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Modelling the continental effect of oxygen isotopes over Eurasia

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ABSTRACT

δ18O in CO2 attracts attention for its ability to decipher the gross fluxes of terrestrial CO2 exchange. This capacity can only be exploited if one knows the impact of the biology and the water isotope cycle on the isotopic composition of atmospheric CO2. To study the processes, we built an integrated global model of δ18O in atmospheric CO2 that calculates 18O in the water cycle pools, the CO2 and the inherent δ18O-CO2 fluxes and transports these in the atmosphere. Within the framework of the European Project EUROSIBERIAN CARBONFLUX, we investigated the 18O processes on the continent of Eurasia north of 40°N. We show that there is a large impoverishment in the water isotopic composition of rain in the continental interior but this is significantly reduced during the growing season. We validated the model directly by comparing it to Net Ecosystem Exchange (NEE) measurements from eddy-flux towers and indirectly by comparing it to atmospheric measurements of CO2 and δ18O-CO2 at 3000 m a.s.l. The model reproduces well the seasonal cycle of CO2 and the phase for δ18O-CO2 but it underestimates the amplitude of that species. Our model further predicts a large diurnal rectifier effect that is different in CO2 than in δ18O-CO2 in eastern Siberia due to very low or even negative leaf discrimination. We analyse the longitudinal gradient in leaf discrimination which reflects primarily the gradient in leaf water isotopic composition. The leaf water δ18O-H2O gradient is itself determined 25% by the meridional gradient in source water and 75% by the change of relative humidity with longitude.

1. Introduction

Our current knowledge does not yet allow to balance the global budget of carbon dioxide within better than ±30% (Prentice et al., 2001). The present consensus is that there is a terrestrial carbon sink in the northern hemisphere. However, the contribution of different geographic locations as well as biomes is not yet clear. For example, it is a matter of debate how this sink is distributed between North America and Eurasia, and more so whether the North American continent and Eurasia are sources or sinks for CO2. Using inversion techniques with current models does not provide an unequivocal distribution of CO2 sources and sinks (e.g. Bousquet et al., 2000; Rayner et al., 1999; Fan et al., 1998; Ciais et al., 1995). Given the existing distribution of CO2 stations, this problem is mathematically underconstrained and therefore needs further and independent constraints.

Though we aim to quantify the biospheric CO2 fluxes better. In the last few years, δ18O in atmospheric CO2 attracted attention to tackle this problem. Observations of the oxygen isotopic composition of CO2 have the capability to distinguish between gross carbon fluxes of the terrestrial biosphere, namely assimilation and respiration. Because of the isotopic exchange of CO2 with water (Mills and Urey, 1940), CO2 exchanged by the biospheric fluxes carries the isotopic
signature of different water pools. These pools are the leaf water and the soil moisture, respectively. $\delta^{18}O$ of atmospheric CO$_2$ is therefore primarily a convolution of CO$_2$ gross fluxes and water isotope signatures. In constraining the isotopic signature of the water pools interacting with carbon dioxide, we can estimate separately the gross CO$_2$ biosphere fluxes.

The global $^{18}O$ signature of H$_2$O in the unsaturated soil layer is about $-10\%$ on average (vs. VSMOW, see below). However, the geographical distribution of the water isotopes is mainly controlled by the rainfall of air masses transported to the interior of the continents. This mechanism produces extremely strong continental gradients of the isotopic composition of precipitation and of the corresponding soil water, and makes a detailed modelling of the abovementioned rain-out mechanism necessary. Subsequently when water evaporates, lighter molecules are preferentially evaporated leaving a heavier water pool. For example, water at the evaporating site of leaves is enriched by about 10–20% (vs. soil water). Net assimilation [which is the Gross Primary Productivity (GPP) minus leaf respiration and equals about 0.85 GPP] and net ecosystem respiration fluxes are around 100 GtC yr$^{-1}$ (Schimel et al., 1996) whereas net exchange between biosphere and atmosphere (NBP) is only 0–2 GtC yr$^{-1}$ globally. This means that the resulting $\delta^{18}O$-CO$_2$ signal is the result of (mainly) two huge opposing fluxes that we name hereafter “isofluxes”, even more pronounced than for CO$_2$ (Ciais et al., 1997a,b; Farquhar et al., 1993). From the convolution, it originates that a small change in the CO$_2$ gross fluxes leads to a big change in the isofluxes. The biospheric gross fluxes should therefore be deducible (Peylin, 1999). Eurasia is the largest land mass in the northern hemisphere. Its role in the global carbon cycle is important but is not known precisely. In addition to CO$_2$ fluxes, there is a special depletion in rainfall, and subsequent water pools in the biosphere in the interior of Eurasia (Dansgaard, 1964) which could lead to peculiar effects in $\delta^{18}O$. Namely, one might expect leaf water $\delta^{18}O$-H$_2$O to be more negative than the atmosphere over most of the continental land mass, yielding photosynthesis to diminish $\delta^{18}O$ in atmospheric CO$_2$ rather than increasing it as usually expected. We have built a consistent global model of $\delta^{18}O$ in atmospheric CO$_2$ which calculates at each timestep the water isotopic composition of the different water pools (in leaves, in soils, and in the atmosphere), the CO$_2$ and CO$_{18}O$ fluxes from and into the atmosphere, and finally the atmospheric CO$_2$ concentration together with its $\delta^{18}O$-CO$_2$ value, including transport. Hence, we built a tool to investigate the CO$_2$ cycle together with the $\delta^{18}O$ cycles in CO$_2$ and water because the exchange of CO$_2$ and water is fully coupled in the model. Here we investigate the distribution of CO$_2$ with $\delta^{18}O$ over the interior of Eurasia. We compare our model results with literature data and measurements made within the European Project EUROSIBERIAN CARBONFLUX.

