaluation also presents a good match for the \(\Delta^{2}\)H over the entire water column in the Atlantic Ocean, reflecting the influence of the different water masses.
1 Introduction

Stable water isotopologues (D/H, δ18O, δ17O, δ2H) expressed hereafter in the usual delta notation with respect to V-SMOW scale (Dansgaard, 1964) are important tracers of the hydrological cycle and are measured in a large variety of archives to reconstruct climate variations. At first order, the δ2H and δ18O isotopic ratios of precipitation measured in ice core can be used as a proxy of past temperature at the drilling site (e.g. Johnsen et al., 1972; Lorius et al., 1979; Jouzel, 2003). As they present the same variations, and following observations at lower latitudes, we can derive a second-order parameter called deuterium excess (d-excess) from the difference between the δ2H and δ18O. During evaporation, kinetic non-equilibrium processes affect the relationship between oxygen and hydrogen isotopes and lead to a deviation from the global Meteoric Water Line (MWL), which represents the linear relationship between δ2H and δ18O (Craig, 1961; Dansgaard, 1964):

\[
d\text{-excess} = \delta^2H - 8 \times \delta^{18}O
\]

This parameter is a classical polar ice-core tracer that can be used to provide additional constraints on past climates and changes in the atmospheric water cycle. The deuterium excess is conventionally interpreted in terms of temperature at the moisture source, or shifts in moisture origin (Stenni et al., 2001; Vimeux et al., 2002; Masson-Delmotte et al., 2005) even if it can also be impacted by local temperature (Masson-Delmotte et al., 2008a) and by mixing along trajectory (Hendricks et al., 2000; Sodemann et al., 2008). Modelling studies such as Risi et al. (2013) also suggested that the d-excess is controlled by convective processes and rain re-evaporation at the tropics and by effect of distillation and mixing between vapors from different origins at high latitudes. Recently, Landais et al. (2021) also shown using the first 800 000 years d-excess record that precipitation, seasonality and moisture source regions changes in the past can complicate the interpretation of the d-excess.

Following experiment developments for an accurate measurement of δ18O abundance (Barkan and Luz, 2007; Landais et al., 2008), a second-order parameter, the δ17O-excess, has been defined as:

\[
\delta^{17}\text{O}\text{-excess} = \ln \left( \frac{\delta^{17}\text{O}}{1000} + 1 \right) - 0.528 \times \ln \left( \frac{\delta^{18}\text{O}}{1000} + 1 \right)
\]

The δ17O-excess is then multiplied by 10 to express it in per mil since magnitudes are very small (Landais et al., 2008). Note that we used the logarithm notation for δ17O-excess following Luz and Barkan (2005). This definition makes it very sensitive to mixing between vapors of different origins (Risi et al., 2010).

The δ17O-excess is commonly used in ice core based paleoclimate studies to give information on the relative humidity over the ocean (e.g. Landais et al., 2008, 2018; Risi et al., 2010; Steig et al., 2012). This proxy is controlled by kinetic fractionation during evaporation, and similarly to d-excess, is very sensitive to empirical parameters determining the supersaturation in polar
clouds (Landais et al., 2012; Winkler et al., 2012). Since influences of temperature or condensation altitude on $^{17}$O-excess are expected to be insignificant in contrast to d-excess, measurements of $^{17}$O-excess have an added value with respect to d-excess and can be used to disentangle the parameters (temperature, relative humidity) that affects the water isotopic composition. For example, Risi et al. (2010) shown that the different behaviors of d-excess and $^{17}$O-excess in polar regions could be related to fractionation processes along the distillation pathway form the evaporative source to polar region that affect more the d-excess than the $^{17}$O-excess, with $^{17}$O-excess recording more the signal from low latitudes during surface evaporation. Modelling the $^{17}$O-excess is still very challenging since it depends on complex processes that have to be properly reproduced in the climate models. To date, only the LMDZ4 model has included the $^{17}$O-excess (Risi et al., 2013). However, even if the processes that control the $^{17}$O-excess are more complex than those controlling the d-excess, the combination of the d-excess, $^{17}$O-excess and $^{18}$O could bring new information on the understanding of past changes in local temperature, moisture origin and conditions at the moisture source.

Among the new proxies to document the water isotopic ratio in precipitation, the hydrogen isotope composition of plant lipids wax (alkanes) has been found to reflect predominantly local continental rainfall fluctuations (e.g. Schefuß et al., 2005; Collins et al., 2013; Kuechler et al., 2013). Isotopic changes are primarily controlled by moisture loss by evapotranspiration, soil waters conditions and precipitations rates, but the vegetation and isotopic enrichment effects are also to consider (Hou et al., 2008; Sachse et al., 2012; Kahmen et al., 2013a,b). Another method has also been developed to extract the fossil water (fluid inclusions) of speleothem records (Vonhof et al., 2006; van Breukelen et al., 2008). It then becomes possible to realize hydrogen and oxygen stable isotope analyses of fossil precipitation waters and document the deuterium excess values in the past, outside the limited region of ice core presence.

Similarly to continental records, the isotopologues in ocean surface waters track regional freshwater balance and then the hydrological cycle (Craig and Gordon, 1965). Water isotopologues in seawater can therefore be used as a proxy for salinity since surface freshwater exchanges are important in determining the variability of both variables. Seawater oxygen isotope concentration preserved in carbonate from organisms such as foraminifera allows estimations of past regional and qualitative changes in salinity and ocean circulation (Schmidt et al., 2007; Caley et al., 2011). It has been suggested that combining seawater hydrogen isotopes ($\delta^2$H obtained from alkenones or other biomarkers) with oxygen isotope ($\delta^{18}$O obtained from zooplankton calcite shells of foraminifera) could be a promising way to quantitatively estimate salinity variability (Rohling, 2007; LeGrande and Schmidt, 2011, Leduc et al., 2013; Caley and Roche, 2015).

With the emergence of new paleoproxy to document water isotopologues in atmospheric and oceanic components of the climate system, the necessity to develop and use isotope-enabled models, and in particular coupled ocean-atmosphere models, as never been greater (e.g. Risi et al., 2012; Werner et al., 2016; Cauquoin et al., 2019). These later allow more complex assumptions related to paleoclimatic proxies to be examined (LeGrande and Schmidt, 2006; Schmidt et al., 2007). For example, the simulation of the climate and its associated isotopic signal can provide a “transfer function” between the isotopic signal...
and the considered climate variable such as precipitation rate/water isotopes in precipitation or salinity/water isotopes in seawater relationships.

