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A NUMERICAL EXAMINATION OF ¹⁴CO₂ CHAMBER METHODOLOGIES FOR SAMPLING AT THE SOIL SURFACE

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ABSTRACT. Radiocarbon is an exceptionally useful tool for studying soil-respired CO₂, providing information about soil carbon turnover rates, depths of production, and the biological sources of production through partitioning. Unfortunately, little work has been done to thoroughly investigate the possibility of inherent biases present in current measurement techniques, like those present in $\delta^{13}CO_2$ methodologies, caused by disturbances to the soil's natural diffusive regime. This study investigates the degree of bias present in four ¹⁴C sampling chamber methods using a three-dimensional numerical soil-atmosphere CO₂ diffusion model. The four chambers were tested in an idealized, surrogate reality by assessing measurement bias with varying $\Delta^{14}C$ and $\delta^{13}C$ signatures of production, collar lengths, soil biological productivity rates, and soil diffusivities. The static and Iso-FD chambers showed almost no isotopic measurement bias, significantly outperforming dynamic chambers, which demonstrated biases up to 200‰ in some modeled scenarios. The study also showed that ¹³C and ¹⁴C diffusive fractionation are not a constant multiple of one another, but that the $\delta^{13}C$ correction still works in diffusive scenarios because the change in fractionation is not large enough to impact measured $\Delta^{14}C$ values during chamber equilibration.

INTRODUCTION

The radioactive isotope of carbon (14 C) is an exceptionally useful tool for studying soil-respired CO₃, providing information about the biological sources of production through partitioning (Gaudinski et al. 2000; Trumbore 2000; Hahn et al. 2006; Schuur and Trumbore 2006; Hicks Pries et al. 2013). In recent years, many studies have utilized partitioning techniques, both physical and isotopic, as tools for separating sources of soil respiration, to understand how soil respiration sources may be affected by the future changing climate (Hanson et al. 2000; Högberg et al. 2001; Singh et al. 2003; Lee et al. 2003; Kuzyakov 2006; Moyes et al. 2010; Bond-Lamberty et al. 2011; Drake et al. 2012; Gomez-Casanovas et al. 2012; Risk et al. 2012). Source partitioning with isotopes has an advantage over physical partitioning as it is typically involves less disturbance than physical partitioning. However, in natural abundance isotopic partitioning studies, ¹⁴C can be a more sensitive tool than $\delta^{13}C$. The difference between autotrophic and heterotrophic $\delta^{13}C$ signatures of soil-respired CO₂ is only a few permil (‰) (except in C₃-C₄ vegetation shifted studies), whereas there can be a much larger separation between Δ^{14} C source signatures, especially in systems where slow decomposition or long-term storage accentuate isotopic differences (Trumbore 2006). A peak in atmospheric Δ^{14} C signatures in 1963 caused by nuclear weapons testing has allowed researchers to utilize ¹⁴C as a tracer to distinguish whether carbon substrates were utilized pre- or post-bomb, because post-bomb signatures are distinctive given their relative ¹⁴C enrichment (Levin and Hesshaimer 2000). Autotrophic respiration consumes new carbon, so its ¹⁴C signature will reflect current atmospheric ¹⁴CO₂ signatures, whereas heterotrophic signatures will reflect the age of the substrates that the heterotrophs consume, which can be very new or quite old (Gaudinski et al. 2000; Phillips et al. 2013).

Despite the potential utility of ¹⁴CO₂ as a tool for investigating soil-respired CO₂, little work has been done to thoroughly investigate the possibility of biases inherent to existing measurement techniques, because the high cost of analysis naturally drives researchers to focus effort on the ecological aspect of studies, rather than error or uncertainty testing. In the case of δ^{13} C, Cerling et al. (1991) demonstrated that although mass differences in ¹²C and ¹³C isotopologues cause ¹²C to diffuse 1.0044 times faster through the soil, if the soil is at a diffusive steady-state, the δ^{13} C of production should match the δ^{13} C of surface flux. Soils are, however, rarely at a diffusive steady-state, and