2. Experimental and model set-up

2.1. The model

The model combines the Atmospheric General Circulation Model (AGCM) ECHAM, including transport of inert tracers and the water isotope cycle, the biosphere model BETHY, and a newly built $^{18}O$-CO$_2$ flux module (Fig. 1). ECHAM is the general circulation model of the Max-Planck Institut für Meteorologie (MPI-MET) in Hamburg, a state-of-the-art global climate model used in several studies (e.g. Arpe et al., 1994; Roeckner et al., 1992) [for a full model description see Modellbetreuungsgruppe (1994)]. In this study, we use the spectral model ECHAM in the T21 resolution corresponding to a physical grid of $5.6^\circ \times 5.6^\circ$ (40 min time step). The model has 19 vertical layers from surface pressure up to a pressure level of 30 hPa. It includes an inert tracer transport scheme which transports the tracer identical to the water vapour with the semi-Lagrangian advection scheme according to Rasch and Williamson (1990).

The water isotope module (WFRAC) was implemented in cycle 3 of ECHAM by Hoffmann et al. (1998) and later built into cycle 4 by Werner (2000). Herein, the water isotopes are treated exactly in parallel to the model’s moisture. However, at each phase transition a temperature-dependent fractionation (Majoube, 1971) is applied to the water isotopes, continuously depleting the vapour phase compared to the liquid or solid phase.

We interfaced the process-based model of terrestrial vegetation activity, BETHY, to the AGCM ECHAM. It calculates at each time step of the AGCM, CO$_2$ fluxes of the terrestrial biosphere together with further diagnostic variables such as stomatal conductance and vegetation temperature. The BETHY online model uses a conservative vegetation distribution map (Wilson and Henderson-Sellers, 1985) adjusted to
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Fig. 1. Information flow of the coupled model. ECHAM is the atmospheric general circulation model (Modellbetreuungsgruppe, 1994); BETHY is the biosphere model (Knorr and Heimann, 2001a,b); other CO₂ sources are monthly input fields, for example ocean fluxes; WFRAC is the water isotope module (Werner, 2000); OFRAC is the newly build CO₂ isotopomer flux module; and δ¹⁸O etc. stands for the transported CO₂ and δ¹⁸O in the atmosphere.

include C4 vegetation (Knorr and Heimann, 2001a). In order to obtain realistic evapotranspiration rates, vegetation temperatures and, therefore, realistic stomatal conductances from the ECHAM energy and water budgets, we recompute the latent and sensible heat fluxes from the canopy. Since plants constantly lose water through their stomatal pores while photosynthesising, water availability is related closely to carbon uptake. A detailed description of the original model can be found in Knorr and Heimann (2001a,b). To interface BETHY to ECHAM, we mainly removed the BETHY internal weather generator and soil model. These variables are taken directly from ECHAM. Deficiencies due to interfacing and not coupling BETHY to ECHAM and other points will be discussed in a subsequent paper.

The ¹⁸O-CO₂ flux module (OFRAC) follows the overall picture of Farquhar et al. (1993) and Ciais et al. (1997). The main difference between the approaches used in Farquhar et al. (1993), Ciais et al. (1997a,b) and Peylin et al. (1999) and our model is the more credible estimation of the ¹⁸O-CO₂ fluxes. We use a time step more adapted to physiological processes and compute the ¹⁸O-CO₂ fluxes interactively in dependence on the synchronous CO₁⁸O concentration in the atmosphere. Each time step, i.e. each 40 min, BETHY computes assimilation and respiration CO₂ fluxes. The ¹⁸O-CO₂ flux module OFRAC computes for every CO₂ flux a corresponding CO₁⁸O flux, namely to assimilation, respiration, ocean, fossil fuel and biomass burning. The corresponding CO₁⁸O fluxes are the product of the CO₂ fluxes with the isotope ratio of the CO₂ dissolved in water and the fractionation occurring at the transition from one compartment to the other (see e.g. Ciais et al., 1997). CO₁⁸O is then transported together with CO₂ by the AGCM ECHAM as inert tracer. It thus permits one to calculate δ¹⁸O in atmospheric CO₂ each time step, and δ¹⁸O-CO₂ can feedback semi-implicitly on the CO₁⁸O fluxes.
2.2. The experimental data

In the EUROSIBERIAN CARBONFLUX project are four different aircraft measurement sites for CO₂ concentrations together with δ¹⁸O-CO₂ outside the atmospheric boundary layer (ABL): Orléans, France (47°55′N, 1°54′E), Tver, Russia (56°28′N, 32°56′E), Syktyvkar, Russia (61°40′N, 50°45′E) and Zotino Russia (60°45′N, 89°24′E) (Levin et al., 2002).

In Tver and Zotino there are also eddy-covariance flux towers installed measuring Net Ecosystem Exchange (NEE) (Milyukova et al., 2002; Tchebakova et al., 2002). We include three other sites for NEE from the EUROFLUX network further west to have a meridional coverage: Sarrebourg, France (48°40′N, 7°05′E) (Granier, 2002), Flakaliden, Sweden (64°14′N, 19°46′E) (Lindroth, 2002), and Hyytiälä, Finland (64°51′N, 24°17′E) (Vesala, 2002; Suni, T., personal communication).

