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The impact of soil microorganisms on the global budget of δ^{18} O in atmospheric CO₂

Lisa Wingate^{a,b,1,2}, Jérôme Ogée^{b,1,2}, Matthias Cuntz^{c,1,3}, Bernard Genty^{d,e}, Ilja Reiter^{d,e}, Ulli Seibt^f, Dan Yakir^g, Kadmiel Maseyk^{f,g}, Elise G. Pendall^h, Margaret M. Barbourⁱ, Behzad Mortazavi^{j,k}, Régis Burlett^b, Philippe Peylin^f, John Miller^{I,m}, Maurizio Mencuccini^a, Jee H. Shimⁿ, John Huntⁱ, and John Grace^a

^aSchool of GeoSciences, University of Edinburgh, Edinburgh EH9 3JN, United Kingdom; ^bUnité de Recherche 1263 Ecologie Fonctionnelle et Physique de l'Environnement, Institut National de la Recherche Agronomique, 33130 Villenave d'Ornon, France; ^cMax Planck Institute for Biogeochemistry, 07701 Jena, Germany; ^dLaboratoire d'Ecophysiologie Moléculaire des Plantes, Institut de Biologie Environnementale et de Biotechnologie, Service de Biologie Végétale et de Microbiologie Environnementale, Commissariat à l'Energie Atomique, 13108 Saint-Paul-lez-Durance, France; ^eUnité Mixte de Recherche Biologie Végétale et Microbiologie Environnementales, Centre National de la Recherche Scientifique, 13108 Saint-Paul-lez-Durance, France; ^fUnité Mixte de Recherche 7618 Biogéochimie et Ecologie des Milieux Continentaux, Centre National de la Recherche Scientifique/Université Pierre et Marie Curie, 78850 Thivernal-Grignon, France; ⁹Department of Environmental Sciences and Energy Research, Weizmann Institute of Science, Rehovot, 76100, Israel; ^hDepartment of Botany, University of Wyoming, Laramie, WY 82071; ¹Landcare Research, P.O. Box 40, Lincoln 7640, New Zealand; ¹Department of Biological Sciences, University of Alabama, Tuscaloosa, AL 35487; ^kDauphin Island Sea Lab, Dauphin Island, AL 36528; ¹National Oceanic and Atmospheric Administration Earth System Research Laboratory, 325 Broadway R/GMD1, Boulder, CO 80305; ^mCooperative Institute for Research in Environmental Sciences, University of Colorado, Boulder, CO 80309; and ^mDepartment of Forest, Rangeland, and Watershed Stewardship, Colorado State University, Fort Collins, CO 80523

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Improved global estimates of terrestrial photosynthesis and respiration are critical for predicting the rate of change in atmospheric CO₂. The oxygen isotopic composition of atmospheric CO₂ can be used to estimate these fluxes because oxygen isotopic exchange between CO₂ and water creates distinct isotopic flux signatures. The enzyme carbonic anhydrase (CA) is known to accelerate this exchange in leaves, but the possibility of CA activity in soils is commonly neglected. Here, we report widespread accelerated soil CO₂ hydration. Exchange was 10-300 times faster than the uncatalyzed rate, consistent with typical population sizes for CAcontaining soil microorganisms. Including accelerated soil hydration in global model simulations modifies contributions from soil and foliage to the global CO18O budget and eliminates persistent discrepancies existing between model and atmospheric observations. This enhanced soil hydration also increases the differences between the isotopic signatures of photosynthesis and respiration, particularly in the tropics, increasing the precision of CO₂ gross fluxes obtained by using the δ^{18} O of atmospheric CO₂ by 50%.