Since the initial works of Joussaume et al. (1984) and Jouzel et al. (1987), much progress has been done in atmospheric general circulation models (AGCMs) (e.g. Hoffmann et al., 1998; Noone and Simmonds, 2002; Mathieu et al., 2002; Risi et al., 2010; Werner et al., 2011) that can simulate accurately the $\delta^{18}O$ of precipitation. The subsequent development of water isotopes modules in oceanic general circulation models (OGCMs) (Schmidt, 1998; Delaygue et al., 2000; Xu et al., 2012) opens the possibility for coupled simulations of present and past climates, conserving water isotopes through the hydrosphere (Schmidt et al., 2007; Zhou et al., 2008; Tindall et al., 2009; Werner et al., 2016; Cauquoin et al., 2019). In general, General Circulation Models (GCMs) have been used exclusively to simulate separately water isotopes in the atmospheric and oceanic components. Given the computing resources needed to run coupled climate models, applying intermediate complexity coupled climate models with water isotopes like iLOVECLIM to long-term palaeoclimate perspectives still appears quite suitable (e.g. Caley et al., 2014). It could allow to improve our understanding of the relationship between water isotopologues, second-order parameter (like d-excess) and climate over a broad range of simulated climate changes.

Oxygen isotopes (18, 16) have been implemented in iLOVECLIM, allowing fully coupled atmosphere-ocean simulations. The detailed implementation of oxygen isotopes in iLOVECLIM and the evaluation against observed data in water samples and carbonates can be found in Roche (2013), Roche and Caley (2013) and Caley and Roche (2013). In the present manuscript, we present the design and the validation of $\delta^2H$ water isotopes as well as deuterium excess and $\delta^{17}O$-excess in the coupled climate model iLOVECLIM for the atmospheric and oceanic components. The agreements and differences from the direct comparison between modelling results under pre-industrial conditions with (1) multiple datasets and (2) several isotope-enabled GCMs results for the atmosphere and the ocean components will be discussed to determine the potential and the interest of using iLOVECLIM for palaeoclimatic studies.

2 Description of the water isotopic scheme in iLOVECLIM

2.1 Atmospheric component ECBilt

The iLOVECLIM model (version 1.1.5) is a derivative of the LOVECLIM-1.2 climate model extensively described in Goosse et al. (2010). It is composed of an atmospheric, oceanic, land surface and vegetation component. The atmospheric component ECBilt is a quasi-geostrophic model with a T21 spectral grid (resolution of 5.6° in latitude and longitude) with a complete description of the water cycle from evaporation, condensation to precipitation. The timestep of the atmospheric component is 6 hours. It is subdivided in three vertical layers at 800, 500 and 200 hPa with the humidity contained only in the first layer and representative of the total humidity content of the atmosphere. Evaporative water fluxes are added to this humid layer and vertical advection is computed. Water fluxes crossing the limit between the humid and dry layers are rained out instantly as...
convective rain. For specific humidity of the humid layer larger than 80 % (set as the saturation humidity at given temperature),
the excess water is removed as large-scale precipitation. If large scale precipitation occurs with negative temperatures, excess
precipitation is removed as large-scale snowfall.

With regards to water isotopes, the main development lies in the atmospheric component in which evaporation, condensation
and the existence of different phases (liquid and solid) all affect the isotopic conditions of the water isotopes. The methodology
used to trace the hydrogen water isotopes in ECBilt is identical to the description made in Roche (2013) for the oxygen water
isotopes. We used the same equations presented for the $^{18}$O in Roche (2013) but with adapted fractionation coefficients for the
hydrogen and for $^{17}$O. We present in this section the equations for the heavy/light isotope ratios. Additional information on
general water scheme formulation can be found in Roche (2013).

In ECBilt, the water isotopic quantity is expressed as a single tracer of water similarly to Merlivat and Jouzel (1979). For
$^{1}H^{2}^{16}O / ^{1}H^{2}^{18}O$, it is defined as a function of the quantity of precipitable water for the whole atmospheric column ($q$ which
depends on the mass of the water, the surface area of the cell and the water density) and of the ratio ($R^{H}$) between the number
of moles of $^{1}H^{2}^{16}O$ and the number of moles of $^{1}H^{2}^{18}O$:

$$ q^{H} = q \times R^{H} $$

(3)

The isotopic composition then changes within the water cycle, from evaporation to precipitation. The evaporation term for
hydrogen water isotopes cannot be simply written like for the humidity because there is no vertical discretization for water
isotopes in the model. The solution adopted by Roche (2013) is to compute the water isotopic ratio in the evaporation using a
Craig and Gordon (1965) type-model in the formulation adapted by Cappa et al. (2003). The hydrogen isotopic ratio of
evaporating moisture can then be written as:

$$ R^{H}_E = \alpha^{*, diff} \left( \frac{R^{H}_{eq} - h^{*}_{a} R^{H}_{a}}{1 - h^{*}_{a}} \right) $$

(4)

where $R^{H}_{eq}$ is the isotopic ratio at equilibrium with the ocean, $R^{H}_{a}$, the isotopic ratio of the humidity in the atmosphere and $h^{*}_{a}$
is an apparent relative humidity value for the atmosphere. $\alpha^{*, diff}$ is a ratio of molecular diffusivity and defined for the hydrogen
such as:

$$ \alpha^{*, diff} = \left( \frac{D^{H}}{D} \right)^{n} $$

(5)
with $D$ the molecular diffusivity of water $\text{H}_2\text{H}^{16}\text{O}$, $D$ the molecular diffusivity of water $\text{H}_2\text{O}$ and $n$ a coefficient that varies with turbulence and evaporative surface (Brutsaert, 1975; Mathieu and Bariac, 1996). The molecular diffusivity ratio for $\text{H}_2\text{H}^{16}\text{O} / \text{H}_2\text{O}$ is set to 0.9755 (Merlivat, 1978) and 0.9855 for $\text{H}_2\text{O} / \text{H}_2^{16}\text{O}$ (Barkan and Luz, 2007).

Since ECBilt only includes three layers, it is supposed that precipitation always forms in isotopic equilibrium with the surrounding moisture with instantaneous rainout to the surface. The precipitations (convective and large scale) and snow are in equilibrium with isotopic values at 650, 800 hPa and 650 hPa respectively. When computing the precipitation and snow fractionation schemes (see Roche, 2013), we take into account the temperature, the fractionation coefficients between the different water phases for the hydrogen, an enhanced kinetic fractionation at high latitude (Merlivat and Jouzel, 1979) and the ratio of hydrogen isotopes in vapor. In these equations, the hydrogen fractionation coefficient between liquid water and vapor is taken from Majoube (1971a) and depends on the temperature:

$$
\alpha_{\text{H}_{2}\text{O} / \text{H}_{2}\text{O}} = \exp \left( \frac{24844}{T^2} - \frac{76.248}{T} + 0.052612 \right) 
$$

(6)

For $\text{O}$, the fractionation between liquid water and vapor is calculated from Majoube (1971a), Barkan and Luz (2005; 2007):

$$
\alpha_{\text{H}_{2}^{18}\text{O} / \text{H}_{2}\text{O}} = \exp \left( \frac{1137}{T^2} - \frac{0.4156}{T} - 0.0020667 \right) \times 0.529
$$

(7)

The equilibrium fractionation coefficient between solid water and water vapor for hydrogen is taken from Merlivat and Nief (1967) and depend on the temperature as well:

$$
\alpha_{\text{H}_{2}\text{O} / \text{H}_{2}\text{H}^{16}\text{O}} = \exp \left( \frac{16289}{T^2} - 0.0945 \right)
$$

(8)

For $\text{O}$, the fractionation between solid water and vapor is calculated from Majoube (1971b), Barkan and Luz (2005) and Barkan and Luz (2007):

$$
\alpha_{\text{H}_{2}^{18}\text{O} / \text{H}_{2}\text{H}^{16}\text{O}} = \exp \left( \frac{11.839}{T^2} - 0.028224 \right) \times 0.528
$$

(9)
2.2 Ocean and land surface components

The oceanic component CLIO has a 3x3° horizontal resolution, 20 vertical layers and a free surface. All the variables are calculated with a daily timestep. In the ocean, the water isotopes are mass conserving and act as passive tracers under equilibrium fractionation ignoring the small fractionation implied by the presence of sea-ice (Craig and Gordon, 1965).

