Response of methane emissions from wetlands to the Last Glacial Maximum and an idealized Dansgaard–Oeschger climate event: insights from two models of different complexity

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Abstract. The role of different sources and sinks of CH4 in changes in atmospheric methane ([CH4]) concentration during the last 100 000 yr is still not fully understood. In particular, the magnitude of the change in wetland CH4 emissions at the Last Glacial Maximum (LGM) relative to the pre-industrial period (PI), as well as during abrupt climatic warming or Dansgaard–Oeschger (D–O) events of the last glacial period, is largely unconstrained. In the present study, we aim to understand the uncertainties related to the parameterization of the wetland CH4 emission models relevant to these time periods by using two wetland models of different complexity (SDGVM and ORCHIDEE). These models have been forced by identical climate fields from low-resolution coupled atmosphere–ocean general circulation model (FAMOUS) simulations of these time periods. Both emission models simulate a large decrease in emissions during LGM in comparison to PI consistent with ice core observations and previous modelling studies. The global reduction is much larger in ORCHIDEE than in SDGVM (respectively −67 and −46 %), and whilst the differences can be partially explained by different model sensitivities to temperature, the major reason for spatial differences between the models is the inclusion of freezing of soil water in ORCHIDEE and the resultant impact on methanogenesis substrate availability in boreal regions. Besides, a sensitivity test performed with ORCHIDEE in which the methanogenesis substrate sensitivity to the precipitations is modified to be more realistic gives a LGM reduction of −36 %. The range of the global LGM decrease is still prone to uncertainty, and here we underline its sensitivity to different process parameterizations. Over the course of an idealized D–O warming, the magnitude of the change in wetland CH4 emissions simulated by the two models at global scale is very similar at around 15 Tg yr−1, but this is only around 25 % of the ice-core measured changes in [CH4]. The two models do show regional differences in emission sensitivity to climate with much larger magnitudes of northern and southern tropical anomalies in ORCHIDEE. However, the simulated northern and southern tropical anomalies partially compensate each other in both models limiting the net flux change. Future work may need to consider the inclusion of more detailed wetland processes (e.g. linked to permafrost or tropical floodplains), other non-wetland CH4 sources or different patterns of D–O climate change in order to be able to reconcile emission estimates with the ice-core data for rapid CH4 events.

1 Introduction

Reconstructions from polar ice cores show that the atmospheric CH4 concentration ([CH4]) has varied greatly as a function of past climate changes. Spectral analyses
performed by Loulergue et al. (2008) indicate that the variability in [CH$_4$] over the last 800 kyr is dominated by the 100 kyr glacial–interglacial cycles and by the precessional component of Milankovitch cycles. Suggested underlying mechanisms involve a link between wetland extent and northern ice sheet dynamics as well as between the strengths of tropical sources/sinks and tropical climate patterns, for example through monsoon systems and via the position of the intertropical convergence zone (ITCZ) (Loulergue et al., 2008; Singarayer et al., 2011).

Large uncertainty remains surrounding to what extent the main natural source (wetlands) contributed to the glacial–interglacial change in [CH$_4$], and whilst earlier bottom-up modelling studies could not explain the glacial–interglacial change in [CH$_4$] with a reduction in wetland CH$_4$ emissions alone in response to cooling and change in hydrological cycle (Kaplan et al., 2006; Valdes et al., 2005), more recent studies suggest that a modification in sink strength is neither required (Weber et al., 2010) nor reproduced by atmospheric models suggesting that a modification in sink strength is neither required (Weber et al., 2010) nor reproduced by atmospheric chemistry model simulations (Levine et al., 2011).

Overlaid on the glacial–interglacial changes, the climate of the Pleistocene was also prone to a strong climatic variability on a millennial timescale, the most extreme feature of which is a series of abrupt jumps in Greenland temperature of between 8 and 16 °C over the course of 10–40 yr (e.g. Wolff et al., 2010). This abrupt warming called Dansgaard–Oeschger (D–O) events also have counterparts in the ice-core records (Kaplan et al., 2006; Valdes et al., 2005), more recent studies suggest that a modification in sink strength is neither required (Weber et al., 2010) nor reproduced by atmospheric chemistry model simulations (Levine et al., 2011).}

To our knowledge, only one bottom-up modelling study (Hopcroft et al., 2011) investigated changes in global wetland CH$_4$ emissions during transient D–O events. The results of this study suggest that atmospheric changes driven by modifications of the AMOC induced CH$_4$ variations from natural wetlands that are too small to explain the variations in [CH$_4$] observed in ice-cores during D–O events. But the CH$_4$ emission model in the global vegetation model (the Sheffield DGVM, called SDGVM hereafter) used in Hopcroft et al. (2011) is relatively simple, and the low sensitivity of SDGVM to climate change is one of the reasons advanced by the authors to explain the mismatch with the ice-core data. The scope of the present paper is to perform simulations using the same climate fields from an idealized D–O event as in Hopcroft et al. (2011) but with a process-oriented and recently developed wetland emission model, the ORCHIDEE-WET model (Ringeval et al., 2010, 2011).

An intercomparison of the wetland CH$_4$ emissions simulated during an idealized D–O event between the two models (SDGVM and ORCHIDEE-WET) is performed. Through this intercomparison, our purpose is to evaluate and understand a potential difference of modelled wetland CH$_4$ emission sensitivity to climate change between the models.

Because (i) ORCHIDEE-WET has never been used to simulate the change in wetland CH$_4$ emissions between the LGM and pre-industrial period (PI) and (ii) given the uncertainty remaining of the contribution of wetland emissions to LGM climate conditions, we will perform the intercomparison at first on the LGM–PI difference. Then we will focus on one idealized D–O event.

In Sect. 2, we describe the climate simulations and the two used wetland CH$_4$ emission models. The intercomparison between the two models on both the LGM–PI transition and idealized D–O event is performed in Sect. 3. Finally the results are discussed in Sect. 4.

2 Methods

2.1 Climate simulations

The climate simulated by the Fast Met Office UK Universities Simulator (FAMOUS: Smith et al., 2008) has been...
used to force two global vegetation models (GVMs), namely the Sheffield DGVM (SDGVM) and version of the ORCHIDEE model able to simulate the wetland CH\textsubscript{4} emissions, ORCHIDEE-WET (see Sect. 2.2). FAMOUS is a low-resolution version (5° × 7.5°) of the UK Met Office’s HadCM3 coupled general circulation model. FAMOUS has been used to simulate two equilibrium climates representative of the PI and the LGM. The boundary conditions relevant to the LGM are the 21 kyr orbital configuration, the atmospheric concentrations of major greenhouse gases, the glaciation extent and the sea level, and these follow the PMIP protocol (Braconnot et al., 2007, http://pmip2.lsce.ipsl.fr/). The LGM climate of FAMOUS (Fig. A1) bears many similarities to that simulated by HadCM3 (Singarayer and Valdes, 2010), with a global mean cooling of 4.6 °C, that is similar to HadCM3 and intermediate in the range of cooling magnitudes simulated with other coupled atmosphere–ocean GCMs analysed in PMIP2 (Braconnot et al., 2007). The simulated cooling is more intense over Greenland than in HadCM3, in better agreement with ice-core reconstructions. It also shows warming over the North Pacific and Alaska, the latter in reasonable agreement with terrestrial pollen-based mean annual temperature reconstructions (Bartlein et al., 2010). Over the tropics where proxy-based reconstructions are sparser, differences with HadCM3 are less pronounced, though regional differences in the patterns of the change in the ITCZ and hence precipitation are prominent, particularly in the eastern Pacific and South America. In addition to these two equilibrium simulations, starting from LGM conditions, freshwater forcing has been applied in the Atlantic Ocean in order to perturb the AMOC (Atlantic meridional overturning circulation) and to simulate the space/time patterns of climate during the simulated course of an idealized transient D–O event. During this idealized event, the Greenland temperature is characterized by a cold period (analogous to a Heinrich stadial, HS) then by a warm period (analogous to a Greenland interstadial, GI). The reader is referred to Sect. 2.3 of Hopcroft et al. (2011) for a full description of the FAMOUS simulations and resulting climate. The imposed freshwater forcing, the FAMOUS-simulated AMOC and Greenland temperature are given in Fig. A2. Hopcroft et al. (2011) also performed different simulations modifying the background climate from which the freshwater forcing is applied. Given the larger computational cost of the ORCHIDEE-WET model in comparison to SDGVM, we will use the outputs of only one FAMOUS climate simulation to force the GVMs: the D–O simulation with LGM background, i.e. the reference D–O simulation in Hopcroft et al. (2011).