We also compare the modelled isotopic signature in precipitation to stations of the Global Network for Isotopes in Precipitation (GNIP) of the International Atomic Energy Agency (IAEA), Vienna, Austria (IAEA/WMO, 2001). The GNIP database incorporates more than 10⁴ measurements of the three isotopes deuterium, tritium, and ¹⁸O-H₂O in precipitation and gives monthly mean values for the measurement period. The GNIP stations are not identical to the above locations of CO₂ measurements, so we selected GNIP stations nearby. These are namely Thonon-les-Bains, France (46°13′N, 6°16′E), St. Petersburg, Russia (59°34′N, 30°10′E), Kirov, Russia (58°23′N, 49°22′E), and Enisejsk, Russia (58°16′N, 92°05′E).

All δ¹⁸O-H₂O values of water reported here are relative to the standard Vienna SMOW (VSMOW) (Baertschi, 1976) and δ¹⁸O-CO₂ relative to Vienna Pee Dee belemnite (VPDB-CO₂) (Allison et al., 1995).

3. Results and discussion

Figure 2 shows the investigation area together with the locations of the measurement stations. However, we used only the area between 40°N and 70°N for

Fig. 2. Two different representations of the investigation area. In the upper frame are indicated the measurement stations of water isotopes (squares), CO₂ fluxes (cross) and atmospheric CO₂ with isotopes (circles) (ORL, Orléans; THO, Thonon-les-Bains; SAR, Sarrebourg; FLA, Flakaliden; HYY, Hyytiälä; S.P, St. Petersburg; TVE, Tver; SYT, Syktyvkar; KIR, Kirov; ZOT, Zotino; ENI, Enisejsk). In the lower frame, the boxes are the grids of the model and the colours with numbers stand for each vegetation class used (4–15 of 23 overall in the model).
the calculation of the longitudinal gradients. But, one should keep in mind that the model is run globally, so it fully accounts for lateral input/output of H₂O, CO₂ and isotopes to/from the Eurasian domain in this study. In the upper plate is also indicated the border of the land masses in the model. Combining the upper with the lower panel gives a clear picture of the used model resolution. In the bottom plate, we show the vegetation distribution which is used in BETHY. There is mostly tundra vegetation in north Siberia and a mixture of evergreen and deciduous conifers over temperate and boreal Eurasia as well as in east Siberia. The southern part is dominated by grasses (C3 and C4) and crops. It is apparent in the vegetation distribution that, apart from the longitudinal gradient in climate due to continentality, a gradient in latitude exists which also plays a role for δ¹⁸O-CO₂.

The δ¹⁸O signal in CO₂ is mainly the convolution of the water isotopic composition with the CO₂ fluxes. Hence, both parts of the convolution have to be validated. We compare first the water isotopes with measurements from the GNIP database and next CO₂ net fluxes with measurements of eddy-flux towers. We will then compare the seasonal cycle of CO₂ and δ¹⁸O-CO₂ in the free troposphere, namely at ca. 3000 m a.s.l.

3.1. Water isotopes

On a global scale, a detailed comparison of the water isotope module of ECHAM with global observations of the IAEA observations gave an excellent agreement of the spatial and temporal patterns of the simulated and observed isotope fields (Hoffmann et al., 1998). To illustrate this, we show here some typical results of the monthly δ¹⁸O-H₂O in precipitation over Eurasia (Fig. 3). Note that all IAEA stations in Russia are actually closed now, so the presented data include no recent water isotopic composition. Nevertheless, there is no obvious trend in time in the water isotopic composition, so we can use the older data for isotopes in precipitation as an isotope climatology. In Fig. 4 one can clearly see the continental impoverishment in rainfall isotope data at the four selected IAEA stations distributed along the eastward transport trajectory of water vapour over Eurasia. The continental gradient and its origin (the rainout mechanism) is already described in the first review of the IAEA network (Dansgaard, 1964). One can also see that the isotopic composition of the soil water is close to rain water in our model. The difference originates mainly from different infiltration and runoff of winter and summer precipitation (Hoffmann et al., 1998). Figure 3 illustrates the increasing continentality, which is also linked to an increasing seasonal amplitude, not only in temperature but also in water isotopes (up to 120°E approximately). A larger seasonal amplitude of δ¹⁸O-H₂O in precipitation is consistently observed (and simulated by ECHAM) in higher latitudes compared to lower latitudes and in the interior of the continents compared to regions under a marine influence. The increasing seasonality of the water isotopes, moving into the continent, mainly lowers the winter values (summer, about −8 to −10‰ both in St. Petersburg and in Enisejsk; winter, −16‰ in St. Petersburg and −24‰ in Enisejsk). As a consequence, the spatial gradient of the water isotopes, strongly expressed in the annual mean and even more in winter, is comparatively weaker during plants’ growing season in late spring and summer.

3.2. Net ecosystem exchange

Figure 5 shows a comparison of the measured Net Ecosystem Exchange (NEE) on eddy-flux towers with the modelled fluxes of BETHY. Such a comparison should be treated with caution because NEE measurements are representative of a very small area (<1 km²) compared to our model grid. It is, however, a useful semi-quantitative information of the BETHY model performance. Included is the modelled NEE

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Fig. 4. Longitudinal gradient of water isotopes and temperature over Eurasia of yearly and summer means. The modelled meridional summer and yearly mean rain and soil water are shown in comparison to the four IAEA stations of Fig. 3. The summer composition of soil water is not shown because it is very similar to the yearly mean. The model output is processed like the station data, but the monthly means of the model are already rainfall weighted means.