carbon cycle | water cycle | carbonic anhydrase | oxygen isotopes | terrestrial biosphere

he Earth's climate system is intimately connected to the movement of water and carbon across the planetary surface. As global warming proceeds, it is expected that photosynthetic CO₂ uptake will increase in colder regions of the world and diminish in those regions that are already warm and dry (1). At the same time, warming is expected to increase microbial activity, at least where water is not limiting, and therefore lead to an enhanced breakdown of organic matter in the soil, producing a large respiratory flux of CO₂ back to the atmosphere (2). Because terrestrial ecosystems presently sequester about a quarter of the CO₂ emissions associated with fossil fuel burning $(7.1 \,\mathrm{GtC}\,\mathrm{y}^{-1})$ (1), it is critical that we understand how large-scale, climate-driven changes will affect the carbon sequestration of the terrestrial biosphere. Currently, the precise response of terrestrial CO₂ sources and sinks to changes in climate remains uncertain (3) and its understanding requires the ability to quantify the amount of CO₂ taken up during photosynthesis separately from the amount released by respiration.

atmospheric CO2 during photosynthetic and respiratory CO2 exchange, via an isotopic exchange during CO_2 hydration (7): $CO_{2aq} + H_2^{18}O \rightleftharpoons CO^{18}O_{aq} + H_2O$. Despite the short residence time of CO_2 in leaves, CO_2 involved in photosynthesis is nearly completely relabeled by ¹⁸O-enriched leaf water because of the enzyme carbonic anhydrase (CA; EC 4.2.1.1), a very efficient catalyst of CO_2 hydration and isotopic exchange (4, 5, 8, 9). Typically the δ^{18} O of leaf and soil water pools are very different. There is a tendency for the heavier molecules of water to accumulate more readily in leaves than in soils during evapotranspiration because of the difference in water pool size (10, 11). Because the CO_2 -H₂O exchange in leaves (associated with photosynthesis) or soils (associated with soil respiration) produces such contrasting ¹⁸O signals, estimates of the amount of CO₂ exchanged during photosynthesis and respiration can in principle be constrained by using the δ^{18} O signal of atmospheric CO₂ (6, 12).

However, our ability to partition gross fluxes of CO₂ may be complicated because the δ^{18} O of soil water (δ_{sw}) can often display a strong vertical gradient at the soil surface because soil evaporation also leads to an enrichment of heavy water molecules in the uppermost layers (13–15). Thus, to determine the δ^{18} O of CO₂ exchanged between soils and the atmosphere accurately it becomes necessary to know the shallowest depth (z_{eq}) where diffusing CO₂ molecules (from the atmosphere or produced by soil respiration; Fig. 1*A*) have enough time to fully equilibrate isotopically with soil water. With increasing temperature and moisture, CO₂ hydration increases relative to the diffusion rate so that z_{eq} moves closer to the surface, and toward more enriched δ^{18} O values (see *Methods*, Eq. 4). Although we know that CA accelerates the rate of hydration in leaves, the possibility of CA activity in soils is commonly neglected (4, 15),

The oxygen isotope composition of atmospheric CO₂ (δ_a) was shown to be a powerful tracer of photosynthetic and respiratory CO₂ fluxes while at the same time providing information on the intensity of water cycling within terrestrial ecosystems (4–6). This tracing property occurs because the oxygen isotope composition (δ^{18} O) of leaf and soil water pools is transferred to

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¹L.W., J.O., and M.C. contributed equally to this work.

²To whom correspondence may be addressed. E-mail: I.wingate@ed.ac.uk or jogee@bordeaux.inra.fr.

³Present address: Helmholtz Centre for Environmental Research, Zentrum für Umweltforschung, 04318 Leipzig, Germany.