2.2 Wetland CH\textsubscript{4} emission models

SDGVM and ORCHIDEE-WET have been forced with FAMOUS climate output to simulate the wetland CH\textsubscript{4} emissions during PI, LGM and over an idealized D–O event. SDGVM and ORCHIDEE-WET are two models of different complexity and, regarding wetland CH\textsubscript{4} emissions, have been used for different timescales: paleo-studies for SDGVM (Singarayer et al., 2011; Valdes et al., 2005) and studies over current (Bousquet et al., 2011; Ringeval et al., 2010) or future time period (Koven et al., 2011; Ringeval et al., 2011) for ORCHIDEE-WET.

ORCHIDEE-WET is more process-based than SDGVM in the computation of both the wetland extent dynamic and the CH\textsubscript{4} flux densities. The wetland extent computation is based on a subgrid topographic approach in ORCHIDEE-WET while it corresponds to a simple orographic correction in SDGVM. In ORCHIDEE-WET, the CH\textsubscript{4} flux densities are computed from the process-based Walter et al. (2001) model, which accounts for a soil vertical discretization and for an explicit representation of CH\textsubscript{4} transport from the soil to the atmosphere, whilst in SDGVM transport and vertical discretization are ignored. The wetland extents computed in ORCHIDEE have been evaluated against remote sensing products of inundated area (Ringeval et al., 2012), and the CH\textsubscript{4} flux densities have been optimized with site-level observations (Ringeval et al., 2010). The year-to-year variability of the wetland CH\textsubscript{4} emissions reflects the wetland CH\textsubscript{4} emission sensitivity to the climate variability. The ORCHIDEE-WET simulated year-to-year variability in wetland emissions has been evaluated against top-down estimates over 1990–2000 (succinctly displayed in the Fig. 2 of Ringeval et al., 2011) and is being further analysed over the 1990–2009 period (I. Pison, personal communication, 2012), whilst the interannual variability of SDGVM has not been explored in detail. All of these elements increase our confidence in the modelled wetland emission sensitivity to the climate of ORCHIDEE-WET relative to SDGVM, at least over the current time period. The strategy followed here has been to perform the optimization under the current time period then to apply this model configuration in conditions representative of the LGM and the idealized D–O event.

The SDGVM model has already been described in Hopcroft et al. (2011). Thus, the following paragraphs mainly focus on ORCHIDEE-WET (Sect. 2.2.1) as well as on the differences of methodology used to compute the wetland CH\textsubscript{4} emissions in the two models (Sect. 2.2.2). Similarly, only the ORCHIDEE-WET simulations will be presented (Sect. 2.2.3) and the reader should refer to Hopcroft et al. (2011) for more details of the SDGVM simulations.

2.2.1 The ORCHIDEE-WET model

In the ORCHIDEE-WET model, the wetland CH\textsubscript{4} emissions ($E_{CH_4}$) are computed for each grid cell $g$ and for each time step $t$ through the following equation:

$$E_{CH_4}(g, t) = \sum_{WTD_i} S_{WTD_i}(g, t) \cdot D_{WTD_i}(g, t)$$

with $WTD_i = 0$ and $-3$ cm (1)
where $S_{WTD}$ and $D_{WTD}$ are respectively the extent (given as a grid cell fractional area) and the CH$_4$ flux density of a wetland with a water table depth equal to WTD$_i$. A negative value for WTD$_i$ means that the WTD is below the soil surface. In the present study, not only the emissions of the inundated wetland (i.e. with WTD = 0) are computed as in Ringeval et al. (2011), but also the emissions of wetlands with a mean water table equal to $-3$ cm are also considered. This methodology is close to the one used in Bohn et al. (2007). $S_{WTD}$ and $D_{WTD}$ are computed by the coupling of ORCHIDEE-WET with respectively (i) a TOPMODEL approach and (ii) a slight modification of the Walter et al. (2001) model.

For each grid cell and at each time step, ORCHIDEE-WET simulates a soil water content resulting from a hydrologic budget accounting for some inputs (snowmelt and rainfall not intercepted by the canopy) and losses (soil evaporation, transpiration, sublimation, deep drainage, and surface runoff). This soil water content could be used to express a mean soil water deficit over the grid cell. This deficit is defined as a gap between the simulated soil water content and the maximum soil water content in the model, i.e., the soil field capacity. The coupling between ORCHIDEE-WET and TOPMODEL allows us to distribute the mean soil water deficit over each grid cell as function of the subgrid topographic index distribution. This leads to diagnosis of the fraction of the grid cell with a deficit equal to 0. Then, the inundated wetland extents are computed from these “field capacity extents”. In Ringeval et al. (2011), remote sensing data of inundated extent were used to do this computation. The mean climatology (average of 1993–2000) of the modelled field capacity extents were normalized to the same climatology of Prigent et al. (2007) data, and only the simulated temporal variability was kept. In order to prevent the use of current remote sensing data over paleo-timescale, a parameterization has been introduced. Briefly, the parameterization consists in a shift of the topographic index distribution in each grid cell. The shift value is the same for all grid cells and has been optimized to simulate a current global wetland fraction close to 4% (Prigent et al., 2007) at 1° resolution when forced by the 1960–1991 CRU (http://www.cru.uea.ac.uk/) climatology. The reader is referred to Ringeval et al. (2012) for more details. The coarse-resolution effect on the wetland extent simulation through the coupling between ORCHIDEE-WET and TOPMODEL is illustrated in Fig. A3. In particular, the loss of information in subgrid topographic index distribution due to the coarse resolution (Fig. A3d) leads to substantial difference in the wetland extents when compared to the a posteriori gridded of the 1° simulated wetland extent (Fig. A3b). Nevertheless, the global wetland extent simulated at FAMOUS resolution is close to the value given by Prigent et al. (2007) dataset at the same resolution (Fig. A3c). As in Bohn et al. (2007), the coupling with TOPMODEL has been extended to compute wetland extents with a negative water table depth value. In Eq. (1), $S_{-3\text{ cm}}$ values are taken as extents given by TOPMODEL with a deficit between 0 and $-6$ cm.

The CH$_4$ flux densities are computed using a slight modification of the Walter et al. (2001) model. As in Ringeval et al. (2010), the main modification of the original model concerns the use of the labile carbon pool simulated by ORCHIDEE-WET ($C_L$) to approach the methanogenesis substrate in such a way that the production rate for a soil layer $z$ and a time $t$ ($\text{Prod}(t, z)$) is defined as follows:

$$\text{Prod}(t, z) = \alpha_0 \cdot f_{\text{org}}(z) \cdot C_L(t) \cdot H(T(t, z)) \cdot Q_{10}^{(T(t, z) - T_{\text{ref}})/10}$$

where $f_{\text{org}}$ is a function that vertically distributes the carbon in the soil, $H(T(t, z))$ is the Heaviside step function for the temperature, and $T_{\text{ref}}$ varies in space as in the Walter et al. (2001) model. In ORCHIDEE, the turnover time of the labile carbon pool is equal to 55 days. In the present study, $T_{\text{ref}}$ is defined as the mean surface temperature computed by ORCHIDEE-WET when forced by the 1960–1991 CRU (http://www.cru.uea.ac.uk/) climatology. The parameter $\alpha_0$, which contains both the fraction of the labile carbon pool that could be used as methanogenesis substrate and the base rate at $T_{\text{ref}}$, has been optimized against three sites (Abisko, Michigan or Panama) and then extrapolated at large band scale (Ringeval et al., 2010). As in Ringeval et al. (2011), identification of each grid cell to a wetland type (i.e. sharing the same optimized $\alpha_0$ as Abisko, Michigan or Panama) is based on a criterion of vegetation type. Through this interpolation, the assumption that the base rate at $T_{\text{ref}}$ co-varies with the amount of substrate is implicitly made. The optimization of $\alpha_0$ has been performed using CRU climatology at FAMOUS resolution at monthly time step. A $Q_{10}$ of 3 has been chosen at global scale for the methanogenesis sensitivity to temperature as in Ringeval et al. (2011). This value allows us to match the observed seasonal cycle of CH$_4$ flux densities on both boreal and temperate sites.

### 2.2.2 Differences between SDGVM and ORCHIDEE-WET

Figure 1 summarizes the differences between the CH$_4$ emission parameterizations of the two models. As succinctly mentioned above, the major differences between the two models are related to the computation of both the fractional area covered by a CH$_4$ emitting wetland ($S$) and the CH$_4$ flux densities ($D$). We describe the differences in the case of a given grid cell $g$ at time step $t$.