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research looking at δ^{13} C has shown that non-steady-state transport fractionations can be induced by soil transport (Nickerson and Risk 2009a; Moyes et al. 2010) and headspace sampling (Ohlsson 2010), where non-steady-state chambers can induce a bias of 4‰ and steady-state chambers up to 15‰ caused by lateral diffusion (Nickerson and Risk 2009b). The magnitude of these biases could potentially overprint the δ^{13} CO₂ signatures of biological flux (Kayler et al. 2010; Moyes et al. 2010; Phillips et al. 2010), making partitioning difficult. These time-dependent fractionations ("dynamic fractionations"; Nickerson and Risk 2009a) will also be present and will potentially cause biases when attempting to measure ¹⁴CO₂, because like ¹³CO₂, ¹⁴CO₂ has a different diffusion coefficient than ¹²CO₂, diffusing 1.0088 times slower (Wang et al. 1994). Currently, to calculate Δ^{14} C, researchers use δ^{13} C to correct for potential steady-state mass-dependent fractionations (Stuiver and Polach 1977). The δ^{13} C correction will not account for dynamic fractionations, however, because the assumption that ¹⁴C fractionation is a constant multiple of ¹³C fractionation may not hold in typical measurement conditions where the soils are rarely at steady-state. It will therefore be of interest to the ¹⁴C community to know if steady-state and non-steady-state chamber methods used to measure Δ^{14} C of soil-respired CO₂ induce bias in a similar magnitude to those of δ^{13} C.

In order for researchers to understand bias that may have been induced by ¹⁴CO₂ chamber methodologies in past studies, and to help decide which chamber method is the most robust for future studies, we utilized a three-dimensional numerical soil-atmosphere model to simulate soil-atmosphere-chamber exchange of CO₂ isotopologues, including ¹⁴CO₂, to compare isotopic signatures measured by various chamber methods to the natural steady-state. We also examined the effect of chamber-specific protocols, such as pump speed, stable isotopic signature calculations, and mixing model type, to determine the effect of these on the final estimates of stable and radioactive isotopic signature. Additionally, we investigated whether Δ^{14} C of soil flux changed during equilibration, and the amount by which ¹³CO₂ and ¹⁴CO₂ fractionation factors differ from one another during chamber equilibration. This was done in order to understand whether a biased stable isotopic value for flux used as a correction in the calculation of Δ^{14} C will affect the results. We predict that static and dynamic chambers will induce more bias measuring Δ^{14} C of flux than they do for δ^{13} C, as the diffusive fractionation factor for ¹⁴CO₂ is larger (Stuiver and Polach 1977), and furthermore because additional error could be related to the δ^{13} C correction in a dynamic environment.

METHODS

Soil-Atmosphere Model

A three-dimensional coupled soil-atmosphere-chamber model was used to explore biases inherent in current ${}^{14}\text{CO}_2$ chamber methodologies. The model simulates CO₂ produced in the soil, diffusion to the atmosphere, and in some cases into a chamber on the surface of the modeled soil. Model physics and structure are further described in Creelman et al. (2013); the present version of the model adds ${}^{13}\text{CO}_2$ and ${}^{14}\text{CO}_2$ functionality in addition to ${}^{12}\text{CO}_2$.

Diffusion of each carbon isotopologue of CO_2 (${}^{12}CO_2$, ${}^{13}CO_2$, and ${}^{14}CO_2$) is modeled according to Fick's law with its own specific atmospheric concentration, production, and diffusion rate. The simulations described here build on related work by Risk and Kellman (2008) and Nickerson and Risk (2009b), which used similar Fick's law-based models to examine the impacts of non-steady-state conditions on the soil surface flux of stable C isotopologues of CO_2 . The model reproduces patterns of atmospheric invasion and soil air enrichment described in seminal papers on natural steady-state soil isotopic composition (Cerling et al. 1991; Wang et al. 1994; Davidson 1995; Amundson et al. 1998). The use of numerical simulation allows us to address both steady-state and time-dependent (non-steady-state) system dynamics that are not encompassed in the analytical solutions presented in the earlier research. Drivers of these non-steady-state dynamics include soils undergoing rapid day-night changes in respiration rate, wind-induced advection, or temporary disruption of the natural steady-state regime owing to emplacement of a chamber on the surface (Risk and Kellman 2008; Venterea and Baker 2008; Bowling et al. 2009; Nickerson and Risk 2009a,b; Kayler et al. 2010; Phillips et al. 2010; Creelman et al. 2013). These settings have been shown to induce transient fractionations that are not predicted by the steady-state literature.

The model assumes that radioactive decay is negligible for ¹⁴CO₂ over the timescale of the measurements (Cerling et al. 1991; Wang et al. 1994). We produced model runs for each isotopologue where the resulting concentrations and fluxes of each isotopic species are used to calculate total CO₂, δ^{13} C, and Δ^{14} C signatures of the soil CO₂ and surface flux at each time step.

Simulations were performed using a set of soil production rates, soil diffusivity coefficients, chamber collar lengths (downward extent of the collar below the soil), and δ^{13} C and Δ^{14} C signatures of biological production (Table 1). In all model scenarios, the δ^{13} C and Δ^{14} C of the atmosphere were 8.8‰ and 100‰, respectively, which are in the range of most typical environments.