Fig. 5. Net ecosystem exchange (NEE) modelled and measured with eddy-flux towers in Sarrebourg, Flakaliden, Hyytiälä, Tver, and Zotino.

(at monthly time steps) of Peylin et al. (1999), which comes from SIB2 (Sellers et al., 1996a,b), for comparison. BETHY’s net fluxes are too small in amplitude but they are realistic for the phase of NEE at almost all flux tower sites. The same is true for the fluxes of SIB2 but with a slightly larger, though more realistic amplitude. BETHY has a larger net flux amplitude only at the most western station, Sarrebourg. Flakaliden is the northernmost station which shows a very sharp decrease of NEE in May probably at the time when the temperature in Flakaliden passes the threshold temperature for assimilation for the dominant species, *Picea abies*, there. This behaviour is captured by BETHY but is not as pronounced as in the measurements. There is no such abrupt behaviour in Hyytiälä whereas the model still shows a sudden beginning of assimilation. (Hyytiälä falls in the same grid box as Flakaliden in the model.) In Tver, both models are very comparable but BETHY reproduces better the large positive flux to the atmosphere at the end of the growing season. Both models predict too much respiration in Zotino at the end of the year but SIB2 simulates the minimum value in June well. One should recall that NEE is actually the sum of two opposing fluxes, of assimilation and ecosystem respiration, where the maximum of assimilation ranges from 3 μmol (CO₂) m⁻² s⁻¹ in Flakaliden to 7.5 μmol (CO₂) m⁻² s⁻¹ in Tver. The amplitude and the phase of NEE are very sensitive to the phasing of the two fluxes, especially in regions with a contrasted seasonal climate. In contrast, the western part of Eurasia, with a mild climate and hence not as
pronounced seasonal variations as in the interior, is much more insensitive to the respective phasing of assimilation and respiration.

3.3. Seasonal cycle of atmospheric measurements

CO₂ and the CO¹⁸O fluxes emitted into the atmosphere are transported via advection, convection and turbulent diffusion. There is no known fractionation associated with these processes, so CO₂ and CO¹⁸O are transported as passive tracers. We have not closed the carbon cycle budget globally in our model, so we can not strictly compare the absolute values. One flux omitted in the model is deforestation but there is probably only a small seasonality associated with it. So we can still compare the simulated seasonal cycle, both of CO₂ and δ¹⁸O-CO₂, with the observations. The upper panel of Fig. 6 shows the mean seasonal cycle of CO₂ in 3000 m a.s.l. as measured (circles) (Levin et al., 2002) and modelled by ECHAM/BETHY (solid line) and Peylin et al. (1999) (dashed line). The atmospheric CO₂ concentrations of Peylin et al. are the monthly SIB2 fluxes transported in the TM2 atmospheric transport model (Heimann, 1995). The seasonal cycle of CO₂ is well reproduced by both models. The amplitudes of the data and both models are almost the same. Nevertheless at Syktyvkar and Zotino ECHAM/BETHY seems to produce slightly more realistic amplitudes than Peylin et al. and thus seems to capture the seasonal cycle of CO₂ better even if its not significant. The lower panel of Fig. 6 shows the mean seasonal cycle in δ¹⁸O-CO₂ with measurements discussed in Levin et al. (2002). The CO¹⁸O fluxes of Peylin et al. are the synthesis of the SIB2-CO₂ fluxes with the GISS water isotopes (Jouzel et al., 1987) on a monthly mean basis and the transport of TM2. Both models follow the phase of the cycle, but Peylin et al. have a higher amplitude. One can clearly see that the amplitude in the CO₂ seasonal cycle remains roughly constant when going into the interior of the continent, from Tver to Zotino. Only Orléans is much lower, reflecting its maritime influence. In contrast, the amplitude of δ¹⁸O-CO₂ increases from Orléans to Syktyvkar and then decreases in Zotino again. This behaviour is not well understood (Levin et al., 2002). There are several possible explanations: 1. There are different transport patterns west and east of the Urals, i.e. west of the Urals is more influenced by the Azores high and the Icelandic low, and east of the Urals is mostly influenced by the Siberian high (winter) and low (summer) (Aizen et al., 2001). Zotino is therefore more influenced by northern latitudes (Levin et al., 2002); 2. The free troposphere is much more influenced by ABL air west of the Ural than east of them, but measurements (Lloyd et al., 2002; Ramonet et al., 2002) and simulations with regional scale models (Chevillard et al., 2002; Kjellström et al., 2002) indicate the contrary; 3. There is a positive gradient of CO₂ gross (not necessarily net fluxes, cf. CO₂ seasonal cycle) and hence δ¹⁸O-CO₂ fluxes from the Atlantic to the Urals and a negative gradient from the Urals to the Pacific; 4. There is no gradient of the CO₂ gross fluxes but there is one in discrimination. Unfortunately, our model does not fully resolve the Ural mountains in its influence on weather regimes because it has a smoothed orography and a rather coarse resolution.