The methodology in SDGVM leads to a binary estimation of $S$, but allows the simulated WTD in a wetland to vary at monthly timescale from +10 to −10 cm. By contrast, the more process-oriented approach in ORCHIDEE-WET, through its coupling with TOPMODEL, allows $S$ to vary continuously between 0 and 1. Nevertheless, only two WTD classes (0 and $-3$ cm; cf. Eq. 1) are considered in a given grid cell in ORCHIDEE-WET.
In SDGVM, the wetland extent, S, is equal either to 0 or to \( f_{\text{max}} \) depending on a criterion varying with the latitude. This criterion is the value of the surface air temperature in boreal regions: the wetland presence in a given year starts for monthly temperature above 5 °C according to Fung et al. (1991). In the non-seasonally frozen environments, a positive difference between precipitation and evapotranspiration is required to reach \( f_{\text{max}} \). \( f_{\text{max}} \) is equal to the product of the entire area of the wetland and an orographic correction. In the wetland covered fraction \( S \in \{ > 0 \} \) of a grid cell, a WTD value is computed following the Cao et al. (1996) relationship applied to the SDGVM soil water content. As in the Cao et al. (1996) model, the wetlands with the highest water table position allowed by the model (+10 cm) are considered as inundated and the others are called non-inundated wetlands – a distinction determining the CH\(_4\) flux density parameterization.

Regarding the CH\(_4\) flux density, the main differences between SDGVM and ORCHIDEE-WET relate to the soil physics vertical discretization, the accounting for CH\(_4\) transport from the soil to the atmosphere, the proxy of the methanogenesis substrate supply and the parameterization of the methanogenesis sensitivity to the temperature.

In SDGVM, the CH\(_4\) flux density for a given grid cell is estimated by the difference between a production and an oxidation rate. The methanogenesis rate is a function of the surface air temperature (T), the calculated WTD and the heterotrophic respiration through

\[
\text{Prod}(t) = P_0 \cdot R_H(t) \cdot f(WTD(t)) \cdot Q_{10}(T_{\text{ref}}) \cdot Q_{10}^{(T(t) - T_{\text{ref}})/10}
\]

where \( P_0 \) is a constant factor used to compute the base methanogenesis rate from the heterotrophic respiration \( R_H \). Thus, \( R_H \) could be considered as the proxy for the methanogenesis substrate as applied in the Cao et al. (1996) approach. \( f(WTD) \) is equal to 1 in the case of the inundated wetland and decreases exponentially when the WTD decreases. The methanogenesis sensitivity to the temperature is parameterized using a \( Q_{10} \) formulation with a \( Q_{10} = 1.5 \) and a global constant reference temperature \( T_{\text{ref}} = 30^\circ \text{C} \).

The oxidation rate is a given percent of the production (0.9) for non-inundated wetland and a function of gross primary production in the other wetlands. In ORCHIDEE-WET, following Ringeval et al. (2010), the flux density at the atmosphere/surface interface is the result of three processes: production in the soil layers below the WTD, oxidation above the WTD and transport by diffusion, ebullition and through plant aerenchyma. A soil vertical discretization is used as in
the Walter et al. (2001) model. As for SDGVM, a $Q_{10}$ formulation defines the methanogenesis sensitivity to the temperature, but the $Q_{10}$ is here equal to 3 and the reference temperature ($T_{ref}$ of the Eq. 2) varies in space. In ORCHIDEE-WET, oxidation only occurs in the soil layers above the WTD and is also a function of temperature.

The comparison of the results of the two models driven by glacial–interglacial and D–O climate changes will first focus on the change in emissions between different time periods. Then, to better understand the reason of eventual differences between the two models, we will compare the change in the different components of the wetland CH$_4$ emissions (i.e. the wetland extents and the CH$_4$ flux densities per unit of wetland) as in the Eq. (1). To compare more easily each component between the two models, we will compute them in the case of a saturated wetland. Thus, we will compare between the two models both the saturated wetland extent ($S_0$) and the CH$_4$ flux density for a saturated wetland ($D_0$). These two variables are direct outputs of ORCHIDEE-WET (WTD = 0 in Eq. 1). For SDGVM, the saturated wetland extents are defined a posteriori as the wetland extents with a water table depth above the soil surface (and below +10 cm, which is the prescribed maximum value). They encompass the so-called inundated wetlands by Cao et al. (1996) and the non-inundated wetlands with a WTD between +10 cm and 0. The 0 value is used as a threshold, because it is the maximum value allowed by the TOPMODEL approach (following Saulnier and Datin, 2004) used in ORCHIDEE-WET. For SDGVM, the CH$_4$ flux densities for a saturated wetland are approached by dividing the simulated CH$_4$ flux densities by $f$(WTD), i.e. by the function used to decrease the potential methanogenesis rate depending on the WTD value.

### 2.2.3 The ORCHIDEE-WET simulations

The boundary conditions for the ORCHIDEE-WET simulations concern the soil texture (fractions of sand, silt and clay), the vegetation distribution and the orography/topography.

The soil texture maps come from the ISLSCP data (http://badc.nerc.ac.uk/data/islscp/hydro.html) and are the same as the one used as input of SDGVM in Hopcroft et al. (2011). In ORCHIDEE-WET, a given grid cell represents the heterogeneous vegetation using a “mosaic” of 10 natural plant functional types (PFTs) and bare soil. The fraction of the grid cell occupied by each PFT is either calculated (and thus variable in time) or prescribed (Krinner et al., 2005). In the present study, dynamic vegetation is not activated; thus, vegetation maps are used as fixed boundary conditions. However, this does not prevent accounting for wetland extent dynamics (see above), which is decoupled from the vegetation dynamics in the ORCHIDEE-WET model. For the PI period, the vegetation map from the HYDE 3.0 database (Klein Goldewijk et al., 2007) is used. The prescribed LGM vegetation comes from Woillée et al. (2011). Contrary to SDGVM, the LGM vegetation is static during the entire transient D–O simulations in ORCHIDEE-WET.

The mean altitude of each grid cell is taken from the orography boundary condition used in FAMOUS and is derived from the ICE-5G data (Peltier, 2004). The mean altitude is used to derive the surface atmospheric pressure. The altitude is considered constant during the D–O run. Concerning the subgrid topography necessary as input of ORCHIDEE-WET through its coupling with TOPMODEL, we use the current subgrid topography given by HYDRO1k (http://webgis.wr.usgs.gov/globalgis/metadata/hydro1k.htm) for all the simulated time periods. Thus, it is implicitly assumed that the large-scale (i.e. the mean altitude of each grid cell) and the small-scale topography are independent. As described in Decharme and Douville (2007) and Ringeval et al. (2012), the spatial distribution of the topographic indices in each grid cell is derived from the mean, standard deviation, and skewness of the actual distribution using a three parameter gamma distribution. For the grid cells for which the continental fraction increases from PI to LGM, the same statistical variables as for the PI are used to extend the subgrid distribution to the new land part of the grid cell. For entirely new grid cells under LGM conditions, we use the subgrid topography distribution of the closest grid cell existing under PI conditions. This strategy had to be applied only for few grid cells as compared to the previous case (increase of continental fraction in PI existing grid cells). An optimal alternative would have been to use the bathymetry data (e.g. http://www.ngdc.noaa.gov/mgg/bathymetry/relief.html) for these new grid cells (J. O. Kaplan, personal communication, 2011). No changes in the topography/orography are accounted for during the transient D–O run for the two models, and they are fixed to LGM conditions. Note also that the sea level is considered constant during the D–O transient run as in Hopcroft et al. (2011).

A spin-up run of several thousand of years was performed to bring all ORCHIDEE-WET carbon pools to their long-term equilibrium values for both the PI and LGM conditions. A 30-yr interannual simulation was then carried out for both the PI and the LGM with year-to-year variability deriving from FAMOUS and is used to perform the intercomparison with SDGVM.

In the present study, three ORCHIDEE-WET configurations named hereafter V0, V1 and V2 have been used (see Table 1). V0 is the standard ORCHIDEE-WET set-up. The aim of the two other configurations is either to estimate the contribution of different parameterizations to potential differences between ORCHIDEE-V0 and SDGVM, or to better understand the wetland CH$_4$ emission sensitivity to climate in ORCHIDEE-WET.

V1 differs from V0 by a change in the parameterization of the methanogenesis sensitivity to temperature. The same sensitivity as in SDGVM is used in V1. It corresponds to a spatially invariant $T_{ref}$ equal to 30°C and $Q_{10} = 1.5$. As

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Table 1. Description of the different ORCHIDEE-WET simulations (ORCHIDEE-V0, V1, V2 and opt).