Table 1 Model parameters.	
Variable	Ranges
Diffusivity (m ² /s)	$10^{-8}, 5 \times 10^{-8}, 10^{-7}, 5 \times 10^{-7}, 10^{-6}, 5 \times 10^{-6}$
Productivity (µmol/m²/s)	0.1, 1, 5
Collar length (cm)	0, 2, 4, 8
δ^{13} C of production (‰)	-30, -20, -15
Δ^{14} C of production (‰)	-500, -200, 0, 200, 500

Calculation of Isotopic Signatures

The model output calculates isotopic signatures using delta notation. $\delta^{13}C$ is presented in ‰ as

$$\delta^{13}C = \left(\begin{pmatrix} R_{s} \\ R_{std} \end{pmatrix} - 1 \right) \times 1000 \tag{1}$$

where R_s is the ¹³C/¹²C ratio of the sample and R_{std} is the ¹³C/¹²C ratio of the PDB standard. For radiocarbon, Δ^{14} C is also presented in ‰ using the value of fraction modern (*FM*) of the sample:

$$\Delta^{14}C = (FM - 1) \times 1000$$
⁽²⁾

FM is calculated using A_s , the measured activity of the sample, where the model outputs a ¹⁴C/¹²C ratio, which is equivalent to A_s to the fourth decimal place (Southon 2011). A_{abs} is the measured activity of the oxalic acid standard, with a model parameterization value of 1.2511e⁻¹², where A_{abs} is 0.95 times the activity of the oxalic acid standard in 1950, corrected to a δ^{13} C of –19‰ (Stuiver and Polach 1977):

$$FM = \frac{\left(\frac{A_s}{A_{abs}}\right) \left(1 - \frac{25}{1000}\right)^2}{\left(1 + \frac{\delta^{13}C_s}{1000}\right)^2}$$
(3)

where $\delta^{13}C_s$ is the $\delta^{13}C$ signature of the sample, and 25 is the sample activity corrected for $\delta^{13}C$ isotopic fractionation.

Chamber Descriptions

Four chambers utilized in previous studies were simulated, and are described below and represented graphically in Figure 1. All calculations to determine chamber δ^{13} C and Δ^{14} C from the model output were performed using the equations provided in the original studies. All results were converted to Δ^{14} C notation in cases where the results were calculated in percent modern or fraction modern, to facilitate comparison of the results between different chambers.



Figure 1 The four chambers that were modeled: (A) A dynamic chamber with a pump speed set to 0.5 L/min (Gaudinski et al. 2000); (B) A dynamic chamber with a pump speed that matches the flux rate (Schuur and Trumbore 2006); (C) A static chamber where CO_2 accumulates for 30 min prior to sampling (Hahn et al. 2006); (D) A diffusion-driven dynamic chamber, which is based on the isotopic-forced diffusion (Iso-FD) design for a $\delta^{13}C$ chamber presented in Nickerson et al. (2013).

Dynamic Chambers

Dynamic chambers are a type of steady-state chamber often employed in soil flux studies, including ¹⁴C studies. They theoretically minimize alteration to the soil CO₂ concentration gradient by flowing atmospheric air or a CO₂-free gas through the chamber, which decreases the headspace concentration in order to maintain chamber concentrations closer to the surrounding atmosphere (Rayment and Jarvis 1997). We chose two dynamic chambers to model. Chamber A has a volume of ~11.5 L and is based on the chamber presented in Gaudinski et al. (2000). It utilized a pump with a constant flow rate of 0.5 L/min, and soda lime column to decrease the headspace concentration, regardless of the soil flux value at the time of measurement, just before diverting the flow path into a molecular sieve trap to collect 2 mg C for ¹⁴C analysis. For modeling purposes, we assumed that the soda lime and molecular trap were perfectly efficient at removing CO₂ from the air stream, and further, that there was no isotopic fractionation associated with either process. Stable isotopic (δ^{13} C) signatures of surface flux were used to correct the Δ^{14} C of the headspace for mass-dependent fractionation and for incomplete stripping of atmospheric CO₂ during the trapping period using the following equation:

$$X = \frac{\delta^{13}C_{measured} - \delta^{13}C_{soil}}{\delta^{13}C_{atmosphere} - \delta^{13}C_{soil}}$$
(4)

where X is the fraction of remnant atmospheric air in the sample, $\delta^{13}C_{atmosphere}$ is the atmospheric $\delta^{13}C$ signature, and $\delta^{13}C_{soil}$ is the $\delta^{13}C$ signature of soil respiration. $\Delta^{14}C$ is then calculated as

$$\Delta^{14}C_{soil} = \frac{\Delta^{14}C_{measured} - X \times \Delta^{14}C_{atmosphere}}{1 - X}$$
(5)

Chamber B differs from Chamber A by using an infrared gas analyzer in-line with the pump to determine the flux rate into the headspace prior to CO_2 scrubbing. The pump speed was adjusted to match the soil flux rate, thereby maintaining near natural steady-state conditions in the chamber (Schuur and Trumbore 2006). The same equations for X and $\Delta^{14}C_{soil}$ that were used for Chamber A were also used for Chamber B. The modeled volume (11.5 L) and surface area (0.058 m²) of Chamber B were based on the values used in Schuur and Trumbore (2006).