For the land surface model, the isotopes water implementation in the bucket follows the same procedure as for the water. If re-evaporation occurs on land, it is assumed to be at equilibrium (without fractionation). A snow layer is also taken into account. Above a given threshold, the isotopic water and snow contents in the soil and snow buckets are routed to the ocean without fractionation.

2.3 Simulation setup

We present results of a 5,000 years equilibrium run under fixed pre-industrial boundary conditions. The atmospheric pCO$_2$ is chosen to be 280 ppm, methane concentration is 760 ppb and nitrous oxide concentration is 270 ppb. The orbital configuration is calculated from Berger (1978) with constant year 1950. We use present-day land sea mask, freshwater routing and interactive vegetation. With regards to the water isotopes, the atmospheric moisture is initialized at 0 and the δ$^2$H at 0‰. The consistency of our integration is checked by ensuring that the water isotopes are fully conserved in our coupled system. The model has been run at T21 spatial resolution and the output are computed with an annual timestep.

To investigate the seasonal variations of the model in comparison to the observations, and to estimate the range/dispersion of the modelled results, we performed a 100 years simulation starting from the equilibrium run, with monthly outputs for the climate and the isotopes. This simulation is investigated in Section 3.1.4.

2.4 Observational data and water isotopes-enabled GCMs

To allow for comparison and discussion with iLOVECLIM results, global hydrogen and d-excess isotopic datasets for the atmosphere from the Global Network of Isotopes in Precipitation (GNIP) dataset (IAEA, 2023) and Masson-Delmotte et al. (2008b) have been used. The original GNIP dataset has been subsampled to keep only the stations where the isotopic composition has been reported for a minimum of 3 calendar years within the period 1961-2008. To evaluate the seasonal evolution of the model, we looked at the evolution of precipitations and atmospheric isotopic composition at several locations distributed on multiple continents to reflect the variety of climate: Pretoria (25.73°S, 28.18°E), Belem (1.43°S, 48.48°W), Ankara (39.95°N, 32.88°E) and Reykjavik (64.13°N, 21.92°W). Present day measurements of δ$^{17}$O-excess from multiple studies (Landais et al., 2008, 2010, 2012; Luz and Barkan, 2010; Uemura et al., 2010; Winkler et al., 2012; Pang et al., 2015; Tian et al., 2021) have also been used. Note that the data of Uemura et al. (2010) are for the vapor and not the precipitation and does not allow for a direct model-data comparison.
The GISS global seawater isotope database (Schmidt et al., 1999) has been used to compare the $\delta^1$H and d-excess with the ocean component in the model. We looked at the surface distribution of the isotopes for the first oceanic layer at 5 m depth in the model and selected GISS sea water values between 0 and 10 m to be representative of the surface.

To evaluate our model results against water isotopes-enabled GCMs, we used several model outputs: ECHAM5-wiso (Steiger et al., 2018), GISS (Schmidt et al., 2007), LMDZ4 (Risi et al., 2010, Risi et al., 2013), MIROC (Kurita et al., 2011), CAM (Lee et al., 2007) and MPI-ESM-wiso (Cauquoin et al., 2020). The GISS, LMDZ4, MIROC and CAM data are from the Stable Water Isotope Intercomparison Group, Phase 2 (SWING2) (Risi et al., 2012). $\delta^1$H$_{\text{water}}$ in MPI-ESM-wiso has been calculated from $\delta^{18}O_{\text{ seawater}}$ and d-excess outputs.

3 Results and discussion

3.1 Water isotopic composition in the atmosphere

3.1.1 Annual $\delta^1$H$_{\text{precipitation}}$

The annual mean modelled distribution of $\delta^1$H$_{\text{precipitation}}$ is presented in comparison to observations on Fig. 1a. The latitudinal gradient from the poles to the equator is correctly reproduced in the model with depleted values at high latitudes (cold and dry regions) and enriched values at lower latitudes. Regions like central Africa and northern region of South America show however differences with the data since the modelled $\delta^1$H$_{\text{precipitation}}$ is underestimated in comparison to the few measurements available. This could be due to one of the well-known iLOVECLIM biases that is the overestimation of the precipitation in these regions. The west coast of South America also presents discrepancies between the model and the GNIP data (Fig. 1a). This could be related to the coarse model resolution that may not perfectly reproduce the observed $\delta^1$H$_{\text{precipitation}}$ since the value is representative of a larger area. Finally, the modelled $\delta^1$H$_{\text{precipitation}}$ over northern America and Europe is higher than the observations. The difference in atmospheric isotopic composition of precipitation over land and ocean is however well reproduced in the model with values closed to zero over the Pacific, Atlantic and Indian oceans and depleted values lower than -50 ‰ and -80 ‰ respectively over the Arctic and Austral oceans (Fig. 1a).

We also compared the zonal distribution of several water isotopes-enabled GCMs for results that co-locate with observations. From mid to low latitudes, all models show similar $\delta^1$H$_{\text{precipitation}}$ with iLOVECLIM being more enriched than the other GCMs below 20°S and above 30°N. The model however reproduces the global trend of depleted values at high latitudes and enriched values at low latitudes, as observed in the data (Fig. 2a). At high latitudes, iLOVECLIM models an isotopic composition that is too enriched compared to the one in ECHAM5-wiso, GISS, LMDZ4, MIROC and CAM models, as well as in the GNIP data with values between up to -453 ‰ (Fig. 2a). These very low measured values over Antarctica can be explained by the low temperature (with a continental effect) and by other influences like moisture transport or the distance from the coast that add complexity in modelling this region (Fig. 2a). Since iLOVECLIM only have three vertical layers in comparison to the 19 to 26 vertical layers for the other GCMs, we cannot properly reproduce the isotopic variations at these latitudes as a...
consequence of the non-conservative behaviour of the advection scheme at very low moisture content. However, no model is able to correctly reproduce these very low values as observed in the measurements. All the GCMs model higher values, between -305 ‰ and -365 ‰.