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<th>ORCHIDEE-WET – V0</th>
<th>ORCHIDEE-WET – V1</th>
<th>ORCHIDEE-WET – V2</th>
<th>ORCHIDEE-WET – opt</th>
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<tr>
<td>General description</td>
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<td>As ORCHIDEE-WET – V0 with the same methanogenesis sensitivity to the temperature as SDGVM</td>
<td>As ORCHIDEE-WET – V1 with prescribed soil water content to compute the CH$_4$ flux densities</td>
<td>“Optimal” version</td>
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<tr>
<td>$Q_{10}/T_{\text{ref}}$</td>
<td>$Q_{10} = 3$ $T_{\text{ref}} = 30^\circ\text{C}$ everywhere</td>
<td>$Q_{10} = 1.5$ $T_{\text{ref}} = 30^\circ\text{C}$ everywhere</td>
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<td>$Q_{10} = 3.0$ $T_{\text{ref}} = 30^\circ\text{C}$ everywhere</td>
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<tr>
<td>Way to compute $\alpha_0$ and value for the different latitude bands (temperate, boreal, tropical) (in $10^{-6}$ m$^2$ month$^{-1}$)</td>
<td>Optimized against three sites then extrapolated at latitude band scale as in Ringeval et al. (2010) (1.1, 2.2, 17.5)</td>
<td>Optimized against three sites then extrapolated at latitude band scale as in Ringeval et al. (2010) (5.5, 8.5, 20.1)</td>
<td>For each sites, equal to $\alpha_0$(ORCHIDEE-V1) $C_{\text{sol}}$(ORCHIDEE-V2)/$C_{\text{sol}}$(ORCHIDEE-V1) then extrapolated at latitude band scale (6.9, 5.4, 24.5)</td>
<td>For each sites, equal to $\alpha_0$(ORCHIDEE-V1) $C_{\text{sol}}$(ORCHIDEE-Vopt)/$C_{\text{sol}}$(ORCHIDEE-V1) then extrapolated at latitude band scale (34.8, 21.1, 37.6)</td>
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<tr>
<td>Soil water conditions used to compute the CH$_4$ flux densities</td>
<td>Computed by the model</td>
<td>Computed by the model</td>
<td>Prescribed: constant in time and space and equal to the maximum soil water content in the model</td>
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explained in Sect. 2.2.1, the parameter $\alpha_0$, which represents the fraction of the labile carbon pool that could be used as methanogenesis substrate, accounts also for a methanogenesis base rate at $T_{\text{ref}}$. A change in $T_{\text{ref}}$ and $Q_{10}$ requires a new optimization of $\alpha_0$ that we performed on the same three sites as for ORCHIDEE-V0 (see Table 1). The fact that $T_{\text{ref}}$ is constant could lead to differences in the spatial distribution of emissions in each latitude band sharing the same $\alpha_0$ value as compared to V0. Through the V1 simulation, we aim to estimate the role played by the $Q_{10}$ formulation on the difference between SDGVM and ORCHIDEE-V0. V1 is closer to SDGVM than V0. Note, however that using such a low $Q_{10}$ value in ORCHIDEE-WET leads to poor agreement between the simulated and observed seasonal cycles of CH$_4$ flux densities for present-day site-level observations (not shown).

Finally, ORCHIDEE-V2 differs from V1 through the prescription of the maximum soil water content in each grid cell and at each time step to compute the carbon cycle. That means we fixed the soil moisture at its field capacity everywhere regardless of the soil water budget, in the computation of the different carbon pools. The computation of the wetland extent is not affected by this operation: the ORCHIDEE-V1 modelled wetland fractions are combined with such CH$_4$ flux densities to compute the CH$_4$ emissions. Moreover, the values of the water table used to compute the CH$_4$ flux densities are the same in V0 and V1 (cf. Eq. 1). However, prescribing maximum soil water content during the carbon cycle computation allows removing what we consider as a bias of the methodology used in ORCHIDEE-WET (see also Sect. 4). In fact, in ORCHIDEE-WET, a subgrid approach (TOPMODEL) is used to diagnose the wetland fraction of each grid cell. But this subgrid treatment has no effect on the carbon cycle computation and in particular on the moisture dependence of decomposition. Indeed, there is no wetland PFT and thus no subgrid wetland/non-wetland distinction to compute the carbon cycle variables. Instead of this, the mean value of the labile soil carbon content over the grid cell ($C_L$ in the Eq. 2) is used as the wetland substrate. Thus the methanogenesis substrate is sensitive to change in precipitation in the model while it would be less sensitive in
The reality for a continually saturated wetland. The strategy used in V2 allows us to treat each grid cell as a saturated wetland and to remove the effects of the temporal variability in the soil water content on the carbon cycle variables necessary to compute the CH$_4$ flux densities. This leads to a modification of the latitudinal distribution of the wetland CH$_4$ emissions as compared to V1 and in particular to lower boreal emissions (cf. Fig. 2). $a_0$ is not again optimized on sites, but a correction is applied to the V1 value (cf. Table 1). Note finally that using constant soil field capacity conditions has an effect not only on the substrate but also on the surface temperature and on the net primary productivity (NPP), which could both additionally modify CH$_4$ emissions, through the methanogenesis and transport respectively. However, these effects are estimated to be of second order in comparison to the effect on the substrate (not shown).

In order to better compare our simulation results with wetland CH$_4$ emission estimates from ice-core data, a final ORCHIDEE simulation, hereafter called ORCHIDEE-opt, has been performed. In this simulation, as in ORCHIDEE-V2, the substrate sensitivity to precipitation is removed. However, a space-constant $T_{ref}$ is used as well as a $Q_{10}$ equal to 3. The aim of the previous configurations (V1 and V2) was either to estimate the contribution of different parameterizations to potential differences between ORCHIDEE-V0 and SDGVM, or to better understand the wetland CH$_4$ emission sensitivity to climate in ORCHIDEE-WET. The aim of ORCHIDEE-opt is to provide our best ORCHIDEE estimates of the change in wetland emissions. These estimates will be exclusively discussed in Sect. 4.

3 Results

3.1 LGM–PI

3.1.1 Magnitude and latitudinal distribution of the LGM–PI change in emissions

The basic parameterization of the two models leads to larger simulated PI emissions in ORCHIDEE-WET than in SDGVM (275 vs. 197 Tg yr$^{-1}$; Table 2) but with a similar latitudinal distribution at FAMOUS resolution (Fig. 2, Fig. 3, left side). The ORCHIDEE-WET PI emissions are slightly higher than previous estimates (e.g. Chappellaz et al., 1993). This apparent over-estimation against commonly accepted values was also obtained over the period 1990–2000 (Ringleval et al., 2011). Contrary to some other studies (e.g. Spahni et al., 2011), the global ORCHIDEE-WET emissions had not been calibrated to match other estimates. Instead of this, the model has been independently tuned to reproduce the wetland extent against remote sensing data and the CH$_4$ flux densities against sites measurements. This underlines the uncertainty linked to the contribution of the wetlands to the global CH$_4$ budget (Melton et al., 2012a and S. Kirschke, personal communication, 2012).

Both (i) the global magnitude and (ii) the latitudinal distribution of the LGM–PI change are different between SDGVM and ORCHIDEE-V0. The global decrease of emissions in LGM as compared to PI is higher in ORCHIDEE-V0 than in SDGVM (respectively, −67 vs. −46 %) (cf. Table 2 for values in both Tg yr$^{-1}$ and percent). Both models lead to higher decrease during LGM than the range given by Weber et al. (2010) (35–42 %), which focused on the effect of the uncertainty in the LGM climate modelling on the wetland CH$_4$ emissions using a very simple wetland CH$_4$ emission parameterization (see Sect. 4). Here the lower LGM wetland CH$_4$ emissions simulated by ORCHIDEE-WET could completely explain the observed change in [CH$_4$] as suggested by previous top-down studies, e.g. Chappellaz et al. (1997) and Dallenbach et al. (2000), while these approaches have their own limitation (see e.g. Baumgartner et al., 2012).

In ORCHIDEE-V0, the LGM decrease of CH$_4$ emissions in northern latitudes (> 30° N) is higher than the decrease in tropics (30° S–30° N) while they are of the same magnitude in both latitude bands in SDGVM (Table 2; Fig. 2, bottom panel). Boreal wetland emissions are almost shut down in ORCHIDEE-WET (decrease of 88–97 % in emissions northwards of 60° N), which seems to be in agreement with the large drop of boreal wetland emissions in LGM inferred by Fischer et al. (2008) using CH$_4$ isotopic information from ice cores.
Table 2. PI, LGM and LGM–PI wetland CH₄ emissions for SDGVM and ORCHIDEE. In the top row, the global PI and LGM emissions are given in Tg yr⁻¹. For ORCHIDEE-WET, the first number in brackets corresponds to the emissions from saturated wetland while the second number refers to the emissions from non-saturated wetlands. In the bottom row, the LGM–PI change (in percent) is given for different latitudinal bands. In the bottom row, numbers in square brackets correspond to tests performed to evaluate the sensitivity to change in latitudinal distribution due to modification of α₀ (please refer to the text in Sect. 3.1.1).