Static Chamber

Static chambers are a type of non-steady-state chamber in which CO_2 from soil is allowed to accumulate without interference in the chamber headspace. In the case of ¹⁴C sampling, after the accumulation period, a molecular sieve trap or sampling flask captures gas from the headspace. For modeling scenarios, the static chamber, Chamber C, is based on the one presented by Hahn et al. (2006). The chamber was deployed for 30 min to allow CO_2 accumulation in the headspace before sampling, thereby ensuring that enough CO_2 is captured in the sampling flask for $\Delta^{14}C$ analysis. For consistency, we also modeled this chamber with a volume of 11.5 L. Isotopic signatures ($\delta^{13}C$ and $\Delta^{14}C$) of flux were calculated using a standard two-source mixing model:

$$\delta^{13}C_{respired} = \frac{\delta^{13}C_{chamber} \times \left[CO_2\right]_{chamber} - \delta^{13}C_{freeair} \times \left[CO_2\right]_{freeair}}{\left[CO_2\right]_{chamber} - \left[CO_2\right]_{freeair}}$$
(6)

where $\delta^{13}C_{respired}$ is the stable isotopic signature of respired CO₂, $\delta^{13}C_{chamber}$ and $[CO_2]_{chamber}$ are the stable isotopic signature and CO₂ concentration of chamber air, respectively, and $\delta^{13}C_{freeair}$ and $[CO_2]_{freeair}$ are the stable isotopic signature and CO₂ concentration of the free air (near-surface atmosphere). A similar equation can be constructed to calculate the chamber estimate respired ¹⁴C activity, which can then be converted to the $\Delta^{14}C$ signature (Hahn et al. 2006).

Isotopic-Forced Diffusion Chamber

The isotopic-forced diffusion (Iso-FD) chamber is similar in design to a dynamic chamber, but rather than mass outflow, the air exchange between the chamber and atmosphere is regulated by a diffusive membrane (Figure 1D). This chamber design has been tested as a tool for sampling $\delta^{13}CO_2$ (Nickerson et al. 2013) and is based on the forced diffusion (FD) bulk CO_2 flux chamber presented by Risk et al. (2011). Here, we evaluated its theoretical performance for $\Delta^{14}C$ sampling (Chamber D). The principle of FD operation is to restrict exchange between the chamber and the atmosphere passively using membranes of known diffusivities. The offset in CO_2 concentration and isotopic abundance between the chamber and the surrounding atmosphere can then be related to soil flux rate and composition. Membranes of particular diffusivities and panel surface areas were chosen for the specific chamber geometry in order to obtain the ideal amount of CO_2 buildup in the chamber for measurements with the smallest error (Creelman et al. 2013). As is the case with the dynamic chamber, FD chamber interior concentrations are an intermediate between the atmosphere and soil, so the technique uses an atmospheric reference chamber, which has no soil inlet, alongside the FD chamber, to correct for changes in the atmospheric reference were coupled to molecular

sieve traps. Δ^{14} C signatures were calculated from the model output for this chamber using the following equation, modified from that presented in Nickerson et al. (2013):

$$\frac{F_{in}^{^{14}C}}{F_{in}^{^{12}C}} = \frac{1}{1.0088} \frac{(C_{FD}^{^{14}C} - C_{atm}^{^{14}C})}{(C_{FD}^{^{12}C} - C_{atm}^{^{12}C})}$$
(7)

where $F_{in}^{^{14}C} / F_{in}^{^{12}C}$ is equivalent to A_s and can be represented in delta notation with Equations 2 and 3, 1.0088 is the diffusive fractionation factor for $^{14}CO_2$ (Wang et al. 1994), and C_{FD} and C_{atm} are the concentrations of each isotopologue present in the Iso-FD chamber and atmospheric reference chamber. Again, for consistency the modeled volume and surface area of the chamber were 11.5 L and 0.058 m², respectively.

Error Analysis

Propagation of uncertainty (error) was calculated for the chambers using the standard partial derivative form (Ku 1966):

$$S_{f} = \sqrt{\left(\frac{\partial f}{\partial x}\right)^{2} S_{x}^{2} + \left(\frac{\partial f}{\partial y}\right)^{2} S_{y}^{2} + \left(\frac{\partial f}{\partial z}\right)^{2} S_{z}^{2} + \dots}$$
(8)

where s_f is the absolute error in the function f, which is composed of the variables x, y, z, and so on, each with variable specific uncertainty s_x, s_y, s_z . For the specific equations used for each chamber, see <u>Appendices A and B</u> in the online Supplementary material.