In order to further evaluate our model results against water isotopes-enabled models and the observations, we analysed the standard deviation (SD), correlation R and root mean square error (RMSE), combined in a Taylor Diagram (Fig. 3). In all these figures, we removed Antarctic values for the reason explained above. We observe for the $\delta^2$H$_{\text{precipitation}}$ that ECHAM5-wiso is the model that has the best correlation coefficient with the observation ($R=0.64$ vs $R=0.56$ for iLOVECLIM). The different GCMs have close correlation coefficient (between 0.59 and 0.64), standard deviation (between 40.21 and 46.43) and RMSE (between 34.94 and 39.82). The iLOVECLIM model presents a lower standard deviation (SD=29.93) and RMSE than the other models (Fig. 3a). However, considering the close metrics between all models, iLOVECLIM presents the advantage to run faster than other GCMs and is perfectly justified for the use of long-term global climate simulation.

Figure 1: Model-data evaluation of the annual mean isotope distributions. (a) $\delta^2$H in precipitation, (b) d-excess and (c) $^{17}$O-excess in iLOVECLIM. The model results are compared to observations (in circles).
The mean annual d-excess distribution is derived from the oxygen and hydrogen isotopic composition. To evaluate the accuracy of the model, we compare the model results to the observations. As observed for the \( \delta^2H_{\text{precipitation}} \), the d-excess results to the observations. The model values match the GNIP observations and correctly locate with the observations. The gray points.

### 3.1.2 Annual deuterium excess

The mean annual d-excess distribution is derived from the oxygen and hydrogen isotopic composition. To evaluate the accuracy of the model, we compare the model results to the observations. As observed for the \( \delta^2H_{\text{precipitation}} \), the d-excess

### Figure 2: Multi-model zonal (a) \( \delta^2H_{\text{precipitation}} \) and (b) d-excess comparison. The model results (in color) are compared against the observations (in grey). The dotted curved line indicates the reference line.

### Figure 3: Taylor diagram representing (a) \( \delta^2H_{\text{precipitation}} \) and (b) d-excess and (c) \(^{18}O\)-excess values for different climate models (iLOVECLIM, LMDZ4, ECHAMS-wiso, CAM, GISS and MIROC) without Antarctic values. The simulated values are plotted against the observations. The dotted curved line indicates the reference line (standard deviation of the observation) and the bold grey contours represent RMSE values.
presents a latitudinal gradient with depleted negative values to the poles and enriched positive values to the equator (Fig. 1b and Fig. 2b). The modelled values fit well with the observations at global scale. Differences between the model and the observations remain for some regions like over India where the modelled d-excess is slightly higher than the observations. This is also observed in the zonal distribution were the modelled d-excess in iLOVECLIM is higher than observation and other GCMs from mid to low latitudes (Fig. 2b). The modelled d-excess over Greenland, and especially the coastal areas, is negative whereas the few available data points indicate positive values that are up to 20 ‰ higher. Similarly to the annual $\delta^2$H$_{\text{precipitation}}$ distribution, the d-excess over Antarctica is not correctly reproduced in the model and presents outliers values in the coastal regions. The local data show values between 5 and 10 ‰ whereas the model calculates values ranging from -10 to 25 ‰ or higher in the region of Adélie Land (Fig. 1b). For a better zonal multi-model comparison, we decided to exclude these outlier values from the Figure 2b. In comparison to the observations, the LMDZ4, GISS and CAM models have enriched values from mid to high latitudes, whereas ECHAM5-wiso has systematically depleted values. The MIROC model is the only one that shows a different trend in the zonal distribution of the d-excess, with higher values in the high latitudes and depleted values to the equator. Over the ocean, few d-excess data points are available but the model presents an overall good agreement with the GNIP data with mean values ranging from -10 ‰ over the Arctic and Austal oceans to 17 ‰ over the Atlantic and Pacific oceans. A maximum in d-excess is reach over the Arabian sea with 20.6 ‰.

In comparison to the measurements for the atmosphere, iLOVECLIM has a correlation coefficient that is in the range of others models (0.34 to 0.52), but has a higher SD compared to the observations and other GCMs. The CAM model has the best correlation coefficient with the observations whereas LMDZ4 has the closest standard deviation relative to the observations (Fig. 3b). Within all models, MIROC is the one with the lowest SD and RMSE. However, considering the general low correlation coefficient for all models, they all do not perfectly reproduce the d-excess variations as observed in the data. iLOVECLIM however presents the advantage to run faster than the other GCMs and could be used to investigate past changes in d-excess in global transient simulations.

The relationship between the d-excess and the $\delta^2$H$_{\text{precipitation}}$ can be investigated and shows that it is partially driven by high latitudes values, mainly in Antarctica, as presented in Fig. 4. From the globally available data, a relationship between d-excess and $\delta^2$H$_{\text{precipitation}}$ exists with high d-excess value (~15 ‰) for highly depleted $\delta^2$H$_{\text{precipitation}}$ values (around -400 and 0 ‰), whereas lower d-excess is observed for mean $\delta^2$H$_{\text{precipitation}}$ between -250 and -300 ‰. The low $\delta^2$H$_{\text{precipitation}}$ values correspond to high latitudes values, mostly corresponding to Antarctic values, that drive the relationship between d-excess and $\delta^2$H$_{\text{precipitation}}$ ($R^2=0.50$ when considering all values, $R^2=0.10$ for values without the high latitudes). Similar relationship between the d-excess and $\delta^2$H$_{\text{precipitation}}$ is observed in the iLOVECLIM model. Highest d-excess values are obtained for low $\delta^2$H$_{\text{precipitation}}$ values (around -200 ‰) and lower d-excess for intermediate $\delta^2$H$_{\text{precipitation}}$ (Fig. 4). The shape of the regression curves is however different between the data and the model because of outlier modelled d-excess values that are too high in the model. These data points mainly correspond to Antarctic values as already observed on Fig. 1.
Antarctic isotopic values are not computed correctly due to issues in the conservation of water in the advection scheme at very low humidity content, a fact that was already highlighted in Roche (2013). Improving the conservation in the spectral advection scheme is beyond the scope of the present study. We thus removed these Antarctic values in the following to investigate the isotopic trend without the influence of this region. This results in a better agreement between the data and iLOVECLIM model (with a correlation coefficient of 0.71), even if differences are observed with generally lower d-excess value in the model than in the data for low $\delta^2$H precipitation (Fig. 4).

For the d-excess, the range of modelled values can be large for some locations (as already seen in Fig. 1). Thus, we can evaluate the ability of the model to reproduce the d-excess in comparison to the observed data, as presented in Fig. 5. The distribution of most d-excess values is centred around values between 8-18‰. Low correlation coefficient is obtained due to outlier d-excess values but statistical significance between the model and the data is obtained with a p-value of 3e-4 (<0.001). This attests of a good representation of the d-excess in the model (excluding Antarctic values). This is also supported by the modelled d-excess in LMDZ4 that presents similar values than in iLOVECLIM (Fig. 5). However, considering the larger dispersion of the values in our model compared to LMDZ4 and to the fact that the uncertainties on the d-excess measurements are large, the relationship between model and data might vary and get closer to the expected 1:1 line.