<table>
<thead>
<tr>
<th>Tg yr⁻¹</th>
<th>SDGVM</th>
<th>ORCHIDEE-WET V0</th>
<th>ORCHIDEE-WET V1</th>
<th>ORCHIDEE-WET V2</th>
<th>ORCHIDEE-WET opt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Global PI</td>
<td>197</td>
<td>275 (203 + 72)</td>
<td>259 (191 + 68)</td>
<td>236 (174 + 62)</td>
<td>229 (168 + 61)</td>
</tr>
<tr>
<td>Global LGM</td>
<td>106</td>
<td>90 (65 + 25)</td>
<td>128 (92 + 36)</td>
<td>161 (117 + 44)</td>
<td>146 (106 + 40)</td>
</tr>
<tr>
<td>LGM–PI (%)</td>
<td>SDGVM</td>
<td>ORCHIDEE-WET V0</td>
<td>ORCHIDEE-WET V1</td>
<td>ORCHIDEE-WET V2</td>
<td>ORCHIDEE-WET opt</td>
</tr>
<tr>
<td>Global</td>
<td>−46 %</td>
<td>−67 %</td>
<td>−51 % [−50 %]</td>
<td>−32 % [−35 %]</td>
<td>−36 % [−38 %]</td>
</tr>
<tr>
<td>&gt; 30° N</td>
<td>−41 %</td>
<td>−87 %</td>
<td>−75 % [−73 %]</td>
<td>−52 % [−64 %]</td>
<td>−45 % [−58 %]</td>
</tr>
<tr>
<td>30° S–30° N</td>
<td>−48 %</td>
<td>−57 %</td>
<td>−39 % [−39 %]</td>
<td>−25 % [−23 %]</td>
<td>−32 % [−30 %]</td>
</tr>
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</table>

Fig. 3. Left panels: PI emissions (Tg yr⁻¹) for SDGVM (a) and for the different ORCHIDEEWET versions (c, e, g). Right panels: LGM-PI change (Tg yr⁻¹) for each model (b: SDGVM and d, f, h: different ORCHIDEE-WET versions). The same scaling as for Fig. 2 is applied to the ORCHIDEE-WET plots. Grey areas correspond to grid cells without any vegetation.
The different LGM–PI change at global scale between the two models could be reconciled by prescribing the same $Q_{10}$ formulation to quantify the methanogenesis sensitivity to temperature in each model (Table 2, SDGVM and ORCHIDEE-V0). This underlines the large consequence of the uncertainty relating to particular key parameters. Contrary to the global magnitude of the LGM–PI difference, the latitudinal distribution of this difference cannot be easily reconciled between the two models: the decrease in the > 30° N region is higher than the one in 30° S–30° N whatever the ORCHIDEE-WET version and in contrast with SDGVM (Table 2). The amount of $\alpha_0$ change from one ORCHIDEE version to the other one is not the same for the three big latitude bands sharing the same $\alpha_0$ parameter. This could contribute to modify the contribution of each latitude band to the global signal. A posteriori sensitivity tests have been performed by applying correcting scaling factors to obtain, in each version, exactly the same contribution of each big latitude band to the global PI emissions as in V0. The LGM–PI difference has been computed using such “correcting” emissions and is indicated in Table 2. This correction then ensures that the primary influence of the change between the versions is not related to change in $\alpha_0$.

### 3.1.2 Factors explaining the difference in the LGM–PI change in emissions between the two models

To explain the differences between the two models, we examine the two components of the emissions as explained in Sect. 2.2.3: the saturated wetland extent ($S_0$) vs. the CH$_4$ flux density for a saturated wetland ($D_0$). We focus also on the drivers of the sensitivity of each component to the climate. At first, Fig. 4 shows the components of the PI emissions for both models (top panels for SDGVM and bottom ones for ORCHIDEE). The contribution of each component to the PI emissions is very different between SDGVM and ORCHIDEE-WET (Fig. 4). Indeed, the mean yearly PI $D_0$ over the globe for SDGVM is about half the ORCHIDEE-V0 value (respectively 47.4 and 87.5 g CH$_4$ m$^{-2}$ yr$^{-1}$). Given the relatively similar global PI emissions between the two models (Fig. 2), the opposite relationship is obtained for the mean saturated wetland component ($S_0$). In the Figs. 5 and 6, the LGM–PI change of each component will be expressed in percent of its PI value because of the difference of PI value between the two models.

The role played by the changes in wetland extent in explaining the LGM–PI difference in emissions is weak in the two models (Figs. 3 and 5). Thus, the impact of the additional complexity of the hydrological scheme employed in ORCHIDEE-WET compared to in SDGVM has only a limited effect on the LGM–PI difference in CH$_4$ emissions. The change in wetland extent between LGM and PI is partially due to the change in both continental ice sheets (decrease of land area available for wetlands) and continental shelves (increase in land area available for wetlands), which are named “geographic effects” in Weber et al. (2010). The contribution of the “geographic effects” to the change in emission is close in the two models. Indeed, the gain of emissions during PI over areas covered by continental ice-sheets during LGM is 16 and 19 Tg yr$^{-1}$ for SDGVM and ORCHIDEE, respectively. In the same way, the loss of emissions during PI due to shrinking continental shelves area (higher sea level) is of 13 and 11 Tg yr$^{-1}$ for SDGVM and ORCHIDEE,

![Fig. 4. Components of the PI emissions for each model (SDGVM and ORCHIDEE-V0).](image-url)
respectively. These effects nearly oppose each other, and together do not contribute to the global difference between the two models. However, the simple treatment used to estimate the subgrid topography of the new land surface during the LGM (i.e. the extrapolation from nearby land grid cells) does not allow a comprehensive analysis of the role of coastal shelf regions in LGM–PI wetland CH₄ emissions.

The lower LGM emissions in boreal regions in ORCHIDEE-WET as compared to PI values are mainly explained by a drop in the CH₄ flux densities (Fig. 5). In ORCHIDEE, the main driver of the LGM–PI CH₄ flux densities change is the decrease from PI to LGM in substrate supply (Fig. 6). Indeed, the effect of the change in temperature on the methanogenesis rate is low (compare ORCHIDEE-V0 and ORCHIDEE-V1 in Fig. 5). The large decrease in the substrate availability computed by ORCHIDEE in LGM as compared to PI is not simulated by SDGVM and explains the difference of behaviour between the two models. Between the two equilibrium states (LGM and PI), the change in substrate supply is caused by a change of input, reflecting change in the NPP. In ORCHIDEE, the large decrease in NPP in LGM seems to be mainly driven by a change in NPP flux density per vegetation type rather than by a change in vegetation coverage (Fig. A4a–b). In particular, the increase in summer vegetation moisture stress in ORCHIDEE-WET from PI to LGM is a major contributor to the decrease in NPP (see Fig. A4). Soil freezing processes that limit the availability of liquid water to plants are accounted for in ORCHIDEE.
and not in SDGVM, which could explain the lower NPP and substrate availability in ORCHIDEE compared to SDGVM. Note that the contribution of climate vs. CO\textsubscript{2} in the LGM–PI change in NPP has not been compared between the two models. The contribution of each process in ORCHIDEE is discussed in Woillez et al. (2011, Fig. 15) while the model version used in this latter study does not include representation of soil water freeze/thaw.

It seems that the difference of chosen proxy for the substrate between the two models (heterotrophic respiration (HR) in SDGVM vs. labile carbon pool in ORCHIDEE) plays a minor role: in ORCHIDEE, the LGM–PI change in HR is similar to the change in active carbon pool (not shown).

3.2 D–O events

3.2.1 Change in emissions at global scale and contribution of the tropics vs. extra-tropics

We now analyse the changes in the wetland CH\textsubscript{4} emissions over one idealized D–O event and in particular during two key-periods relative to the LGM: (i) the cold period corresponding to the AMOC off phase and here denoted as analogous to a Heinrich stadial (HS) and (ii) the warm period corresponding to the strong overturning behaviour and denoted as analogous to a Greenland interstadial (GI). The LGM, the HS and the GI periods are respectively delimited by the following transient simulation years: 1–30, 151–180 and 301–330.