RESULTS AND DISCUSSION

Dynamic Chambers (A and B)

Simulations of Chamber A, which used a constant pump speed of 0.5 L/min, showed that headspace CO_2 concentrations could differ substantially from the atmosphere unless the soil flux rates were well matched to the CO_2 scrubbing rates. Assuming steady-state concentrations at the soil surface were initially in the range of about 380–1000 ppm CO_2 , using a constant pump speed of 0.5 L/min, the optimal soil flux rate into the chamber should have been 2–6 µmol m⁻² s⁻¹ in order to maintain steady-state concentrations. When soil flux rates fell in this range, our simulation results showed ¹⁴C errors were at their lowest (Figure 2A). This suggested that the main driver of error for Chamber A is the constant pump rate, which is unable to maintain a near steady-state concentration during measurement when fluxes are too low or too high to match the rate of CO_2 removal.

 $\Delta^{14}CO_2$ errors were exacerbated when Chamber A was placed on virtual porous soils, and where flux rates were low (Figure 2A). This was due to a combination of overpumping and lateral diffusion, a problem that is exacerbated in soils of high soil diffusivity. In low-diffusivity soils, however, the chamber performed relatively well because of the limited feedback between the chamber and the soil, which created an effective 1D diffusion pathway eliminating lateral diffusion errors. Results from simulations with varying soil collar depth helped support the interpretation that lateral diffusion contributed to ¹⁴C measurement errors. When the collar depth was increased at fixed levels of diffusivity and production, $\Delta^{14}C$ errors decreased linearly (data not shown).

The apparent ¹⁴C errors associated with Chamber A were also affected by similarity in isotopic values of soil CO₂ production and atmospheric CO₂. This increase in bias arose because of the chamber-based errors in the estimate of X (Equation 4). Any error in the chamber estimate becomes amplified because the two δ^{13} C values are differenced in the denominator of the calculation. As the



-6.5

-7.0

-7.5

-8.0

3 4 5

1 2

0

-10

-20

-30



3

4 5

2

Production (µmol/m²/sec)

-6.5

-7.0

-7.5

-8.0

-6.5

-7.5

-8.0

ź

i.

4 5

3

B -7.0-

 δ^{13} C signature of production became more enriched, overestimates in Δ^{14} C began to appear in lower porosity soils (i.e. soils with higher moisture/bulk density). Similarly, biases also became larger in porous soils with low biological productivity (Figure 2A).

When the specified $\Delta^{14}CO_2$ of production in the model was more enriched, in porous soils with low fluxes, the chamber-measured signatures were more depleted than they should have been. When the signatures of production were more depleted, the chamber yielded more enriched values than it should have (Figure 3A). Near the point where soil production and atmospheric $\Delta^{14}CO_2$ were similar, the error in measurement was minimized. This minimized error was likely an *apparent* minima caused by the lack of distinction between source and ambient $\Delta^{14}CO_2$ signatures.

For Chamber B, in which pump speed was regulated to hold CO_2 at a constant concentration, modeled errors varied with soil porosity and productivity. In soils with high diffusivity, $\Delta^{14}CO_2$ was slightly underestimated, and slightly overestimated in soils with low diffusivities and low biological production rates. The adjustment of the pump speed to the ambient atmospheric concentration led to the maintenance of near steady-state concentrations in the chamber during the measurement period, with some deviation from the true steady-state caused by soil diffusivity (and therefore soil collar length) and the slight stratification of the model atmosphere, causing the slight over- and underestimates.

Similar to the Chamber A results, if the δ^{13} C signature of production and atmosphere were considerably different, Δ^{14} CO₂ deviations from the true isotopic value occurred in soils with low productivity. However, as the δ^{13} C signature of production approached that of the atmosphere (more enriched), deviations similarly became apparent in highly productive soils (Figure 2B). In soils when the Δ^{14} CO₂ of production was depleted relative to the atmosphere, Chamber B gave slight overestimates of Δ^{14} CO₂. If the Δ^{14} CO₂ of production were to be enriched more than the range of our



Figure 3 Contour plots of model output error of Δ^{14} C (‰) of Chamber A (column A) and Chamber B (column B) with a simulated δ^{13} C of production of -30%, collar length of 2 cm, and Δ^{14} C of production of -500%, -200%, 0%, 200%, and 500% in rows 1, 2, 3, 4, and 5 respectively. Soil production (µmol/m²/s) is on the *x* axis and soil diffusivity (logscaled where axis numbers are 10^{y} ; m²/s) on the *y* axis. Note the scale differences between Chamber A and Chamber B.

model simulations, the chamber would produce slight underestimates across all diffusivities, again in soils with lower productivity (Figure 3B).