Figure 2: Global relationship between d-excess and $\delta^2$H in precipitation. High latitude values (above 60°N and below 60°S) are presented with the red triangles for the data and with the light blue triangles for iLOVECLIM. Data for other regions are presented with the orange circles for the measurements and with the dark blue circles for the model. Regression curves for the data and the model, without high latitudes values, are also shown in orange and dark blue.
3.1.3 $^{17}$O-excess distribution

Modelled $^{17}$O-excess shares common pattern with $^{18}$O (itself presenting the same spatial pattern than $^{18}$O, see Appendix A) with depleted values over the high latitudes of the Northern Hemisphere and higher values over land (Fig. 1c). The $^{17}$O-excess presents values between 0 and 100 permeg over the Atlantic Ocean, that are lower than in the Indian and Pacific oceans. In comparison to the LMDZ4 model that is currently the only GCM to include the $^{17}$O ($\text{Risi et al., 2013}$), iLOVECLIM presents higher values for most of the latitudes, due to these high values over the ocean. The latitudinal gradient is also larger in LMDZ4 that has relatively homogenous values between 70°S and 90°N. The model reproduces $^{17}$O-excess values that are close to observations over North America, Europe and Africa (Fig. 1c). But $^{17}$O-excess over the Arabian Sea and northern Canada has probably too negative values. Similarly to d-excess and due to the outlined problem in modelling this region, the $^{17}$O-excess modelled over Antarctica present a wide range of values from high negative to high positive and does not fit with ice core measurements.

Figure 3: Relationship between the modelled d-excess in iLOVECLIM (blue) and in LMDZ4 (red) versus measurements without Antarctic values. The errors bars associated with the data are shown at 2σ. The 1:1 line is shown with the black dashed line. Regression lines for iLOVECLIM and LMDZ4 are in dark blue and red respectively with the confidence bands.

Comparison can be done between model and observations for the $^{17}$O-excess (Fig. 6), even if few data exist. A wide dispersion of the $^{17}$O-excess values (excluding values in Antarctica) is observed, with no clear relationship, mostly due to some highly negative values (up to -116‰) in the model, but statistically significant with a p-value of 0.0023 (<0.05). $^{17}$O-excess has also been previously modelled in LMDZ4 ($\text{Risi et al., 2013}$) and presents a lower dispersion of the values than iLOVECLIM, even negative values of the C ice core measurements. 17 Canada has probably too negative values. Similarly to d-excess, the values over land are more depleted than values over the ocean. In comparison to Luz and Barkan (2010) data available over North America, Europe, Asia and Indonesia, iLOVECLIM mostly presents enriched values of permil, even if some agreements are observed between the model and the data. These discrepancies can be explained by the fact that the most of the data is punctual and reflect seasonal conditions whereas the model outputs are annual mean $^{17}$O values, and/or due to the uncertainties associated with the measured values.

Figure 6a presents the relationship between modelled and measured $^{17}$O$_{\text{precipitation}}$ (excluding values in Antarctica). Most of the values modelled in iLOVECLIM are grouped around enriched isotopic values, but the correlation remains low. The p-value is calculated to 0.079 (>0.05), which is slightly higher than the 5 % limit, indicating a close statistical significant between the two parameters. In comparison to LMDZ that is currently the only GCM to include the $^{17}$O ($\text{Risi et al., 2013}$), iLOVECLIM results are in good agreement with most of the values between -2 and -5‰, leading to similar linear trend between the model and the data (Fig. 6a). One point with negative value of -8.7‰ in LMDZ gets closer to the 1:1 line than iLOVECLIM (with -5.8‰). However, considering the large
if most of the data fit within 10–40 ‰ (Fig. 6). LMDZ4 modelled $^{17}$O-excess is however different than iLOVECLIM and the expected data value (considering the 1:1 line) for extrema values: higher values than data are modelled for depleted values and lower values than data are modelled for enriched values. We observe for the $^{17}$O-excess a low correlation coefficient for iLOVECLIM and a low negative correlation coefficient for LMDZ4 with respect to observations. The standard deviation and root mean square error is better for LMDZ4 than for iLOVECLIM (Fig. 3c), suggesting that our model does not correctly reproduce the $^{17}$O-excess and has a too important dispersion of the values, even if the trend is correct.

Figure 4: Relationship between the iLOVECLIM modelled isotopic values and $^{17}$O-excess measurements, without values in Antarctica. LMDZ4 model results are also presented for comparison with iLOVECLIM. The regression curves between model and data are presented in dark blue for iLOVECLIM and red for LMDZ4 with the confidence bands. The 1:1 line are shown with the black dashed lines.

3.1.4 Seasonal variations

We compare the seasonal model results for precipitation, $\delta^2$Hprecipitation and $d$-excess to the GNIP monthly data at several locations representative of various climate conditions to have a global overview. South Africa (Pretoria), South America (Belem), eastern Mediterranean (Ankara) and northern Atlantic (Reykjavik). GNIP data don’t include $^{17}$O-excess so we don’t represent their comparison. We extracted the model results at the corresponding locations but due to the coarse resolution of the model, regional biases exist as depicted in previous section. We performed a mean over the last 10 years of the simulation and normalized the results (we subtracted the annual mean and divided by the standard deviation for each station) for easier comparison with the data. The seasonal evolution of precipitation and isotopic composition in the model is then not expected...
to perfectly reflect the measurements. We then present the normalized values for both model and GNIP data. There is a good agreement in precipitation at Pretoria and Ankara between the observation and the model that correctly reproduce the seasonal cycle (Fig. 7). For Belem and Reykjavik stations the model shows some differences, namely higher precipitations in September and October at Belem and higher monthly amplitude at Reykjavik. Good correlation is observed for the modelled $\delta^2H_{\text{precipitation}}$ in comparison to observations at Pretoria and Ankara (even if the October value is largely depleted). As for precipitations, the amplitude of $\delta^2H_{\text{precipitation}}$ variations is different between the model and the data at Belem and Reykjavik (Fig. 7). But the overall model behaviour in reproducing seasonal variations of $\delta^2H_{\text{precipitation}}$ can be validated based on these observations, especially when considering that the uncertainties associated with the data can be as large as the measurement itself. The d-excess variations show however larger differences between the model and the observations. The modelled d-excess at Reykjavik shows a good agreement with the observation, while larger amplitude of the variations is observed at Belem (Fig. 7). At Ankara, the modelled d-excess is delayed during summer compared to observations and shows too low values in October. At Pretoria, even if the $\delta^2H_{\text{precipitation}}$ is correctly reproduced in the model, the d-excess presents differences with enriched values between May and September, whereas the data indicates lower values during this period. All these model-data differences could be the result of uncertainties associated to the GNIP data and/or to biases in modelling the d-excess.

Figure 5: Monthly evolution of precipitation (top row), $\delta^2H_{\text{precipitation}}$ (middle row) and d-excess (bottom row) at several stations (different columns for Pretoria, Belem, Ankara and Reykjavik). The red line is the GNIP data measured at the station and the blue line is the ILOVECLIM model result at the corresponding location. The data and model results have been normalized. The error bars for the data are also shown at 2σ.
3.2 Evaluation of the main isotopic effects

3.2.1 Amount effect

The amount effect can be defined as a decrease of the isotopic composition for an increase in the precipitation amount. We investigate this effect in the model and compare it to LMDZ4 and to observations. We only extracted values in the models and for the GNIP stations that cover the tropics, from 0–20°N and 0–20°S, to see if a change in precipitation intensity would lead to a change in the hydrogen isotopic composition of the precipitation. For an easier comparison, we normalized the values (we subtracted the annual mean and divided by the standard deviation).