The amplitude of the change in wetland CH\textsubscript{4} emissions between the warm and cold periods of the D–O is very similar between the two models (SDGVM and ORCHIDEE-V0, Fig. 7, top panel). During the cold period of the D–O (HS), the wetland CH\textsubscript{4} emissions are 5.7 % lower in comparison to LGM values for SDGVM, while this decrease is of 3.3 % for ORCHIDEE. The change during the warm period of the D–O (GI) relative to the LGM value is slightly larger in ORCHIDEE-V0 than in SDGVM (respectively +11.5 and +8.5 %) leading to very similar GI–HS differences for the two models (+14.2 for SDGVM and +14.8 % for ORCHIDEE). Despite this consistent global picture in the two models, the contribution of each latitude band to the global D–O change (Fig. 7, bottom panel) is very different between ORCHIDEE and SDGVM. The northern regions (> 30° N) play a minor role in the global signal in ORCHIDEE-V0 contrary to the situation in SDGVM. For instance, this latitude band explains 53 % of the global GI–LGM difference in emissions in SDGVM and only 28 % in ORCHIDEE-V0. The difference between the two models regarding the latitudinal distribution of the D–O change in emissions can be totally explained by the much lower LGM emissions of the > 30° N band in ORCHIDEE-V0 than in SDGVM. Indeed, the variation of the extra-tropical latitude band normalized by the LGM boreal emissions is higher in ORCHIDEE-V0 than in SDGVM (not shown). In the later case, the higher boreal emission sensitivity to D–O climate changes obtained with ORCHIDEE-V0 is explained by its higher $Q_{10}$ value (3 against 1.5 in SDGVM).

Concerning the tropics, ORCHIDEE-V0 shows a HS–LGM change of the same amplitude as SDGVM but a larger increase in the emissions during GI relative to LGM (+4.2 for SDGVM and +8.2 % for ORCHIDEE-V0). Moreover, it seems the tropical HS–LGM change in ORCHIDEE-WET is very sensitive to the different parameterizations (Fig. 7; differences between V0, V1 and V2) with, surprisingly, a positive HS–LGM difference simulated by ORCHIDEE-V1. Thus, while we obtain the same picture at global scale for the two models, the underlying drivers of the sensitivity of each model are not the same, in particular in the tropics. To investigate this, we now examine the contribution of each component (CH\textsubscript{4} flux densities vs. wetland extent) to the modelled emission change between LGM, HS and GI.

3.2.2 Drivers of the change in emissions

We have carried out sensitivity analysis to assess the contribution of wetland extent vs. CH\textsubscript{4} flux density to the
Fig. 7. Evolution of the CH$_4$ emissions during D–O events at global (top panel), extra-tropical (> 30° N, middle) and tropical (bottom panel) latitude bands (in percent of the global LGM emissions) and computed using moving mean over 20 yr. The evolution of global emissions over the D–O event simulated by the ORCHIDEE-opt version is added as a dashed orange curve. Difference in CH$_4$ emissions between LGM, HS and GI. In each latitude band, we computed the annual CH$_4$ emission anomalies relative to the mean global LGM value for the HS and GI periods using the simulations described above. These anomalies are called VAR in the following. We have also computed the annual CH$_4$ emission anomalies in the case where the wetland extent is prescribed and equal for each grid cell to its mean LGM value in the respective ORCHIDEE and SDGVM simulation. These CH$_4$ emission anomalies are denoted as FIXED. Figure 8 displays scatter plots of FIXED against VAR. In Fig. 8, the two triangles delimited by the x-axis and the 1 : 1 line encompass model behaviour in which both the flux density and the wetland extent anomalies have the same sign. In these triangles, the closer a given point is to the x-axis, the higher the contribution of wetland extent in the emission anomaly. For points that fall outside of these two triangular areas, the models are showing competing influence of wetland area vs. CH$_4$ flux densities.

At the global scale (symbol □ in Fig. 8), the role played by the change in wetland extent in the emission anomaly in SDGVM is smaller than in ORCHIDEE-V0 for both the
warm and cold D–O periods (the symbols are closer to the 1 : 1 line in SDGVM than in ORCHIDEE). The contribution of the changes in wetland extent is small in the two models over the boreal region (⊙) and cannot explain the difference of behaviour between SDGVM and ORCHIDEE-V0 at global scale. Indeed, for each model and in the extra-tropical regions, VAR and FIXED are close. Thus, much of the difference between the two models at global scale is driven by differences in the tropical regions.

In the tropics (△), during HS (blue symbols), accounting for the wetland extent variability leads to a reduction of the CH₄ emission anomaly from −7.1 ± 3.3 (FIXED) to −0.9 ± 4.4 Tg yr⁻¹ (VAR) in ORCHIDEE-V0 and to a change in the emissions from −2.2 ± 3.4 (FIXED) to −2.08 ± 3.4 Tg yr⁻¹ (VAR) in SDGVM. In ORCHIDEE-V0, the HS climate change (in comparison to LGM) leads to both increased emitting areas and decreased flux densities at the same time, with opposite effect on the total CH₄ flux entering the atmosphere. The tropical band can be divided into two sub-regions: the 0–30° N and the 30° S–0 latitudes bands. In each model, these two sub-regions have a very different behaviour regarding the CH₄ emission anomalies during HS: the southern tropical band is characterized by an increase of the CH₄ emissions, while the northern band sees a decrease. This is related to a southward shift in the ITCZ simulated by FAMOUS in response to the AMOC perturbation (see Hopcroft et al., 2011 for more details). SDGVM and ORCHIDEE mainly differ in terms of (i) the intensity of the emission anomaly in each sub-region and (ii) the contribution of the wetland extent in the emission anomaly of the southern tropical band. Indeed, in ORCHIDEE-V0, the HS emission anomalies reach −11.6 ± 1.8 and +10.3 ± 3.8 Tg yr⁻¹ for respectively the northern (▽) and southern tropics (⊙) while they are only about of −6.3 ± 2.4 and +3.1 ± 3.0 Tg yr⁻¹ in SDGVM. In the southern tropics, the wetland extent explains around 90 % of the increase in emissions in ORCHIDEE-V0 and only 35 % for SDGVM. These two characteristics show that, while the same HS anomaly is obtained in the two models for the entire tropical band due to a compensating effect, the underlying processes are different. Moreover, Fig. 8 also demonstrates that the HS CH₄ flux density anomaly in the northern tropics (blue ▽) in ORCHIDEE-V0 is mainly due to the substrate sensitivity to change in precipitation (see the difference of the FIXED values between ORCHIDEE-V0 and ORCHIDEE-V2). In fact, the decrease in precipitation occurring in the northern tropics during HS leads to a decrease in the ORCHIDEE-simulated NPP leading to a drop of the methanogenesis substrate supply. Besides, note that accounting for the dynamic in vegetation in SDGVM during the D–O run has a small impact on the change in productivity and could not explain differences between the two models.

Figure 8 allows us to identify which latitudinal band and process drives the main differences between SDGVM and ORCHIDEE as described in Sect. 3.2.1: the larger positive GI emission anomaly in ORCHIDEE-V0 than in SDGVM and the positive HS emission anomaly obtained with ORCHIDEE-V1.

Regarding the CH₄ emissions during the GI period, the difference between the two models described in Sect. 3.2.1 is driven by the changes in the band 0–30° N (▽). The emission anomaly in this region is about 5.3 ± 2.9 Tg yr⁻¹ where 88 % can be explained by the expansion of wetland in ORCHIDEE-V0 against 2.5 ± 2.6 Tg yr⁻¹ and 63 % in SDGVM.

The positive tropical anomaly obtained in ORCHIDEE-V1 is explained by the fact that a weak change in the magnitude of the anomaly of a given tropical sub-region could strongly modify the net magnitude over the tropics as a whole, given the compensating effect described above. The Fig. 8 shows that the change in the temperature sensitivity formulation from V0 to V1 leads to a small decrease of the positive anomaly in the northern tropics (from −8.7 ± 1.8 to −6.9 ± 1.3 Tg yr⁻¹ for FIXED, i.e. a decrease of ~20 %) but with no modification in the southern tropics. This small change is strong enough to disrupt the balance of the compensating effects between the two sub-tropical regions, resulting in a positive anomaly averaged over the tropics as a whole. This underlines an increased sensitivity in ORCHIDEE-WET compared to SDGVM and thus a potential larger sensitivity to smaller local changes, while this is without any substantial change at global scale in the case described above.