Error analysis performed on Chamber A and B provided further explanation for deviations in isotopic signature when the prescribed isotopic signature of production was similar to that of the atmosphere. Uncertainty estimates for the dynamic chambers consist of two necessary calculations, the first of which is the uncertainty in X, the fraction of remaining atmospheric air in the chamber. Uncertainty in X is largely driven by the difference between $\delta^{13}C_{atm}$ and $\delta^{13}C_{soil}$ (the denominator in Equation 4), where large differences between the two values minimize the error. Shown in Figure 4a are uncertainty estimates in X (unitless) as a function of $\delta^{13}C_{atm}$ and $\delta^{13}C_{soil}$ for measurement errors of 0.1‰ (open squares) in all of the variables in Equation 4, and 0.3‰ measurement error in $\delta^{13}C_{atmosphere}$ and $\delta^{13}C_{measured}$ and 1.0‰ measurement error in $\delta^{13}C_{soil}$ (circles).

Figure 4b shows the subsequent $\Delta^{14}C_{soil}$ uncertainty estimates as a function of X for measurement errors of 10% in $\Delta^{14}C_{measured}$ and $\Delta^{14}C_{atmosphere}$, and error in X of 0.01, displayed in the open squares. While the value of X drives the largest uncertainty in the estimate of $\Delta^{14}C_{soil}$, the spread in the uncertainty data at a given X value is linearly related to the absolute difference between $\Delta^{14}C_{measured}$ and $\Delta^{14}C_{atmosphere}$. Also shown is the uncertainty for measurement errors of 50% in $\Delta^{14}C_{measured}$ and $\Delta^{14}C_{atmosphere}$, and error in X of 0.1 (gray circles).



Figure 4 Uncertainty estimates for the dynamic chambers. (a) Uncertainty estimates in *X*, the fraction of remaining atmospheric air in the chamber (unitless), as a function of $\delta^{13}C_{atm}$ and $\delta^{13}C_{soil}$ for measurement errors of 0.1% in all of the variables in Equation 4 (open squares), and 0.3% measurement error in $\delta^{13}C_{atmosphere}$ and $\delta^{13}C_{measured}$ and 1.0% measurement error in $\delta^{13}C_{soil}$ (circles). (b) $\Delta^{14}C_{soil}$ uncertainty estimates as a function of *X* for measurement errors of 10% in $\Delta^{14}C_{measured}$ and $\Delta^{14}C_{atmosphere}$, and error in *X* of 0.01 (open squares), and the uncertainty for measurement errors of 50% in $\Delta^{14}C_{measured}$ and $\Delta^{14}C_{atmosphere}$, and error in *X* of 0.1 (gray circles).

Based on the uncertainty analysis, measurements of Δ^{14} C done with a dynamic chamber can be made with much more certainty in scenarios where the atmosphere values of δ^{13} C and Δ^{14} C are different (5‰ in the case δ^{13} C) from the δ^{13} C and Δ^{14} C of the soil.

As demonstrated through the comparison of Chambers A and B in Figures 2 and 3, Chamber B per-

formed consistently better than Chamber A in all simulated scenarios. We attribute this increased performance to maintenance of near steady-state conditions through the adjustment of pump speed. With a constant pump speed, there is an optimal range of flux rates when Chamber A can perform well, but it is less applicable than Chamber B across a large range of soil conditions.

Static Chamber (C)

Previous numerical modeling by Nickerson and Risk (2009b) showed a bias towards underestimation of $\delta^{13}CO_2$ using static chamber designs. The combination of a strong concentration gradient between the soil and chamber, and diffusive fractionation caused ${}^{12}CO_2$ to accumulate in the chamber before ${}^{13}CO_2$, and tended to make the signature of the chamber headspace more negative than the equilibrium condition, as ${}^{13}CO_2$ caught up (see Figure 3 of Nickerson and Risk 2009b). We expected to see similar results when this type of chamber was used to measure ${}^{14}CO_2$, but did not. In fact, the static chamber design (Chamber C) performed very well over the whole range of soil conditions, with the deviation from the true isotopic signature of $\Delta^{14}CO_2$ flux being in the numerical error bounds of the model (~1‰ $\Delta^{14}CO_2$).