The seasonal cycle in iLOVECLIM is well reproduced and in agreement with the GNIP data (especially for the precipitations between 0–20°S). In the north tropics (Fig. 8a), the isotopic composition of the precipitation of iLOVECLIM is more depleted during the wet season (i.e. during the boreal summer). The opposite effect is observed in the south tropics (Fig. 8b), with enriched δ²H_{precipitation} during the austral winter, associated with a reduced amount of precipitation. These seasonal variations are explained by the fact that precipitation is mostly affected by secondary evaporation during summer, while condensation is mostly responsible for isotope depletion during winter. So, the δ²H_{precipitation} decreases as precipitation intensity increases. In the model, the minimum depleted δ²H_{precipitation} (maximum enriched δ²H_{precipitation}) is also leading the minimum observed for the GNIP stations of one month (maximum observed for the GNIP stations of two months). This delay between the data and the model is also observed for LMDZ4 for the north tropics, with a lag of one month.

Figure 6: Seasonal variations of the mean precipitation and δ²H_{precipitation} in the tropics, from 0–20°N for (a) and from 0–20°S for (b). The values have been normalized, the solid lines represent the precipitation and the dashed lines the δ²H_{precipitation}. The blue curve presents the iLOVECLIM values, the red curve is for LMDZ4 and the green curve corresponds to the GNIP data.
We further investigate this amount effect by looking at the change in the $\delta^2$H$_{\text{precipitation}}$ as a function of the amount of precipitation. Following Risi et al. (2008; 2010), we looked at the seasonal model variations for nine oceanic tropical GNIP stations (Apia, Barbados, Canton Island, Diego Garcia, Madang, Taguac, Truk, Wake Island and Yap). Since the resolution in iLOVECLIM is T21, the local processes may not be perfectly reproduced and complicate the comparison to local oceanic observation. We then decided to take the best neighbour pixel for each station, by selecting the pixel that was in better agreement with the GNIP data from both precipitation and isotopic composition seasonal cycle. We also do not present observational precipitation values above 350 cm.y$^{-1}$ since in the model precipitations are never higher. Figure 9 presents the relationship between the $\delta^2$H$_{\text{precipitation}}$ and the precipitation for the selected stations in iLOVECLIM and the observation (panel a) and in LMDZ4 (panel b). The isotopic composition of precipitation is enriched for low precipitations and changes toward depleted values as precipitations increase. This amount effect is -0.085‰/cm.y$^{-1}$ for iLOVECLIM, in comparison to -0.139‰/cm.y$^{-1}$ for the GNIP data. The modelled $\delta^2$H$_{\text{precipitation}}$ is however higher than the observations for the same precipitation amount (especially at high precipitations). In contrast, the standard version of LMDZ4 has slightly too depleted $\delta^2$H$_{\text{precipitation}}$ at low precipitations in comparison to the observations as already observed in Risi et al. (2010).

![Monthly $\delta^2$H$_{\text{precipitation}}$ as a function of the precipitation at the location of nine tropical oceanic GNIP stations. (a) iLOVECLIM results in blue compared to GNIP data in green and (b) LMDZ4 results. The error bars for the data are shown at 2σ.](image)

### 3.2.2 Temperature effect

Temperature is supposed to play a role on the hydrogen isotopic composition of precipitation with more depleted values for low temperatures. We investigate in this section this relationship in iLOVECLIM and compare it to the LMDZ4 model. Since in our model the surface temperature is not a prognostic variable, we used the temperature at 650 hPa and took the equivalent temperature in LMDZ4 model at 662 hPa. An enhanced depletion of the $\delta^2$H$_{\text{precipitation}}$ is observed with a decrease of the temperature in both models (Fig. 10a). Differences are however noticed at low temperature (below -15°C), mainly...
corresponding to Antarctic values, with an isotopic composition that is not depleted enough in our model. Antarctic isotopic values are indeed not computed correctly due to issues in the conservation of water in the advection scheme at very low humidity content, as already highlighted in Roche (2013). We then investigated the relationship between modelled and measured $\delta^2$H$_{\text{precipitation}}$, excluding Antarctic values (Fig. 10b). Most of the values are found between 0 and -60‰, with similar distribution in iLOVECLIM and LMDZ4. Depleted values are however more scattered between the two models (and shifted from the 1:1 line) due to the difference in simulating the isotopic composition at low temperature.

Figure 8: (a) Annual mean modelled $\delta^2$H$_{\text{precipitation}}$ as a function of the temperature for iLOVECLIM (blue) and LMDZ4 (red). (b) Annual mean modelled $\delta^2$H$_{\text{precipitation}}$ for iLOVECLIM and LMDZ4 against observations (without Antarctic values). The 1:1 line is shown with the black dashed line. The errors bars associated with the data are shown at 2σ. The regression curves between model and data are presented in dark blue for iLOVECLIM and red for LMDZ4 with the confidence bands.

3.2.3 Continental effect
The continental effect can be defined by a contrast in isotopic value between land and ocean, with more depleted values over land associated with fractionation during continental recycling. To evaluate this effect in iLOVECLIM, we extracted the monthly isotopic composition of precipitation over land and ocean separately, and focus on the tropics between 0-20°N and 0-20°S. We also extracted values from the LMDZ4 and ECHAM5-wiso models and from the GNIP stations that have at least 3 measurements for each month. The total number of points/stations over the continents and oceans for each model (increasing with a higher resolution of the model) and observation is summarized in the Table 1. Instead of representing all points, we decided to divide each tropical region into three zones for the continents (America, Africa and Asia/Indonesia/Australia) and three zones for the oceans (Atlantic, Pacific, Indian) and calculated the monthly mean for the different zones. We then obtained three series of monthly values corresponding to the continental zones (and similarly for the oceanic zones) for 0-20°N, and for 0-20°S.
The contrast in isotopic value between land and ocean, with more depleted values over land is well observed in the GNIP data for both tropical regions (with a median value of -23 ‰ for the continents and -9.9 ‰ for the oceans in the northern tropics, and -27.9 ‰ vs -6.1 ‰ in the southern tropics, Fig. 11a). Even if most of the climate models do not include this fractionation, they reproduce this shift towards depleted values. This continental effect is observed in iLOVECLIM with a median value of -11.6 ‰ over the continents and of -4.6 ‰ over the oceans for the northern tropics and of -17 and -3.2 ‰ over the continents and oceans respectively in the southern tropics (Fig. 11b). The difference between the land and the ocean is however less pronounced than in the GNIP data with depleted values of 7 ‰ in the model compared to the 13.1 ‰ between 0-20°N for the observations (13.8 vs 21.8 ‰ between 0-20°S). This smaller depletion in the isotopic composition over land is also observed in the LMDZ4 model. The modelled median values for LMDZ4 are similar to these obtained with iLOVECLIM, despite the difference in complexity and processes represented in the atmosphere. Among all three models and surprisingly, ECHAM5-wiso which least reproduces this continental effect, despite being the more complex in the representation of the physical processes in the atmosphere.