Even if we can not compare the absolute values with the measurements, the longitudinal gradient between Orléans and Zotino should be comparable. In Table 1 are listed the differences in ECHAM/BETHY and in the data of the mean and amplitudes of CO₂ and δ¹⁸O-CO₂ between Tver, Syktyvkar, Zotino and Orléans. Table 1 indicates that there is practically no difference in the amplitude of the seasonal cycle of CO₂ between the three stations in Russia, but the offset between the means differs considerably Levin et al. (2002) argued that this offset is not significant because the differences do not occur over the whole measurement period and even disappear by the end of 2000. From δ¹³C measurements in CO₂, the authors suggested that there is
though it hinders an accurate inversion of gradient in CO$_2$ concentrations between the surface. NEE is negative (sink), resulting in a small negative growing season, the vertical transport is vigorous and effect (Stephens et al., 1999). During daytime in the effect. We further investigate here the diurnal rectification between surface δ$^{18}$O-CO$_2$ where it reaches the size of Orléans to Syktyvkar and decreases approaching Zotino, whereas the amplitude increases from Orléans over Tver to Syktyvkar and decreases approaching Zotino, where it reaches the size of Orléans again. Our model does not capture the behaviour of the data but produces δ$^{18}$O-CO$_2$ fields that are rather uniform in size and amplitude over the whole of Russia. δ$^{18}$O-CO$_2$ does not seem to change significantly over the continent at 3000 m in the model.

3.4. Diurnal rectification gradient

An interesting phenomenon in the carbon cycle, although it hinders an accurate inversion of fluxes, is the covariation between surface fluxes and vertical or horizontal transport, in a generic manner called the rectifier effect. We further investigate here the diurnal rectifier effect (Stephens et al., 1999). During daytime in the growing season, the vertical transport is vigorous and NEE is negative (sink), resulting in a small negative gradient in CO$_2$ concentrations between the surface and the ABL. During night-time, vertical transport is almost suppressed and NEE is positive (source), inducing a large CO$_2$ accumulation near the ground in the shallow nocturnal boundary layer. If one measures only near the ground, the mean concentration in time is shifted to higher values while the mean mid-ABL concentration is shifted to lower CO$_2$ values. Inverse modelling efforts, taking monthly or yearly mean concentrations, have to take into account this spatial gradient induced by the “diurnal rectifier effect” (DRE). Current inverse models mostly take stations on the ocean, where the DRE is not very large if the continental influence can be neglected (Denning et al., 1996). However, stations nearby or inside Eurasia, like the Shemya Islands (Aleutians) or Ulaan Uum (Mongolia), could be strongly influenced by the DRE, the magnitude and extent of which are yet poorly known. There are even inverse models now which invert CO$_2$ and δ$^{18}$O in atmospheric CO$_2$ where the DRE is ignored (Peypin, 1999). The DRE effect is very strong in summer but of minor influence in winter. In addition to the DRE, there are also different transport patterns between summer and winter, vertically and horizontally which also covary with the seasonal pattern of NEE. This effect can lead to another generation of mean spatial concentration gradients (e.g. Denning et al., 1995) and is referred to here as “seasonal rectifier effect” (SRE). Covariations with vertical transports normally induce an enhancement of the measured concentrations near the ground (Stephens et al., 1999), whereas horizontal transport can lead to an abasement (Ciais et al., 2000; Taylor, 1998). The sign of this SRE can be negative or positive depending on the transport patterns and or the regions considered. Our model shows both rectifier effects which are not distinguishable in a normal simulation. To simplify matters, we will also call the overall effect the diurnal rectifier effect because the DRE makes approximately 80% of the total rectifier effect over land (Denning et al., 1996). Figure 7 illustrates the DRE in ECHAM/BETHY. We show the simulated fields for February, August and the yearly mean CO$_2$ and δ$^{18}$O-CO$_2$ in the first (ca. 30 m above ground) and third levels (ca. 800 m above ground) of the model. The third level lies in the middle of the ABL in summertime but in night- and winter-time

### Table 1. Comparison of absolute modelled and measured concentrations relative to Orléans

| Model | TVE | SYT | ZOT |
|-------|-----|-----|-----|
| Mean CO$_2$ (ppm) | 1.1 | 0.6 | 0.3 |
| Amplitude CO$_2$ (ppm) | 2.8 | 4.9 | 4.6 |
| Mean δ$^{18}$O-CO$_2$ (%) | -0.2 | -0.2 | -0.3 |
| Amplitude δ$^{18}$O-CO$_2$ (%) | 0.2 | 0.1 | 0.2 |

Model Data

| Model | TVE | SYT | ZOT |
|-------|-----|-----|-----|
| Mean CO$_2$ (ppm) | 1.1 ± 2.4 | 3.1 ± 1.9 | 1.2 ± 2.1 |
| Amplitude CO$_2$ (ppm) | 5.2 ± 2.4 | 5.1 ± 1.9 | 4.7 ± 2.1 |
| Mean δ$^{18}$O-CO$_2$ (%) | -0.8 ± 0.3 | -0.8 ± 0.4 | -0.5 ± 0.4 |
| Amplitude δ$^{18}$O-CO$_2$ (%) | 0.4 ± 0.3 | 0.7 ± 0.4 | 0.0 ± 0.4 |

Note: ORL, Orléans; TVE, Tver; SYT, Syktyvkar; ZOT, Zotino. Mean is the difference of total averages and Amplitude is the difference in amplitude of the mean seasonal cycles. The values in parentheses are the ratio of the amplitude of the mean seasonal cycle to the amplitude at Orléans.
Fig. 7. Rectification gradients of CO$_2$ and δ$^{18}$O-CO$_2$ at ground (first column) and third (second column) level of the model, approximately 30 and 800 m respectively, above ground. The third column is the difference between the first and second column. In the third column, the colour bar differs from those used in columns one and two. We suppressed longitude and latitude labelling to make the plot intelligible. Longitude ranges from 15 to 170°W (−15 to 190°) and latitude ranges from 35 to 75°N. We subtracted the global mean concentration from each grid point as a reference. This gives negative values everywhere in δ$^{18}$O-CO$_2$ over Eurasia because of the simulated strong north–south gradient.