4 Discussion and conclusions

Weber et al. (2010) quantified the effect of the uncertainties linked to the LGM climate on the different factors controlling glacial changes in CH₄ production by wetlands. To do so, they used different OAGCM outputs to force a unique and simple formulation of CH₄ production. They found that global methane emissions from wetland were reduced by 35–42 % during the LGM in comparison to the PI. Such a reduction is larger than calculated in earlier bottom-up approaches (between −16 and −27 % for Kaplan et al., 2006; Kaplan, 2002; Valdes et al., 2005) and is attributed to differences in the LGM climate simulations (PMIP2 vs. PMIP1: Braconnot et al., 2007). The LGM reduction found by Weber et al. (2010) is closer to the range of reduction found in studies based on top-down modelling (e.g. Crutzen and Brühl, 1993; Martinerie et al., 1995; Chappellaz et al., 1995; Levine et al., 2011). These studies constrained multi-dimensional chemical transport models with ice core observations and inferred the source terms, finding a reduction in the LGM wetland CH₄ emissions by 40–60 %. The present study deals with a complementary approach to Weber et al. (2010) by using the same climate forcing as input for two different wetland CH₄ emission models. We found a decrease of 46 and 67 % at global scale for respectively SDGVM and the base
ORCHIDEE-WET model (version V0). In order to better compare our simulation results with wetland CH₄ emission estimates from ice-core data, we performed a last ORCHIDEE simulation, hereafter called ORCHIDEE-opt as described in Sect. 2.3. For remaining, in ORCHIDEE-opt, as in ORCHIDEE-V2, the substrate sensitivity to precipitation is removed. However, a space-constant $T_{\text{ref}}$ is used as well as a $Q_{10}$ equal to 3. The aim of ORCHIDEE-opt is to provide our best ORCHIDEE estimates of the change in wetland emissions. This version leads to a decrease in LGM as compared to PI of 36%. Thus, the LGM decrease in emissions simulated by both SDGVM and the optimal ORCHIDEE version is close to the lower limit of the range given by top-down studies as discussed above.

The differences between CH₄ concentrations in Greenland and Antarctica as well as the CH₄ isotopic information from ice cores are additional constraints used in top-down modelling to derive the latitudinal change of (wetland) emissions between LGM and PI. Chappellaz et al. (1997) (based on the inter-hemispheric gradient) and Fischer et al. (2008) (based on isotopic information) lead to the same conclusion of a large decrease of boreal wetland emissions during LGM but do not agree on the magnitude of this decrease. The latitudinal distribution of the LGM–PI difference given by SDGVM is consistent with Chappellaz et al. (1997) (−57% for latitudes > 30° N) while the simulated shut-down of boreal wetland emissions in ORCHIDEE-WET is in agreement with Fischer et al. (2008).

New high-resolution CH₄ records from Greenland and Antarctica suggest the boreal wetlands were not completely shut down (Baumgartner et al., 2012). Using the SDGVM and ORCHIDEE simulations, we computed the relative inter-polar concentration difference of CH₄, noted rIPD hereafter and defined by Eq. (9) of Baumgartner et al. (2012):

$$\text{rIPD} = \frac{2 \cdot (s_n - s_s)}{s_n + s_s} \cdot \frac{1}{1 + 2 \frac{t_{\text{ex}}}{\tau}} \quad (4)$$

where $s_n$ and $s_s$ are the CH₄ source for respectively the Northern (0° N–90° N) and the Southern (0° S–90° S) Hemispheres, $\tau$ is the atmospheric lifetime of CH₄ and $t_{\text{ex}}$ is the interhemispheric mixing time. Figure 9a displays LGM rIPD using SDGVM and ORCHIDEE for different value of $\tau$ and $t_{\text{ex}}$. By assuming a present-day value of $\tau$ (10.0 yr), as suggested by Levine et al. (2011), and $t_{\text{ex}}$ (1.8 yr), Baumgartner et al. (2012) derived from ice cores a rIPD = 3.7 ± 0.7% for LGM. Using the same values for $\tau$ and $t_{\text{ex}}$, we find a rIPD of 5.4% for SDGVM and between −1.0 and 1.7% for ORCHIDEE-V0, V1 and V2. ORCHIDEE-opt gives a rIPD = 3.5%, very close to the value found by Baumgartner et al. (2012). However, the value of rIPD is very sensitive to a small difference in $s_n$ and $s_s$. This is underlined in Fig. 9a by the error bars that give the range of rIPD for SDGVM and ORCHIDEE if 25% of the closest grid cells of Southern Hemisphere to the Equator are accounted for in $s_n$ instead of into $s_s$ (or vice-versa). The rIPD value given by Eq. (4) has to be taken with caution, because only two source regions (corresponding to the two hemispheres) are considered. The two-box split does not therefore account for the basic atmospheric circulation patterns (e.g. Hadley cells), nor does it allow separation of emissions from boreal wetland and northern low latitudes. However, it has the advantage of allowing a simple analytic computation of the rIPD (Baumgartner et al., 2012). Given the uncertainties linked to the latitudinal change of emissions, it is not possible to unambiguously discriminate between SDGVM and ORCHIDEE-opt. However, the comparison between the “observed” rIPD and the rIPD computed using the different ORCHIDEE versions suggests that the modification of the methanogenesis substrate sensitivity to the precipitation in the ORCHIDEE model improves the performance of the model in comparison with the ice-core data (see below).

The intercomparison between two independent models, which account for different processes, could help us to improve our understanding of the potential drivers of the wetland CH₄ emission change during glacial–interglacial transition. While temperature seems to play a small role in the LGM decrease of wetland CH₄ emissions in Weber et
al. (2010), we have shown that modifying the $Q_{10}$ parameterization of the methanogenesis sensitivity is sufficient to reconcile the global LGM–PI change between the two models studied here. The CH$_4$ production sensitivity to temperature is highly uncertain at different spatial scales with large effects on the global distribution of wetland emissions (see e.g. Riley et al., 2011, for the effect on both sites and global scale under current climate). It is however likely that a $Q_{10} = 1.5$ as used in SDGVM is too low. Bringing together the different constraints (i.e. measurements at sites, the distribution of wetland emissions at global scale against top-down estimates, ice-core measurements of LGM–PI change in [CH$_4$]) could help us to reduce the range of plausible $Q_{10}$ values. A further issue is the baseline temperature used in the $Q_{10}$ formulation ($T_{ref}$) and its potential variability in space and time as a way to represent microbial space-scale processes of adaptation (Riley et al., 2011; Z. M. Subin, personal communication, 2011). Additional sensitivity tests with a time variable $T_{ref}$ as in Ringeval et al. (2011) could be performed to evaluate the effect on the simulated change in CH$_4$ emissions between LGM and PI. However, there is still a debate about how influential a microbial community temperature adaptation would be for soil organic matter mineralization. For instance, Rousk et al. (2012) showed that a change in the microbial community (i.e. an adaptation) would be minor as compared to the direct effect of temperature on microbial activity and the indirect effect on the quality of the soil organic matter. Also, discontinuity in the mineralization sensitivity to temperature around 0 °C (Koven et al., 2011) could have a strong effect on the LGM–PI change in emissions and calls for additional tests.

An interesting feature of this work is that the modification of the $Q_{10}$ formulation of the sensitivity temperature (from ORCHIDEE-V0 to ORCHIDEE-V1) does not reconcile the latitudinal distribution of the LGM–PI change between the ORCHIDEE-WET and SDGVM. Other processes are also relatively simply represented in the models (e.g. the constant oxidation related to the plant-transport of CH$_4$ in ORCHIDEE) and could have an effect on the simulated LGM–PI change in wetland CH$_4$ emissions.

In the present study and contrary to Weber et al. (2010), the wetland extent seems to play a small role in explaining the LGM–PI change in emissions. This is particularly true in ORCHIDEE-WET in the boreal regions where the CH$_4$ flux densities collapse and drive the major part of the reduction in emission. This pattern is explained by a large decrease in the ORCHIDEE-simulated methanogenesis substrate. The difference of modelled substrate supply between SDGVM and ORCHIDEE-WET underlies why a modification of the $Q_{10}$ value cannot reconcile the latitudinal distribution of the LGM–PI of the two models. This emphasizes the key role of the substrate supply as suggested by Kaplan (2002). However, the driver of the NPP decrease is still not clear: while Kaplan (2002) explains the low LGM NPP is driven by the reduction in atmospheric CO$_2$ concentration, we highlight the impact of soil freezing on vegetation productivity as a potentially important influence. It is also likely the surface hydrology of models during LGM needs to be improved. For example, consideration of permafrost covered areas, glacial runoff from the Andes and Asian mountains and different hydrological drainage systems may all be important.