<table>
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<tr>
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<th>0-20°N</th>
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<td></td>
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<tr>
<td>iLOVECLIM</td>
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<td>181</td>
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<tr>
<td>LMDZ4</td>
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<td>520</td>
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<tr>
<td>ECHAM5-wiso</td>
<td>4306</td>
<td>5454</td>
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Table 1: Number of GNIP stations and points in the different models that cover land surfaces and oceans in the tropical bands between 0-20°N and 0-20°S.
Figure 9: Box plots of the tropical $\delta^2\text{H}_{\text{precipitation}}$ over the continents (in green) and oceans (in blue). The panels present values from (a) the GNIP data, (b) the iLOVECLIM model, (c) LMDZ4 and (d) ECHAM5-wiso. Values are shown between 0-20°N and between 0-20°S. The horizontal line in the box plots corresponds to the median value.

3.3 Isotopes in ocean water

3.3.1 Surface seawater

The hydrogen isotopic composition has also been modelled in the oceanic component for the sea water. iLOVECLIM models annual mean surface $\delta^2\text{H}_{\text{seawater}}$ with low negative values in the Arctic Ocean, that are mainly enriched compared to the existing data points at high latitudes. (Fig. 12a) This is clearly visible in the zonal distribution (Fig. 13a – with similar methodology than Fig. 2 to take the model outputs that co-locate with the measurements and the use a polynomial regression curve) where the $\delta^2\text{H}_{\text{seawater}}$ trend in iLOVECLIM has too high values for high latitudes compared to the observations and MPI-ESM-wiso. The $\delta^2\text{H}_{\text{seawater}}$ in the Atlantic Ocean is well reproduced in the model with high enriched values close to the tropic and the equator and lower values in the northern and southern part of the ocean, even if the modelled values are slightly different than the observation in the northern Atlantic (Fig. 12a). The Mediterranean Sea presents a good agreement with the
observation with high $\delta^2$H$_{\text{seawater}}$ values. The $\delta^2$H$_{\text{seawater}}$ pattern in the Pacific and Austral oceans is also similar to the observations. However, the western part of the Indian Ocean and Arabian Sea presents lower values of $\sim$10 ‰ in comparison to the GISS data (Fig. 12a). This could be explained by a model bias toward higher precipitations and reduced salinity in this area. Both the iLOVECLIM and the MPI-ESM-wiso models reproduce the zonal distribution from 50°S to 20°N in comparison to the observations. They however present differences, with a generally lower modelled $\delta^2$H$_{\text{seawater}}$ value of the surface water in comparison to the data, and less variability in iLOVECLIM compared to MPI-ESM-wiso (Fig. 13a).

The annual mean surface d-excess in the different oceanic basins is also presented in Fig. 12b with the measurements for comparison. The overall pattern of d-excess is similar to the one of the $\delta^2$H$_{\text{seawater}}$ with high positive values in the Arctic Ocean and lower depleted values in the Atlantic, Pacific, Indian and Austral oceans. The modelled d-excess values from $-2$ to $0$ ‰ in the Atlantic and Pacific oceans match the observations, with a gradient from low to high values from the low to the high latitudes (Fig. 12b and Fig. 13b). The western part of the Indian Ocean and the Arabian Sea again presents different values than the observations. The model calculates a d-excess of $\sim$2 ‰ in the western Indian ocean whereas the data have smaller values. The modelled d-excess even goes up to $14$ ‰ in the Arabian Sea, due to precipitation and humidity effect. Even if a small number of data points exist in the Polar Ocean above 60°N (only few measurements in the Atlantic sector), the model reproduces too high d-excess value in comparison to the observations. This is also the case in the MPI-ESM-wiso model (Fig. 13b). These enriched values could be associated to the absence of sea ice in this simulation, that would lead to fractionation during sea ice formation and depletion of the liquid water isotopic composition. iLOVECLIM also does not include river discharge that are at the origin of depleted isotopic values and could allow for a more depleted d-excess than in our simulation.

The iLOVECLIM model presents however a closer agreement with the measurements from the mid-latitudes to the equator than the MPI-ESM-wiso model (Fig. 13b).

As for $^{17}$O-excess, modelled values are as well highly depleted in the entire Arctic Ocean, Arabian Sea, Mediterranean Sea and along the coast of east and west Africa (Fig. 12c). Apart from the northern part that has negative values similar to the Arctic Ocean, the Atlantic Ocean presents relatively small $^{17}$O-excess variations and match the data with values between 0 and 50 permeg. The Pacific and Indian oceans have higher $^{17}$O-excess values up to 200 permeg, which is higher than observations. However, considering the uncertainties associated with the model and the lack of data does not allow a good model-data evaluation for this proxy.
Figure 12: Model-data comparison of the annual mean isotopic distribution in the ocean. (a) $\delta^2$H of ocean surface water, (b) d-excess of ocean surface water and (c) $^{17}$O-excess of ocean surface water in iLOVECLIM. The model results are compared to measurements (in circles).

Figure 13: Multi-model zonal (a) $\delta^2$H of ocean surface water and (b) d-excess of ocean surface water comparison. The model results (in color) are compared to observations (in grey). The different lines are polynomial regression curves for the model results that co-locate with the observations.

Figure 12: Model-data comparison of the annual mean isotopic distribution in the ocean. (a) $\delta^2$H of ocean surface water, (b) d-excess of ocean surface water and (c) $^{17}$O-excess of ocean surface water in iLOVECLIM. The model results are compared to measurements (in circles).

Figure 13: Multi-model zonal (a) $\delta^2$H of ocean surface water and (b) d-excess of ocean surface water comparison. The model results (in color) are compared to observations (in grey). The different lines are polynomial regression curves for the model results that co-locate with the observations.
3.3.2 Vertical profiles

The model-data comparison of $\delta^2$H and d-excess of sea water can be realized over the entire water column with a cross section in the Atlantic Ocean. We find a general good agreement between the GISS observation and the model from the surface to the bottom with the imprint of the different water masses on the simulated $\delta^2$H (Fig. 14a). The strongest $\delta^2$H enrichment is observed in the upper Atlantic (above 700 m) between 30°S and 45°N with a maximum around 20°N with 4.2 ‰. There are however some differences in the surface water with $\delta^2$H values that are lower than the observations by several permil. Below 700 m, the North Atlantic deep waters (NADW) have lower $\delta^2$H values, between 1.8 and up to 0 ‰ at the bottom of the ocean where they mix with the Antarctic bottom water (AABW) coming from the South with depleted values (Fig. 14a). In the Southern Ocean around 1000 m depth, the Antarctic intermediate waters (AAIW) flow to the north with negative depleted $\delta^2$H values.