Based on subsequent analytical modeling of the chamber isotopic signatures (data not shown), it seems that the marginally increased fractionation factor for ¹⁴CO₂ (1.0044 for ¹³CO₂ and 1.0088 for ¹⁴CO₂) is balanced by a very low ¹⁴CO₂ concentration gradient, leading to a smooth transition from atmospheric to respired isotopic signatures for $\Delta^{14}CO_2$. When the fractionation factor was increased in the analytical model, the nonlinear mixing behavior that was noted in the $\delta^{13}CO_2$ simulations became evident in the $\Delta^{14}CO_2$ results. Similarly, if the fractionation factor was held at 1.0088, but the absolute abundance of ¹⁴CO₂ increased, with ¹²CO₂ and ¹³CO₂ staying the same (and thus the gradients between soil and atmosphere became larger), the nonlinear mixing behavior also became evident.

Figure 5 shows the probable uncertainty for the static chamber. It assumes two error rates, the first with a measurement error of 10% (5% AMS error and 5% sampling and extraction error; Phillips et al. 2013) in Δ^{14} C signatures and 1% in bulk gas concentrations (open squares), and the second with measurement errors of 50% in Δ^{14} C signatures and 5% in bulk gas concentrations (open circles). Most of the uncertainty in the estimation of the ¹⁴C fluxes with this chamber comes from the ratio of the final (C₂) and initial concentrations (C₁). Like the dynamic chambers, measurements made with the static chamber can be done with more certainty when the final chamber measurements are at least twice the value of the initial chamber measurements.



Figure 5 The probable uncertainty for static and Iso-FD chambers assuming two error rates, the first with a measurement error of 10‰ in Δ^{14} C signatures and 1% in bulk gas concentrations (open squares), and the second with measurement errors of 50‰ in Δ^{14} C signatures and 5% in bulk gas concentrations (open circles). C₁ and C₂ are the initial and final concentrations for the static chamber, or for the Iso-FD chamber, they are equivalent to the atmospheric chamber and soil chamber measurements.

Isotopic-Forced Diffusion Chamber (D)

In all simulations with the Iso-FD chamber, the predicted isotopic signature of flux was very near the prescribed value. Similar to the static chamber results, the Iso-FD results were within the numerical error bounds of the model (~1‰ $\Delta^{14}CO_2$). We attribute the accuracy of this chamber to the fact that a diffusive steady-state was maintained during the measurement period, avoiding isotopic disequilibrium created by other chamber designs. Simulations showed that the $\Delta^{14}CO_2$ behavior for this chamber is similar to that described in Nickerson et al. (2013) for $\delta^{13}CO_2$, where a slight buildup of concentration in the chamber headspace produced concentration and isotopic plumes directly below the chamber. However, despite these plumes, sampling errors were negligible.

Assuming the fractionation factor for the Iso-FD calculations is constant and known, the uncertainty for the Iso-FD chamber takes the same form as the static chamber and uses the same calculation variables (Figure 5). For the Iso-FD chamber, most of the uncertainty in the estimation of the ¹⁴C fluxes comes from the ratio of concentration measurements in soil chamber and atmospheric chamber measurements, which can be thought of as equivalent to the ratio of the final (C₂) and initial concentrations (C₁) for the static chamber. As was the case with the dynamic chambers and static chamber, measurements made with Iso-FD chambers have more certainty when the soil chamber concentration measurements are at least twice the value of the atmospheric chamber measurements.

Other Considerations

There are also some other possible biases that should be considered, which are not included in the model. The simulated soil in the model is at steady-state. It has a constant biological production, soil diffusivity, and isotopic signature of production through depth, so no natural non-steady-state (NSS) effects (Nickerson and Risk 2009a) are included in the results. A nonuniform soil with varying diffusivities and production rates through depth would provide different model results than the uniform modeled soil (Venterea and Baker 2008). Nickerson and Risk (2009a) and Moyes et al. (2010) demonstrated the effects of dynamic fractionations, where soil features and processes such as biological production, diffusivity, pore space, and atmospheric concentrations, which have temporal variation, will induce NSS transport conditions that lead to transient changes in the isotopic composition of the soil CO₂ flux. The main driving force behind this is the difference in the diffusion rate between CO₂ isotopologues, which will be slightly amplified when considering ¹⁴CO₂ due to its increased mass. The measured values could therefore be further biased on top of the potential biases induced by the chamber method. Despite the counterbalance of low-¹⁴C concentration gradients shown in the case of the static chamber, these dynamic fractionation effects should be investigated further in order to ensure that they are not causing bias.