This intercomparison discloses some limitations of each model and allows us to suggest different ways of improvement. Regarding ORCHIDEE, we require an improvement to the subgrid computation of the methanogenesis substrate. A limitation is linked to the fact a subgrid computation is performed for the hydrology (through TOPMODEL) but not for the carbon cycle. In this way, the mean carbon over the grid cell is used as proxy of the wetland substrate and this makes the modelled substrate more sensitive to change in precipitation since pre-existing wetland fractions might in reality see less relative change in the soil moisture. This inconsistency between the treatments of hydrology/carbon cycle could be resolved by introducing new wetland plant functional types, which would be restricted to fractional grid cells diagnosed as wetlands using TOPMODEL. Furthermore, we suggest replacing the $Q_{10}$ formulation by an Arrhenius-type equation where the effective activation energy for respiration varies inversely with temperature. In SDGVM, we suggest modifying the contribution of the wetland extent versus CH$_4$ flux densities under PI conditions (see Fig. 4) to more closely satisfy available observations. This could be done by scaling the CH$_4$ flux densities to measurement from sites, and the global wetland extent, e.g. against the value given by Papa et al. (2010) dataset. While the present study does not underline a large effect of the imbalance between the two emission components on the simulated change in emissions in comparison to ORCHIDEE, it may not be the case under other climates. Also, we suggest increasing the value of the $Q_{10}$ for the methanogenesis parameterization. Finally, introducing a simple parameterization of freeze/thaw of soil water could help to more accurately model changes in the methanogenesis substrate availability.

The CH$_4$ emissions simulated by two models at global scale is very similar (GI–HS: 14.2 and 14.8 % relative to LGM emissions respectively for SDGVM and ORCHIDEE-V0). Our best estimate using ORCHIDEE leads to a slightly higher change of 18 % (cf. dash orange curve in Fig. 7, top panel). As described by Hopcroft et al. (2011), the SDGVM-simulated changes in wetland CH$_4$ emissions during the idealized D–O event are too low to explain the measured change in [CH$_4$]. In the present study, the same conclusion is also reached with ORCHIDEE-WET: the likely impact of the simulated emissions on the [CH$_4$] will not differ between the two models. And without any change in the CH$_4$ lifetime, the amplitude of the change in global emissions simulated by the two models (~15 Tg yr$^{-1}$) is much lower than that required (~60 Tg yr$^{-1}$) to match a change of 200 ppb in the [CH$_4$], the upper range of observed D–O events (see...
Fig. 9 of Hopcroft et al., 2011). However the contribution of wetlands to D–O events is still open for debate. For instance, Melton et al. (2012b) estimated that tropical wetlands would increase a maximum of 14 Tg yr$^{-1}$ for a global increase of 64 Tg yr$^{-1}$ in case of the Younger Dryas termination, though this is not usually considered a Dansgaard–Oeschger event.

Our simulations point to two features that could lead to increased D–O forced change in emissions simulated by ORCHIDEE-WET in both the tropics and extra tropics. Firstly, in ORCHIDEE-WET, the LGM boreal emissions are nearly zero. Thus the global D–O changes are almost exclusively explained by the tropics. However, because the $Q_{10}$ of the methanogenesis is larger in ORCHIDEE-WET than in SDGVM, larger boreal emissions at the beginning of the D–O simulations will likely lead to increase the global change during the different phases of the D–O events. Additionally, we have shown that the local (half tropical band scale) anomalies are much larger in ORCHIDEE-WET than in SDGVM (between twice and three times larger). A weak change in the magnitude of the anomaly of a given tropical sub-region could strongly modify the magnitude of the overall total tropical anomaly given the compensating effect described in Sect. 3.2.2. This underlines an increased sensitivity in ORCHIDEE-WET compared with SDGVM and thus a potential larger sensitivity to local changes. Sensitivity FAMOUS simulations with different background conditions (i.e. modifying orbital insolation, global ice volume, greenhouse gases level) have been performed in Hopcroft et al. (2011) and could be used to test these two assumptions relative to the boreal and tropical regions. Relatively little information about the latitudinal change in wetland CH$_4$ emissions during D–O events has been derived from the ice core measurements up to now. Bock et al. (2010) used combined information from the inter-hemispheric gradient and CH$_4$ isotopes to derive source contributions and latitudinal change in emissions of each source between different time periods of the D–O 8 ($\sim$37 kyr BP). According to their modelling approach, the high-latitude wetland emissions strengthened from $\sim$5 to $\sim$32 Tg yr$^{-1}$ from stadial to early-interstadial conditions, whereas tropical wetland emissions strengthened only moderately (from $\sim$84 to $\sim$118 Tg yr$^{-1}$). While the change in boreal emissions between HS and GI is larger in SDGVM than in ORCHIDEE (respectively 9.7 and 5.4 Tg yr$^{-1}$), both models simulate a lower variation of boreal emissions than inferred by Bock et al. (2010). Baumgartner et al. (2012) computed also rIPD for different D–O events and found values of 7.1 ± 0.5, 2.9 ± 2.3 and 6.2 ± 2.4 % for respectively the D–O 2, 3 and 4 using present-day values for $\tau$ and $t_{ex}$. Levine et al. (2012) suggested that the lifetime stayed relatively constant during D–O events, because the effects of both warming and changes in volatile organic compound emission were found to produce effects of approximately equal but opposite sign in their atmospheric chemistry simulations. We compute also rIPD during our idealized GI for SDGVM and the different ORCHIDEE versions (Fig. 9b). Both SDGVM and our best ORCHIDEE estimation are close to the value found by Baumgartner et al. (2012) for D–O 2 and 4. This could suggest that, while the wetland CH$_4$ emission sensitivity to the D–O climate seems to be under-estimated in the models, this under-estimation is homogeneously shared between the Northern and Southern Hemispheres. However, it should also be noted that a 2-box model of the global CH$_4$ sources and atmospheric mixing may not discriminate adequately between the tropical and boreal source regions. This limitation will need to be addressed in future work.

While potential increases in ORCHIDEE-WET simulated wetland CH$_4$ emissions seem to be possible, the results are very similar for the two models over the idealized D–O event. Moreover, contrary to what has been found for the LGM, the ORCHIDEE-simulated change in emissions during D–O shows a relatively low sensitivity to the different parameterizations. This hints at either missing processes related to wetlands, a change in other sources/in the OH sink or alternative D–O mechanism of D–O climate change. The two latter have been discussed in Hopcroft et al. (2011), and we focus here on the first point. The present study as Singarayer et al. (2011) and Hopcroft et al. (2011) underlines the key role of the tropics in controlling the variability in wetland CH$_4$ emissions over paleo-timescales. However, many processes important for tropical wetlands are not accounted for in the current wetland CH$_4$ emission models, which have been developed primarily for the conditions encountered in the most extensively investigated mid-to-northern latitudes. In particular, explicit representation of floodplain hydrology processes in connection with river routing will be required in addition to wetlands saturated from below as represented by ORCHIDEE-WET and SDGVM. These processes are particularly relevant in regions such as the Amazon Basin (Miguez-Macho and Fan, 2012). As suggested by Bock et al. (2010) and discussed above, a change in boreal wetland CH$_4$ emissions appears to be required during some D–O events. In these regions, slow processes such as the exposure of land surface as the ice sheet retreated are clearly not capable of producing such fast variations (Wolff and Spahni, 2007). CH$_4$ emissions associated with permafrost destabilization need to be incorporated into paleo-modelling studies such as the one performed here.
Appendix A

Fig. A1. LGM climate fields simulated by FAMOUS: surface temperature (°C, top panel) and precipitation (mm day⁻¹, bottom panel).

Fig. A2. Applied freshwater forcing to FAMOUS from LGM conditions to perturb the AMOC and mimic D–O events (top panel). Corresponding evolution of the AMOC (middle panel) and Greenland temperature (bottom panel) simulated by FAMOUS.
Fig. A3. Effect of the change in resolution on the Prigent et al. (2007) dataset (inundated areas) (left panels) and on the saturated wetland extents simulated by ORCHIDEE–TOPMODEL (right panels) (expressed in mean annual fraction). The number given in the left bottom corner of each panel corresponds to the global coverage of the wetland extent in %.
Fig. A4. For ORCHIDEE: (a)–(b): prescribed maximum potential PFT coverage: distribution of the main supra-PFT classes for PI (a) and LGM (b) used as input of ORCHIDEE. (c)–(d): Simulated annual maximum PFT coverage: distribution of the main supra-PFT classes for PI (c) and LGM (d). The annual maximum PFT coverage is dependent on NPP. (e)–(f): LGM–PI change in NPP for boreal trees (e) and for C3-grass (d) (in percent of its PI value). (g)–(h): LGM–PI change in JJA humidity stress for C3 grass vegetation for ORCHIDEE-V0 (e) and ORCHIDEE-V2 (f) (in percent of its PI value).
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