outside it. We also excluded the concentrations of fossil fuel and biomass burning of CO$_2$ and δ$^{18}$O-CO$_2$ in Fig. 7 in order to examine only the rectification gradients induced by covariations of biospheric fluxes and transport. The model shows an accumulation of CO$_2$ at the ground level with respect to 800 m over the continent on a monthly average basis. This accumulation ranges from 1 ppm over eastern Siberia in winter up to 30 ppm over some regions over western Russia in summer and to 40 ppm over south-east Asia. The wintertime CO$_2$ accumulation is not very pronounced over Eurasia but it reaches a maximum of 10 ppm in south Siberia. Otherwise, it is quite uniform between 2 and 6 ppm in the rest of the investigation area. In summer, there is a vast area between 40 and 60°N and 30 and 90°E where the monthly mean concentrations at the ground are more than 6–30 ppm higher compared to 800 m. West of it, the DRE again seems to be quite uniform around 3 ppm, and east of it the DRE falls even beyond 0. In summer, the 800 m level lies most of the time in the CBL during the day and indicates the net effect of daytime NEE with reduced CO$_2$ concentrations. However, δ$^{18}$O in atmospheric CO$_2$ shows a similar pattern in winter as CO$_2$ does. It shows a small accumulation of negative δ$^{18}$O-CO$_2$ (of respired CO$_2$) in Europe and western Siberia near the ground, with values around −0.3‰ compared to 800 m. In contrast to western Siberia, we see almost no difference.
between ground- and third-level in $\delta^{18}$O-CO$_2$ in winter in eastern Siberia. During summer there is a drawdown in $\delta^{18}$O-CO$_2$ between 800 m and the surface of more than $-0.4\%$ over the whole domain, reaching values of $-1.5\%$ around St. Petersburg and around Jakutsk in eastern Siberia. This pattern mirrors the DRE of CO$_2$ in Europe and west Siberia, but it is different over eastern Siberia. This surprising signal over eastern Siberia comes from negative $^{18}$O leaf discrimination east of approximately 90$^\circ$ E (see below and Fig. 9c). Already at Zotino the DRE signal for $\delta^{18}$O-CO$_2$ differs from that of CO$_2$. We have plotted in Fig. 8 vertical profiles up to an altitude of approximately 4500 m in February and August. We subtracted at each station the value of the tropopause model level (ca. 12 km) from the model values. Contrasting Orléans and Zotino in August, one can see that a comparable vertical profile in CO$_2$ (ca. 5 ppm), reflecting DRE, leads to a very different profile in $\delta^{18}$O-CO$_2$. This comes from the fact that leaf discrimination at Zotino is around zero during the growing season in the model, so that the signal of $\delta^{18}$O-CO$_2$ is determined mostly by the soil isoflux then (see below and Fig. 9c). One can also notice that the magnitude of the accumulation as well as the difference between ground and 800 m changes between the stations and thus reflects the specific isoflux sign at each measurement station. Current inverse modelling efforts very rarely include the diurnal cycle, so they include only the seasonal covariations between fluxes.
and transport. Thus, for example, the prediction that assimilation in Siberia must be increased in current biosphere models (Peylin, 1999) could be an artefact. On the other hand, choosing a site where leaf discrimination is zero in summer and measuring vertical transport alone, supposing one knows or respectively measures soil variation in vertical transport alone, supposing one determines the diffusion fractionation (see below). This suggests that $\delta^{18}$O-CO$_2$ could be used under these conditions to validate atmospheric transport models to reduce the uncertainties implied by the existence of the DRE.

3.5. Relating the simulated and observed longitudinal gradients to controlling climate variables

Writing the global balance equation for $\delta^{18}$O in atmospheric CO$_2$ (regarding only the biospheric activity) leads to:

$$M \frac{d\delta}{dt} = A\Delta_A + F_R \Delta_R$$  \hspace{1cm} (1)

with

$$\Delta_A = -\varepsilon_1 + \frac{c_s}{c_a - c_s}(\delta_l - \delta_a),$$  \hspace{1cm} (2)

$$\Delta_R = \delta_a - \delta_a + \varepsilon_s.$$  \hspace{1cm} (3)

$M$ is the number of moles of CO$_2$ in the atmosphere, $\delta_a$ is $\delta^{18}$O of atmospheric CO$_2$, $A$ the net assimilation (assimilation minus leaf respiration), $\Delta_A$ the discrimination of assimilation (called leaf discrimination, too), $F_R$ the total biospheric respiration minus leaf respiration (heterotrophic plus autotrophic minus leaf respiration), called net ecosystem respiration, $\Delta_R$ the difference between soil respired $\delta^{18}$O-CO$_2$ and atmospheric $\delta^{18}$O-CO$_2$ (we call it soil “discrimination” even if it is not a real discrimination), $\varepsilon_1$ the diffusion fractionation of CO$_2$ entering and leaving the stomata, $c_s$ the CO$_2$ concentration at the surface of the chloroplasts, $c_a$ the CO$_2$ concentration of the atmosphere (precisely the canopy), $\delta_l$ the $\delta^{18}$O of CO$_2$ dissolved and isotopically equilibrated with leaf water, $\delta_a$ the $\delta^{18}$O of atmospheric CO$_2$, $\delta_s$ the $\delta^{18}$O of CO$_2$ dissolved and equilibrated with soil water, and $\varepsilon_s$ the fractionation during diffusion of CO$_2$ out of the soil. The product of $A$ and $\Delta_A$ is called the isoflux of assimilation or leaf isoflux and $F_R$ times $\Delta_R$ is called the isoflux of respiration or soil isoflux. Peylin et al. (1999) calculated separately the assimilation and respiration fluxes and $\Delta$ of assimilation and respiration, both on a monthly mean basis. They transported the product of flux times $\Delta$ in TM2.