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Fig. 1. Schematic showing the influence of CO₂ hydration rates on vertical profiles of $\delta^{18}O$ in soil air CO2. (A) The net soil-atmosphere CO2 exchange is composed of CO_2 molecules moving from the atmosphere into the soil and back to the atmosphere (i.e., invasion) and further CO2 molecules produced during soil respiration. Because of oxygen isotopic exchange between soil CO2 and water, both invasion and respiration fluxes modify the isotopic composition of atmospheric CO₂, and their ¹⁸O isotopic signature depend on the extent of CA activity in the soil. (B) Typical profile of δ^{18} O in soil air CO₂ (δ_{sa}) for uncatalyzed CO₂ hydration in soil water (enhancement factor $f_{CA} = 1$). (C) Same as in *B* but for catalyzed CO₂ hydration (enhancement factor $f_{CA} \gg 1$). In deep soil layers where vertical gradients of δ_{sw} are weak, the residence time of CO₂ is long enough to reach full isotopic equilibrium with soil water (δ_{sa} = $\delta_{sw} - \varepsilon_{eq}$), where ε_{eq} denotes the isotopic equilibrium fractionation between CO₂ and water (22). Above a certain depth z_{eq} (where, by definition, $\delta_{sa} = \delta_{eq}$), CO2 molecules diffuse too rapidly to fully equilibrate with local soil water. If CO_2 hydration is enhanced because of CA activity ($f_{CA} \gg 1$), the equilibration becomes faster and z_{eq} shallower, thus δ_{eq} becomes more enriched.

because the abundance and location of CA in soils is still somewhat unclear, with only indirect and isolated indications based on measurements of δ^{18} O of soil CO₂ or COS fluxes (14, 16, 17). Substantial CA activity in soils would lead to a faster equilibration of CO₂, moving z_{eq} further toward the surface where soil water is more ¹⁸O enriched (Fig. 1 *B* and *C*). So far global simulations have assumed uncatalyzed CO₂ hydration in soils (18–20) and equilibration depths below the region of strong evaporative enrichment (5, 21).

Results and Discussion

Evidence for Enhanced Soil CO2 Hydration Rates. Here, we demonstrate that, in contrast to current assumptions, the observed rate of soil CO₂ hydration is always substantially faster than the uncatalyzed rate. We compared measurements of depth-resolved soil water $\delta^{18}O(\delta_{sw})$ and observed $\delta^{18}O$ signatures of chamber-based soil CO₂ fluxes (δ_{flux}) in seven different ecosystems that encompass most of the major land biomes, providing a global perspective of ¹⁸O exchange in soils (Table S1 and see Tables S5–S7). From the δ_{sw} data, we determined the depth-resolved $\delta^{18}O$ of soil CO₂ in full equilibrium with soil water (δ_{eq}), equal to $\delta_{sw} - \varepsilon_{eq}$ where ε_{eq} is the temperature-sensitive equilibrium fractionation between CO2 and water (22). Most sites exhibited strong gradients in $\delta_{sw} - \varepsilon_{eq}$ at the soil surface, reflecting the evaporative enrichment of soil water (Fig. 2). From the δ_{flux} data, we determined the δ^{18} O of soil CO₂ at z_{eq} (δ_{eq} , see Fig. 1) for different rates of hydration expressed as an enhancement factor (f_{CA}) with respect to the uncatalyzed CO₂ hydration rate (see Eq. 6 in *Methods*). Increasing f_{CA} shifts z_{eq} toward surface layers (Fig. 2) and δ_{eq} toward δ_{a} . The best estimate for f_{CA} would be one in agreement with both soil water and chamber flux measurements. This is obtained when the point (δ_{eq} , z_{eq}) derived from the chamber data intersects the δ_{eq} curve derived from soil water measurements. At all sites, this intersection occurs for values of f_{CA} between 10 and 300, with the lowest f_{CA} in the cooler temperate ecosystems while higher f_{CA} were found at the Mediterranean and subtropical sites (Fig. 2). As a consequence, the equilibration depth z_{eq} was in most cases within the top 5 cm of the soil, the zone containing the strongest δ_{sw} gradients. A reduction in the effective diffusivity of soil CO2 would also lead to shallower equilibration depths z_{eq} by increasing the residence time of CO₂ in soils, but it would not yield simultaneous solutions for both soil water and CO_2 flux isotope data (14). Thus, an enhanced CO_2 hydration rate is the only plausible mechanism to explain these chamber-based measurements.