The model also assumes that the method used to sample from the chamber, molecular sieve trap, or sampling flask is completely efficient and causes no fractionations. Δ^{14} C static chamber methods can include a capillary tube that attaches a sampling flask to the chamber (Hahn et al. 2006). This method could cause a potential fractionation, where the lighter isotopologue, 12 CO₂, would travel faster than 14 CO₂ through the capillary tube, so the resulting mixture in the sampling flask could potentially be more depleted than the mixture in the chamber headspace. It would be ideal to obtain a quick sample from the chamber, for example, by attaching the sampling flask under vacuum to the chamber and drawing 1 L of sample immediately. This configuration is similar to how the model simulates sampling from the chamber and would work well for the two-point mixing model used for static chambers (Hahn et al. 2006). Another issue not addressed in the model surrounding the static chamber and Iso-FD chamber methods is the possible stratification in the chamber headspace because of the lack of mixing. In the case of the static chamber, concentration stratification com-

bined with capillary tube errors could lead to an even greater bias in the captured gas in the sampling flask, where the gas traveling along the tube is not well mixed. For the Iso-FD chamber, there could also be problems with using molecular sieve traps, because as CO_2 is removed from the chamber headspace, it decreases the concentration in the chamber, causing the chamber to no longer be at steady-state. In the case of dynamic chambers, if the soda lime trap and molecular sieve trap have different trapping efficiencies, then the expected advantage of having a truer steady-state will not be met. If the soda lime and molecular sieve traps also fractionate ¹⁴C, then the already biased results found in this study will be further biased.

Although the Iso-FD and static chambers perform well under all simulated conditions in the numerical model, there are some other things that need to be considered. The two samples needed for these chambers (soil or final chamber measurement, and atmosphere or initial chamber measurement) have to be different enough from one another, to keep the error rate within an acceptable range (Figure 6). For the Iso-FD chamber, this means choosing an appropriate membrane, and for the static chamber, this means leaving the chamber deployed for a long enough period of time. Therefore, in scenarios where the soil chamber or final concentrations and the atmospheric chamber or initial chamber concentrations are quite similar, these chambers would not be ideal.

14 C/12 C



Figure 6 (a) Time series of chamber equilibration for stable carbon and ¹⁴C ratios. Note that the ¹³C/¹²C ratio overshoots the equilibrium value between 2 and 10 hr due to lateral diffusion. This behavior is not present in the ¹⁴C ratios except for when the ¹⁴C signature of production and the ¹⁴C signature of the atmosphere (50‰ for this model simulation) are equal. The insert is a *y*-axis zoom (*y*-axis values are 10^{-12}) of the $\Delta^{14}C$ 50‰ line to display the overshooting behavior. (b) ¹⁴C and stable carbon ratio mixing lines during chamber equilibration for three $\Delta^{14}C$ signatures. A hook in the line signifies a different mixing behavior for ¹³C and ¹⁴C, whereas a straight line means the mixing behavior is the same for both isotopes.

The δ^{13} C correction used to account for mass-dependent fractionations in Δ^{14} C assumes that 14 CO₂ and 13 CO₂ diffusive fractionation are a constant multiple of one another. Figure 6 demonstrates that this assumption is incorrect. In a time series view of chamber equilibration (Figure 6a), 14 C/ 12 C does not behave in the same way as 13 C/ 12 C (lateral diffusion), except for when the 14 C signatures of production and atmosphere are equal (50‰ in this case). In Figure 6b, chamber equilibration mixing

lines of ¹⁴C/¹²C and ¹³C/¹²C demonstrate whether a difference in mixing between the stable and radioactive isotopes exists. A straight line shows that as ¹³C increased in the chamber, ¹⁴C increased in a constant proportion. A curvilinear line or hook, in contrast, shows that ¹⁴C increased at a different rate than ¹³C. Despite the differences in mixing behaviors between the isotopes (¹⁴C fractionation is not a constant multiple of ¹³C fractionation), the changes to the fractionation factor between the two isotopes that arises from lateral diffusion biases is not sufficient to have an impact on the ¹⁴C result. In other words, the nonlinear behavior during chamber CO₂ accumulation is minor enough that assuming a fractionation for ¹⁴C of twice ¹³C fractionation produces errors far smaller than AMS error.

CONCLUSIONS

These model simulations suggested more favorable performance of soil surface sampling chambers than we expected. The static chamber had little sampling error in these simulations, whereas we expected it to have a greater bias than dynamic chambers, like in the case of δ^{13} C (Nickerson and Risk 2009b). Based on the simulation results, the static chamber and Iso-FD chamber performed the best under the range of soil conditions simulated. Other considerations should be made, however, when choosing a sampling method, including the choice between molecular sieve traps or sampling flasks, the cost, and the length of time needed to sample the chamber. This modeling exercise also showed that the assumption that stable and ¹⁴C isotopic diffusion fractionations are a constant multiple of one another through time is not universally true, especially under non-steady-state conditions. The δ^{13} C correction still stands because the changes to the diffusive fractionation during equilibration are not large enough to impact Δ^{14} C, but researchers should still be cautious, and this should be investigated further for non-steady-state soil conditions.

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