Having demonstrated that our model reproduces well the water isotopic composition, the CO$_2$ fluxes as well as the atmospheric CO$_2$ concentrations, one can still argue that the amplitude of the seasonal cycle of $\delta^{18}$O-CO$_2$ is not well represented in our model (Fig. 6). Conversely, the model matches well the $\delta^{18}$O-CO$_2$ values at atmospheric background stations (results not shown here), i.e. stations from e.g. GLOBALVIEW (GLOBALVIEWCO2, 2000). However, one can argue that the $\delta^{18}$O-CO$_2$ values over the continent are too low because of the amplification of small model shortcomings, e.g. too small convection, the missing “weather barrier” of the Ural or too low isofluxes over Siberia, although the correct phasing of $\delta^{18}$O-CO$_2$ in our simulation, together with the rather good fit at atmospheric background stations, indicates that the patterns of assimilation, isofluxes and hence discrimination should be realistic even if the absolute values are not perfectly reproduced. Additionally, none of the existing global models is capable of reproducing $\delta^{18}$O in atmospheric CO$_2$ globally in phase and amplitude up to now.

We show in Fig. 9 the meridional gradient of the CO$_2$ fluxes and isofluxes calculated by the ECHAM/BETHY model. The upper panel (Fig. 9a) shows the annual net assimilation (see above) given in GtC yr$^{-1}$ per longitude band. The black line is the result of our model, again in comparison with Peylin et al. (grey line). Three major points in Fig. 9a are the two maxima in Europe and the difference between BETHY and SIB2 east of 120$^\circ$E. The first maxima (5$^\circ$E) in assimilation represents western Europe up to South Scandinavia, Central Europe, and the Mediterranean Balkan area, from north to south. Here there is a very temperate climate in ECHAM which is favourable for assimilation. The second large maxima in assimilation around 40$^\circ$E comes from the presence of arable crops around the black sea and in western Russia (Fig. 2). The last maxima of assimilation in BETHY at around 130$^\circ$E corresponds to the great agricultural plains in northern China. East of 20$^\circ$E, SIB2 shows practically no gradient in assimilation, and SIB2 shows larger photosynthesis rates than BETHY in the far north-east Siberian tundra. Both flux models show almost identical total net assimilation for the complete Eurasian domain (north of 40$^\circ$N), namely 14.5 GtC yr$^{-1}$, and discarding the peaks in BETHY, both models are roughly similar. The middle panel (Fig. 9b) shows patterns of isofluxes for photosynthesis (straight line) and respiration (dashed line). The lower panel (Fig. 9c) shows $\Delta$,
the “discrimination” of both photosynthesis (straight line) and respiration (dashed line). \( \Delta \) in our model is the division of the isoflux by its \( \Delta \) CO\(_2\) flux and it is therefore a flux-weighted \( \Delta \), where the weighting accounts for temporal variability in fluxes each 40 min. Our \( \Delta \) thus reflect exactly what is “seen” by the atmosphere. In contrast, Peylin et al. calculated first monthly \( \Delta \) and the CO\(_2\) fluxes separately, from which they deduced the isofluxes afterwards. Peylin et al. took only relative humidity and the CO\(_2\) concentration at the surface of the chloroplast assimilation weighted in the calculation of leaf water isotopic composition, whereas all other variables are taken as monthly averages. They predict a small decrease of the leaf isoflux and a decrease in the soil isoflux from west to east inside Eurasia (ca. 5 GtC \( \% \) yr\(^{-1}\)). ECHAM/BETHY shows noticeably different fields, decreasing in assimilation from about 20 GtC \( \% \) yr\(^{-1}\) to small negative values. However, the total isoflux (sum of leaf and soil isoflux) of the biosphere is quite similar for both models over Eurasia. It is remarkable that leaf discrimination (Fig. 9c) in ECHAM/BETHY becomes negative after 90° E (the missing values after 150° E come from the fact that one divides the isoflux by a very small CO\(_2\) flux, which leads to numerical instability and is therefore discarded). Between 50° and 90° E, the difference between ECHAM/BETHY and Peylin et al. comes certainly from the different assimilation weighting procedures. Both models use the so-called Farquhar formulation (Farquhar et al., 1980) for assimilation. If the assimilation between 50° and 90° E is fairly similar in both studies (Fig. 9a), the determining parameters like temperature and humidity should also be quite similar. However, already with temperature, one can see that it will make a big difference in, e.g., the equilibration of CO\(_2\) with water (–0.4‰ per degree increase) if one considers explicitly a diurnal cycle. We try to understand what determines the continentality gradient in Fig. 9c. The leaf discrimination is determined by the isotopic composition of CO\(_2\) equilibrated with leaf water at the evaporating site, \( \delta_i \), the atmospheric \( \delta^{18}O\)-CO\(_2\) value, and the factor \( c_i/(c_a-c_i) \) which is the amplification of leaf fractionation due to back-diffusion [Eq. 2]. The leaf water at the evaporating site is calculated from the formulation of Craig and Gordon:

\[
\delta_i^w = \varepsilon_k^w + \delta^{18}O^{atm} - \varepsilon_k \left( \delta_{sap} - \delta_k^w + \varepsilon_k \right),
\]