Consistency with CA Activities in Soil Microorganisms. The uppermost soil layers host many bacterial, algal, and fungal species that produce intracellular and sometimes extracellular CAs (23–25). Based on a literature survey, we claim that this mixed population



Fig. 2. The δ^{18} O of soil CO₂ at the depth of full equilibration (δ_{eq} , z_{eq} ; see Fig. 1) estimated from chamber flux measurements for different levels of hydration rates (f_{CA}). Depth-resolved soil water data yields the δ^{18} O of CO₂ in isotopic equilibrium with soil water ($\delta_{sw} - \varepsilon_{eq}$). The point at which the two curves intersect indicates the most likely value for the enhancement factor, f_{CA} , listed below the ecosystem type for each site (see Table S1). The horizontal error bars on the squared symbols represent the standard deviation of δ_{eq} values over the number of δ_{flux} measurements (n = 1-15).

of soil microorganisms is responsible for the accelerated soil CO2 hydration. Most soils contain 10³ to 10⁶ algae per g of dry soil, but populations can reach 10^8 algae per g of dry soil (Table S2). Bacterial population sizes are even larger at 10⁸ to 10⁹ cells per g of dry soil (Table S3). At 25 °C, the CO₂ hydration rate in soil algal and cyanobacterial cells can be up to 172,500 times the uncatalyzed rate, comparable to CA activities found in plant chloroplasts (Table S4). With a cell volume of $\approx 100 \ \mu m^3$ and population of 10⁶ per g of dry soil, algae could explain a significant fraction of our observed soil CA activities. Indeed, we found that the presence of algae developing naturally on the surface of a peat soil dramatically enhanced CA activity (Table S4). Laboratory studies have also reported high CA activities $(f_{CA} = 50)$ in bulk soil extracts from subtropical karst forests containing a mixture of bacterial and fungal species (Table S4). Based on a cell volume of $\approx 1 \,\mu \text{m}^3$ and population of 4×10^9 cells per g of dry soil (Table S3), this soil-level f_{CA} value would be consistent with soil bacteria operating at a cell-level CO₂ hydration of 8,000 times the uncatalyzed rate (Table S4). These estimates demonstrate that soil microorganisms are likely to be responsible for enhanced soil CO₂ hydration rates of 20-300 times the uncatalyzed rate.

Impact of Soil CA Activity at the Global Scale. The accelerated hydration of CO₂ in soils has been missing in the mass budget of δ^{18} O in atmospheric CO₂. To explore the impact of soil CA activity on the δ^{18} O of atmospheric CO₂, and its north–south (N-S) gradient, we incorporated this biological process into the global model of δ^{18} O in atmospheric CO₂, Mecbeth (18, 19) (see *Methods*). Simulations were performed over an average year calculated from the 1990s and compared with observations from the worldwide network of atmospheric stations for the same decade (26, 27). Three scenarios are discussed here: one uncatalyzed (abiotic) scenario ($f_{CA} = 1$) and two globally uniform f_{CA} scenarios covering the range of soil chamber estimates ($f_{CA} = 20$ and $f_{CA} = 300$).

The high CA activity scenario ($f_{CA} = 300$) improves the agreement between the modeled and observed N-S gradient in δ_a , particularly when compared with the uncatalyzed, abiotic scenario (Fig. 3). This latitudinal feature in δ_a is largely driven by the N-S gradient in the δ^{18} O of precipitation that creates depleted leaf and soil water pools toward the northern latitudes. In the uncatalyzed scenario ($f_{CA} = 1$), photosynthesis dominates the N-S gradient. Introducing f_{CA} enhances the invasion flux (the number of CO₂ molecules from the atmosphere that equilibrate with soil water and go back to the atmosphere; see Fig. 1*A* and *Methods*, Eq. 5). The contribution of this invasion flux to the N-S gradient increases with f_{CA} and at some latitudes becomes larger than the contribution of respiration.