The oceanic d-excess shows less prominent influence of the main water masses. Above 1000 m, the d-excess goes from 40°S to 40°N with depleted negative values (Fig. 14b). Below 1000 m and from 40°S to the north, the NADW d-excess values are higher with a maximum of 2 ‰ around 25°N and 2000 m depth. The comparison with the GISS data shows that the model reproduces the depleted surface values and the enriched d-excess values below 1800 m even if the latitudinal gradient is more pronounced in the model than in the data. The depth interval from 500 to 1800 m presents a disagreement between the modelled d-excess and the observation values that are consistently lower than in the model (Fig. 14b). This is especially the case for high latitudes of the northern hemisphere where the difference between the model and the data can reach 2 to 3 ‰.

Figure 14: Atlantic zonal mean in iLOVECLIM of (a) $\delta^2$H of seawater, (b) d-excess of seawater and (c) $^{17}$O-excess of seawater, compared to observations.

The oceanic d-excess and $^{17}$O-excess shows less prominent influence of the main water masses. Above 1000 m, the d-excess goes from 40°S to 40°N with depleted negative values (Fig. 14b), and enriched positive values for $^{17}$O-excess (Fig. 14c). Below
1000 m and from 40°S to the north, the NADW d-excess values are higher with a maximum of 2 ‰ around 25°N and 2000 m depth. On the opposite, δ¹⁸O-excess values are lower than in the surface, with minimum values at the same latitude and depth than d-excess. The comparison with the δ²H and d-excess observations shows that the model reproduces the depleted surface values and the enriched d-excess values below 1800 m even if the latitudinal gradient is more pronounced in the model than in the data. The depth interval from 500 to 1800 m presents a disagreement between the modelled d-excess and the observation values that are consistently lower than in the model (Fig. 14b). This is especially the case for high latitudes of the northern hemisphere where the difference between the model and the data can reach 2 to 3 ‰. Since no δ¹⁸O observations exist at depth, we refrain for any further evaluation of the modelled values.

### Conclusions

In this study, we presented the implementation of the δ¹⁸O/H²¹⁸O, δ¹⁷O/H²¹⁷O isotopologues in the intermediate complexity coupled climate model iLOVECLIM. Based on the existing δ¹⁸O water isotopic module and on this new extension, we modelled the d-excess and δ¹⁷O-excess variations to have a general overview of the water isotopes. We evaluated the model isotopic composition for preindustrial for both the atmosphere and the ocean components based on a long equilibrium simulation. For the atmospheric part, we found a good agreement between the model, the observations and several GCMs, with the conservation of the latitudinal gradient (considering the intrinsic biases of iLOVECLIM that could lead to local inconsistencies). The modelled δ²H and δ¹⁸O fit with the global Meteorological Water Line and the main isotopic effect (amount effect, temperature effect and continental effect are well reproduced in the model). The d-excess distribution for the atmosphere is also correctly modelled at global scale in comparison to the observations and several GCMs. The isotopic composition of oxygen and hydrogen over Antarctica present however differences of several permil in comparison to the data because of the complexity of the local processes at play that are simplified in the model. At present, our models-data comparison suggests that iLOVECLIM does not correctly reproduce the δ¹⁷O-excess that has a too important dispersion of the values. Modelling the δ¹⁷O-excess has to be improved in the future versions of the isotopes-enabled models. New measurements are also needed with a reduction of their associated uncertainties. For the ocean, we reproduced a good accordance of the modelled surface δ²H and d-excess in comparison to the existing data, except for some parts of the Arctic region and local areas in the Indian Ocean. This good agreement is conserved over the entire water column in the Atlantic Ocean, with similar δ²H values and distribution between the model and the data, influenced by the main water masses.

Given the computing resources needed to run coupled climate models, applying intermediate complexity coupled climate models with water isotopes such iLOVECLIM to future long-term palaeoclimate perspectives appear very promising. Palaeoclimate simulations during the Holocene, Last Glacial Maximum or transient glacial/interglacial periods are the next logical step to compare model results against past isotopic composition records. New proxies that depend on the water isotopes...
can also be implemented in the model, like the leaf wax isotopic composition, in order to quantify the influence of the respective factors (precipitation, vegetation, humidity…) that control its variations.

Appendix A: δ¹⁷O isotopic composition

The latitudinal gradient and the global distribution for the modelled δ¹⁷O is similar to the one of the δ¹⁸O with depleted values from the equator to the poles (Fig. A1a). Similarly, the values over land are more depleted than over the ocean. In comparison to the available data over North America, Europe, Asia and Indonesia, iLOVECLIM mostly calculates enriched values of several permil, even if some agreements are observed between the model and the data. These discrepancies can be explained by the fact that the most of the data is punctual and reflect seasonal conditions whereas the model outputs are annual mean δ¹⁷O values.

δ¹⁷O of seawater in iLOVECLIM shows values close to zero over the Atlantic, Pacific, Indian and Southern oceans which is consistent with the observations (Fig. A1b). The amplitude of variation is small and around 1 ‰. The coast of east Africa and the Arabian sea present lower values, as well as the northern part of the Atlantic Ocean and the Arctic Sea with negative values up to -4 ‰.

Figure A1: Mean annual spatial distribution of the iLOVECLIM modelled (a) δ¹⁷Oprecipitation and (b) δ¹⁷O of ocean surface. Model results are compared to observations (in circles).

Figure A2 presents the relationship between modelled and measured δ¹⁷Oprecipitation (excluding values in Antarctica). Most of the values modelled in iLOVECLIM are grouped around enriched isotopic values, but the correlation remains low. The p-value is calculated to 0.079 (>0.05), which is slightly higher than the 5 % limit, indicating a close statistical significant between the two parameters. In comparison to LMDZ4 that is currently the only GCM to include the ¹⁷O (Risi et al., 2013), iLOVECLIM results are in good agreement with most of the values between -2 and -5 ‰, leading to similar linear trend.
between the model and the data. One point with negative value of -8.7 ‰ in LMDZ4 gets closer to the 1:1 line than iLOVECLIM (with -5.8 ‰). However, considering the large confidence intervals for both model results, the modelled δ¹⁷O_precipitation in iLOVECLIM could be in agreement with the values obtained in LMDZ4. The differences between the model results and the data could be related to (1) the uncertainties in measuring this proxy, (2) the fact that most of the data is punctual and reflect seasonal conditions whereas the model outputs are annual mean δ¹⁷O values and (3) the low number of measurements to compare with.

Figure A2: Model-data relationship for the δ¹⁷O_precipitation without Antarctic values for the iLOVECLIM (blue) and LMDZ4 (red) models. The regression curves between model and data are presented in dark blue for iLOVECLIM and red for LMDZ4 with the confidence bands.

Author contributions. TE and TC designed the study. DMR realized the model development. TE performed and analysed the simulations with inputs from TC. TE wrote the paper with contributions from all co-authors.

Competing interest. The authors declare that they have no conflict of interest.

Code availability. The iLOVECLIM source code and developments are hosted at http://forge.ipsl.jussieu.fr/ludus (IPSL, 2023) but are not publicly available due to copyright restrictions. Access can be granted on demand by request to D.M. Roche (didier.roche@lsec.ipsl.fr) to those who conduct research in collaboration with the iLOVECLIM user group.
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