\( \varepsilon^w \) is the fractionation of the liquid–vapour phase transition [according to Majoube, (1971)] and \( \delta_{sap} \) the source water isotopic composition, \( \varepsilon_k \) the kinetic fractionation factor (taken constant), \( h \) the relative humidity, and \( \delta_{sap} \) the isotopic composition of water vapour. CO\(_2\) equilibrated with this leaf water is then:

\[
\delta_i = \varepsilon^w + \delta_{eq}.
\]

where \( \delta_{eq} \) is the equilibrium fractionation calculated after Brenninkmeier et al. (1983).

In Fig. 10 are plotted the variables which determine the “discriminations” and hence the isofluxes. We plotted both the assimilation-weighted (black lines) and the plain variables (grey lines). (Assimilation-weighted variables are multiplied with the assimilation flux at each time step, and the yearly sum of these products is then divided by the yearly sum of assimilation. They therefore give more realistic mean values.) One can divide northern Eurasia into two parts: the western part from the Atlantic to the Ural (Europe) and the eastern part from the Ural to the Pacific (Siberia), say from 0 to 60° E and from 60 to 180° E. We observe in Fig. 10 an increase in leaf temperature (Fig. 10a) and thus in \( \delta^{18}O\)-CO\(_2\) (Fig. 10e) equilibrated with leaf water (at the evaporating site) in

![Fig. 10. Variables which determine the \( \delta^{18}O\)-CO\(_2\) discrimination of assimilation. Black lines are assimilation-weighted values, and grey lines are not assimilation-weighted (see text for the difference between assimilation-weighted or not).](image-url)
Europe and a decrease in Siberia. The converse picture is true for relative humidity (Fig. 10c) in the canopy. $\delta^{18}$O-H$_2$O of groundwater (Fig. 10d) decreases by about $-5\%$ over the whole continent, whereas $\delta^{18}$O-H$_2$O vapour minus ground water stays constant despite some fluctuations (Fig. 10d). This leads to an overall decrease in leaf $\delta^{18}$O-H$_2$O across Siberia of around $-20\%$ between the Ural and the Pacific. In this signal, $-5\%$ comes from the decrease in source isotope composition and almost $-15\%$ comes from the increase in relative humidity. The temperature-controlled equilibration of CO$_2$ with water isotopes opposes this big drop. However, the assimilation-weighted leaf temperature (Fig. 10a) decreases by only 10$^\circ$C across Siberia, so it lowers the gradient in $\delta_t$ only by about $-2\%$ (Fig. 10e). Since the model shows no east-west gradient in atmospheric $\delta^{18}$O-CO$_2$ (Fig. 10e), the decrease of $\delta_t$ over Siberia is amplified by $c_t/(c_t - c_s)$ (Fig. 10b) and leads to the strong east-west gradient of leaf discrimination of about $-40\%$ (Fig. 9c).

4. Conclusions

We constructed a model of biospheric CO$_2$ exchange interfaced to an AGCM which also fits with a calculation of the isotopic composition of water. We implemented a new module to calculate, in a fully consistent manner with climate, the cycling of $^{18}$O-H$_2$O in the surface water pools (soil, leaves) and the pertaining $\delta^{18}$O-CO$_2$ fluxes exchanged with the overlaying atmosphere. We studied the space and time distribution of water oxygen isotopes, CO$_2$ biospheric fluxes, atmospheric CO$_2$ and $\delta^{18}$O in atmospheric CO$_2$, $\delta^{18}$O-CO$_2$ fluxes, the isotopic discrimination of assimilation, and the isotopic “discrimination” of respiration with respect to CO$_2$ across northern Eurasia. It is shown that the water isotopic composition is simulated well and shows a realistic meridional negative gradient going from west to east. This gradient is much smaller for summer than for winter, so the relevant gradient in water isotopic composition is only about $-5\%$. The NEE flux that we simulate appears too small in amplitude at the selected flux tower sites, but the comparison can be only semi-quantitative. The seasonal cycle at 3000 m a.s.l. of $\delta^{18}$O in atmospheric CO$_2$ offers an unique integrated measure of the performances of our different modules (NEE, transport, $\delta^{18}$O-CO$_2$, etc., and their interactions). The atmospheric seasonal cycle is modelled very well for CO$_2$, but the amplitude of $\delta^{18}$O-CO$_2$ is underestimated, although the model reproduces the right phase.

We found that the diurnal rectifier effect, implied by covariations of biotic fluxes with vertical transport in the ABL, is very pronounced over Eurasia and different between CO$_2$ and $\delta^{18}$O-CO$_2$. This comes mostly from the fact that leaf discriminations become zero or negative even in the yearly mean east of 90$^\circ$E, due to the continentality.

Together with the isofluxes of $\delta^{18}$O-CO$_2$, we demonstrated that there is a large gradient in leaf discrimination from west to east over Eurasia, and leaf discrimination can even become negative in the most continental area of eastern Siberia. We interpret this signal on one hand from the water isotope gradient but mostly from the gradient in relative humidity that lowers the evaporating leaf water $\delta^{18}$O-H$_2$O and, therefore, the leaf discrimination.

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