Incorporating high CA activity ($f_{CA} = 300$) also reduces the mean value of δ_a by $\approx 1\%$ relative to the abiotic case ($f_{CA} = 1$), bringing the model closer to atmospheric observations. This reduction is the result of complex interactions between δ_a and the δ^{18} O signatures of all component fluxes. The influence of each process on δ_a can be represented by using the concept of isoflux (I_x) defined as the product of a gross CO₂ flux (F_x) and its isotopic composition (δ_x) relative to δ_a : $I_x = F_x (\delta_x - \delta_a)$. Photosynthesis tends to enrich the atmosphere (positive isoflux, $\delta_x > \delta_a$), whereas respiration and soil invasion usually have the opposite effect (negative isoflux, $\delta_x < \delta_a$). Nonbiospheric fluxes (from ocean, fossil fuel, and biomass burning) also tend to deplete δ_a , but to a much lesser extent (19, 28). In the uncatalyzed scenario ($f_{CA} = 1$), photosynthetic (I_A), and respiratory $(I_{\rm R})$ isofluxes balance the nonbiospheric isofluxes globally while the soil invasion isoflux (I_{inv}) remains close to zero at all latitudes (Fig. 3). When f_{CA} is increased, the isotopic signatures of soil invasion and respiration become progressively enriched as a result of the isotopic gradient in δ_{sw} (Fig. 1), but usually remain



Fig. 3. Simulated contributions of different biospheric processes to the N-S gradient in δ_a for uncatalyzed ($f_{CA} = 1$) or enhanced ($f_{CA} \gg 1$) CO₂ hydration rates in the soil, compared with measured δ_a . Enhanced hydration increases the corresponding isofluxes of soil invasion and photosynthesis + respiration, i.e., the isotopic imbalance required for gross flux partitioning. Note that δ_a values are always reported relative to the South Pole and thus do not show the absolute changes in δ_a (-0.1‰ and -1.1‰ for f_{CA} of 20 and 300 relative to $f_{CA} = 1$, respectively).

below δ_a (I_R and I_{inv} remain negative). Most importantly, the mass of atmospheric CO₂ molecules that equilibrate with soil water increases from 25 GtC yr⁻¹ in the uncatalyzed scenario to 450 GtC yr⁻¹ when $f_{CA} = 300$. As a result, I_{inv} becomes very large and negative, reaching nearly the same magnitude as soil respiration. The associated depletion in δ_a is partly compensated by an increase in both I_A and I_R . When $f_{CA} = 300$, the absolute value of I_{inv} increases by 571 GtC % o yr⁻¹, whereas I_R decreases by 226 GtC % o yr⁻¹ and I_A increases by 269 GtC % o yr⁻¹. Soil CA activity thus strongly modifies the relative contribution of photosynthesis and respiration to the CO¹⁸O budget in our global model.

Consequences for the Retrieval of CO₂ Sources and Sinks. Measurements of δ_a have been proposed as one of the few tools available to partition net CO₂ fluxes into photosynthesis and soil respiration, but it critically depends on the existence of sufficient imbalance between I_A and I_R (4–6, 9, 12, 21, 28). In previous studies, where δ_a was prescribed and not dynamically coupled to soil and leaf water pools as in our study, such an imbalance had been restricted to the boreal regions (5, 21, 28). In contrast, we now show a significant isotopic imbalance at nearly all latitudes (photosynthesis + respiration curve in Fig. 3), greatly enhancing the potential of the ¹⁸O approach for partitioning CO₂ fluxes. At the continental scale, there is a strong isotopic imbalance over Europe and North America coinciding with the peak season of photosynthetic activity in the Northern Hemisphere (Fig. 4). Over the tropics, the isotopic imbalance between I_A and I_R increases by up to 50% when $f_{CA} = 300$ and is maintained year-round in many areas (Fig. 4). The uncertainties of tropical gross CO₂ fluxes could thus be reduced by an equivalent amount (12), making the δ^{18} O of atmospheric CO₂ a better tracer for terrestrial gross CO₂ fluxes than previously thought in these regions. From a one-box global mass balance budget and using globally averaged fluxes and isotopic signatures from Mecbeth, we calculated that, neglecting "biotic" invasion associated with soil CA activity in inversion studies leads to errors in the isotope-derived estimates of global photosynthesis by up to 30 GtC yr⁻¹, i.e., $\approx 30\%$ of current estimates (1).



Fig. 4. Global distribution in the extent of isotopic imbalance ($I_A + I_R$) across continental surfaces for June and December simulated by the global model Mecbeth for the most enhanced soil CO₂ hydration scenario ($f_{CA} = 300$). Regions where $I_A + I_R$ is the most different from zero correspond to regions of strong isotopic imbalance where biospheric gross CO₂ fluxes are expected to be the most constrained by δ^{18} O data.

Future Directions for Global Isotope-Enabled Models. This study demonstrates that enhanced rates of CO2 hydration occur at the soil surface and appreciably impact the oxygen isotope composition of atmospheric CO₂. This enhanced exchange in the soil brings into focus our limited ability to predict the isotopic enrichment of soil water near the surface (18, 29), highlighting a need for future improvements in this research area. Also, although we provided the basic observations and parameterization, more work is now needed to further assess the variability in f_{CA} in different ecosystems, plant functional types, or regions within the global model, including attempts to establish the mechanistic basis to underpin the observed differences in CA activity between ecosystems. Developments on these fronts will greatly enhance our capabilities to use the δ^{18} O of atmospheric CO₂ to quantitatively inform us of large-scale changes in the intensity of carbon and water cycling in terrestrial ecosystems.

Methods

Soil CO¹⁸O Budget Equation. In a given soil layer, the number of moles of CO¹⁸O changes as a result of (*i*) CO¹⁸O production during heterotrophic and autotrophic respiration, (*ii*) diffusion of these molecules through the soil layer, and (*iii*) oxygen isotopic exchange with the surrounding soil water (30–32):

$$\theta_{\rm t} \frac{\partial C\mathcal{R}}{\partial t} = \mathcal{R}_{\rm c} S_{\rm c} + \frac{\partial}{\partial z} \left[D_{\rm c,iso} \frac{\partial C\mathcal{R}}{\partial z} \right] + k_{\rm h,iso} B \theta_{\rm w} C(\mathcal{R}_{\rm eq} - \mathcal{R}),$$
[1]

where C [mol·mol⁻¹] is the CO₂ mole fraction in soil air, \Re , \Re_c , and \Re_{eg} are the ¹⁸O/¹⁶O ratios of the CO₂ in soil air, respired CO₂, and CO₂ in isotopic equilibrium with the surrounding soil water, respectively, S_{c} (mol $m^{-3} \cdot s^{-1}$) is the respiration rate density, $D_{c,iso}$ (m²·s⁻¹) is the effective diffusivity of CO¹⁸O in soil air, θ_w (m³·m^-³) is the volumetric soil water content, B is the CO2 solubility coefficient, and θ_t (m³·m⁻³) is the total CO₂ porosity. Denoting by θ_a the soil air porosity we have (31): $\theta_t = \theta_a + B \theta_w$. The solubility coefficient *B* depends on soil temperature T_s (K) according to ref. 33: $B = 1.739 \exp(-0.039(T_s - 273.15) +$ 0.000236(T_s - 273.15)²). \Re_{eq} is related to the $^{18}\text{O}/^{16}\text{O}$ ratio in soil water \Re_{sw} through $\Re_{eq} = (1 + \varepsilon_{eq}) \Re_{sw}$, where $\varepsilon_{eq} = 17.604/T_s - 0.01793$ is the CO₂-H₂O equilibrium fractionation (22). Because there are three oxygen atoms present in the bicarbonate intermediate, the isotopic exchange rate during CO2 hydration equals one-third the hydration rate (7): $k_{h,iso} = f_{CA}k_{h,uncat}/3$, where (34) $k_{h,uncat} = 0.037 \times \exp(0.118(T_s - 298.15))$. In this framework, CA activity is expressed as an enhancement factor (f_{CA}) of the uncatalyzed CO₂ hydration rate ($k_{h,uncat}$). The effective CO¹⁸O diffusivity in soil air is calculated as $D_{c,iso} =$ $D_{c,eff} \alpha_{d}$, where $\alpha_{d} = 0.9913$ is the isotopic discrimination during molecular diffusion of CO₂ in air and $D_{c,eff}$ (m²·s⁻¹) is the effective CO₂ diffusivity in soil air. Several parameterizations of this effective diffusivity exist in the literature that differ mostly for wet soils (35). Results presented in this study use ref. 31: $D_{\rm c.eff} = 0.66 \times \theta_{\rm a} \times 1.4 \cdot 10^{-5} (T_{\rm s}/298.15)^{1.75}.$

Full Equilibration Depth. The budget equation above contains two time scales. One time scale indicates the half-life of CO_2 molecules before being isotopically equilibrated with the surrounding water:

$$\tau_{\rm k} = \ln 2 \cdot \left(\frac{\theta_{\rm t}}{k_{\rm h,iso} B \, \theta_{\rm w}} \right)$$
[2]

and another time scale indicates the time required for a plume of C¹⁸OO molecules to diffuse through the soil over a given distance z:

$$\tau_{\rm d}(z) = \frac{\theta_{\rm t} z^2}{2D_{\rm c.iso}}.$$
 [3]

Full equilibration within a soil layer of thickness z is satisfied when the time scale for isotopic equilibration is smaller than the time scale for diffusion through this layer, i.e., $\tau_{k} \ll \tau_{d}(z)$. When $\tau_{k} = \tau_{d}(z)$, full equilibration can occur if the soil layer has uniform soil temperature, moisture content, and isotopic composition. However, in the top centimeters of the soil, strong gradients of T_{s} , θ_{w} , and \Re_{w} are more likely. The shallowest depth of full equilibration, z_{eq} , must therefore satisfy the inequality: $\tau_{k} < \tau_{d}(z_{eq})$. In the following we will define z_{eq} as: $\tau_{k} = \tau_{d}(z_{eq})/4$, or similarly:

$$z_{\rm eq} = 2 \sqrt{\frac{2\ln 2D_{\rm c,iso}}{k_{\rm h,iso}B\,\theta_{\rm w}}}.$$
 [4]

The factor 4 was determined by matching the value of f_{CA} deduced in Fig. 2 with that obtained from simulations using the full numerical model (Eq. 1), i.e., $f_{CA} \approx 300$ for the Mediterranean evergreen site (14) and $f_{CA} \approx 20$ for the montane evergreen site (15). Eq. 4 with $f_{CA} = 20$ also provides seasonal variations of z_{eq} at the temperate evergreen site that correspond to the depth where $\delta^{18}O$ in soil air $CO_2 (\delta_{sa})$ and $\delta_{sw} - \varepsilon_{eq}$ (estimated using the full numerical model, Eq. 1) start to diverge by >0.3 ‰ (a threshold chosen for practical purposes to represent the overall precision of soil water isotope measurements).

Other studies (14, 35) use a different formulation for $D_{c,iso}$, leading to values of this diffusivity 5-fold smaller in saturated soils. Using this other formulation does not fundamentally change the results presented in Fig. 2.

Soil CO₂ Isoflux. In the steady state, and assuming isothermal and uniform soil water conditions, Eq. 1 can also be solved analytically (30–32). In this framework, the isotopic composition of the soil CO₂ flux δ_{flux} is:

$$\delta_{\rm flux} = \delta_{\rm eq} + \varepsilon_{\rm d,eff} + (\delta_{\rm eq} - \delta_{\rm a}) v_{\rm inv} \frac{C_{\rm a}}{F_{\rm R}},$$
[5]

where $\varepsilon_{d,eff}$ is the effective isotopic fractionation during diffusion, F_R is the soil CO₂ efflux, and $v_{inv} = \sqrt{B\theta_w k_{h,iso} D_{c,iso}}$ has the dimensions of a velocity (m·s⁻¹) that when multiplied by C_a gives the soil invasion flux F_{inv} . The product ($\delta_{flux} - \delta_a$) F_R is called the soil CO₂ isoflux. It can be seen as the sum of two isotope fluxes: a respiration isoflux, $I_R = (\delta_{eq} + \varepsilon_{d,eff} - \delta_a)F_R$, and an invasion isoflux, $I_{inv} = (\delta_{eq} - \delta_a)F_{inv}$, sometimes defined as abiotic because it is independent of

 F_R . Assuming a uniform soil CO₂ production S_c over a soil column of depth z_0 , $\varepsilon_{d,eff}$ can be estimated as (31): $\varepsilon_{d,eff} = \varepsilon_d(1 - z_1/z_0(1 - \exp(-z_0/z_1)))$, where $z_1 = (2\sqrt{2\ln 2})^{-1} z_{eq}$. Eq. 5 can then be inverted to estimate δ_{eq} as a function of δ_{flux} , C_{ar} , δ_{ar} , and F_R measurements:

$$\delta_{\rm eq} = \frac{\delta_{\rm flux} - \varepsilon_{\rm d,eff} + v_{\rm inv}C_{\rm a}/F_{\rm R}\delta_{\rm a}}{1 + v_{\rm inv}C_{\rm a}/F_{\rm R}}.$$
 [6]

Oxygen Isotope Composition of the Net CO₂ Flux from Soil Chambers. The steady-state oxygen isotope signal of the net soil CO₂ flux during chamber closure (δ_{ch}) was calculated by using a simple isotopic mass balance:

$$\delta_{\rm ch} = \frac{\delta_{\rm out} C_{\rm out} - \delta_{\rm in} C_{\rm in}}{C_{\rm out} - C_{\rm in}},$$
[7]

where C_{out} , C_{in} and δ_{out} , δ_{in} are the mole fractions and isotopic compositions of CO₂ in the air leaving and entering the chamber, respectively. In the case of the two sites that used closed chambers (subtropical evergreen and semiarid grassland), C_{out} , C_{in} and δ_{out} , δ_{in} are the mole fractions and isotopic compositions of CO₂ at the start and end of a defined chamber closure period, respectively.

To derive δ_{eq} values from soil chamber data, we use Eq. 6, neglect chamber effects, and make the common assumption that the atmosphere inside the chamber is well mixed ($C_a = C_{out}$ and $\delta_a = \delta_{out}$).

Oxygen Isotope Composition of Soil Water. Depth-resolved soil samples were collected at each experimental site within proximity of the soil chamber and at approximately the same time as gas exchange measurements. In the case of the Mediterranean evergreen, subtropical evergreen, and both temperate ever-

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green sites, soil water was extracted cryogenically from bulk soil samples and $\delta^{18}O$ analysis of CO₂ equilibrated with the extracted water was completed (14). For the montane evergreen, subtropical deciduous, and semiarid grassland sites CO₂ with a known isotopic composition was equilibrated directly with fresh soil samples and stored in gas-tight containers for 12 h. Equilibrated CO₂ was then sampled from the container and analyzed for its $\delta^{18}O$ composition (15).

Global Model Simulations. The global model Mecbeth calculates the sources and sinks of CO₂, water, and their respective isotopes and transports them in the atmosphere (18, 19). It merges a description of the biospheric energy, water, and carbon fluxes with a global climate and water isotope model. The atmosphere and biosphere are dynamically coupled to account for feedbacks of the accelerated equilibration of CO₂ with soil water on δ_a and the isotopic signatures of leaf and other fluxes. The model parameterization of soil water isotopes was improved in this study to provide depth-resolved descriptions of soil water and soil water isotopes (35), a necessary step if CA activity occurs in soils containing strong vertical gradients in δ_{sw} (14). Several soil layers of varying thickness were included in the model. The most important upper layers relevant to this study consisted of a top layer at 0–6 cm and another layer at 6–20 cm.

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