

Modeling soil CO₂ production and transport with dynamic source and diffusion terms: testing the steady-state assumption using DETECT v1.0

Edmund M. Ryan^{1,2}, Kiona Ogle^{2,3,4,5}, Heather Kropp⁶, Kimberly E. Samuels-Crow³, Yolima Carrillo⁷, and Elise Pendall⁷

¹Lancaster Environment Centre, Lancaster University, Lancaster, UK

²School of Life Sciences, Arizona State University, Tempe, Arizona, USA

³School of Informatics, Computing, and Cyber Systems, Northern Arizona University, Flagstaff, Arizona, USA

⁴Center for Ecosystem Science and Society, Northern Arizona University, Flagstaff, Arizona, USA

⁵Department of Biological Sciences, Northern Arizona University, Flagstaff, Arizona, USA

⁶Department of Geography, Colgate University, Hamilton, New York, USA

⁷Hawkesbury Institute for the Environment, Western Sydney University, NSW, Australia

Correspondence: Edmund M. Ryan (edmund.ryan@lancaster.ac.uk)

Received: 7 September 2017 - Discussion started: 10 October 2017

Revised: 15 February 2018 - Accepted: 21 February 2018 - Published: 28 May 2018

Abstract. The flux of CO_2 from the soil to the atmosphere (soil respiration, R_{soil}) is a major component of the global carbon (C) cycle. Methods to measure and model R_{soil}, or partition it into different components, often rely on the assumption that soil CO₂ concentrations and fluxes are in steady state, implying that R_{soil} is equal to the rate at which CO₂ is produced by soil microbial and root respiration. Recent research, however, questions the validity of this assumption. Thus, the aim of this work was two-fold: (1) to describe a non-steady state (NSS) soil CO₂ transport and production model, DETECT, and (2) to use this model to evaluate the environmental conditions under which R_{soil} and CO_2 production are likely in NSS. The backbone of DETECT is a non-homogeneous, partial differential equation (PDE) that describes production and transport of soil CO₂, which we solve numerically at fine spatial and temporal resolution (e.g., 0.01 m increments down to 1 m, every 6 h). Production of soil CO₂ is simulated for every depth and time increment as the sum of root respiration and microbial decomposition of soil organic matter. Both of these factors can be driven by current and antecedent soil water content and temperature, which can also vary by time and depth. We also analytically solved the ordinary differential equation (ODE) corresponding to the steady-state (SS) solution to the PDE model. We applied the DETECT NSS and SS models to the six-month growing season period representative of a native grassland in Wyoming. Simulation experiments were conducted with both model versions to evaluate factors that could affect departure from SS, such as (1) varying soil texture; (2) shifting the timing or frequency of precipitation; and (3) with and without the environmental antecedent drivers. For a coarsetextured soil, R_{soil} from the SS model closely matched that of the NSS model. However, in a fine-textured (clay) soil, growing season R_{soil} was ~ 3 % higher under the assumption of NSS (versus SS). These differences were exaggerated in clay soil at daily time scales whereby R_{soil} under the SS assumption deviated from NSS by up to 35 % on average in the 10 days following a major precipitation event. Incorporation of antecedent drivers increased the magnitude of R_{soil} by 15 to 37 % for coarse- and fine-textured soils, respectively. However, the responses of R_{soil} to the timing of precipitation and antecedent drivers did not differ between SS and NSS assumptions. In summary, the assumption of SS conditions can be violated depending on soil type and soil moisture status, as affected by precipitation inputs. The DETECT model provides a framework for accommodating NSS conditions to better predict R_{soil} and associated soil carbon cycling processes.

1 Introduction

The flux of CO₂ to the atmosphere from the soil (i.e., soil respiration, R_{soil}) is one of the largest fluxes in the global carbon (C) cycle, and when aggregated globally over an entire year it is approximately 10 times the annual amount of CO₂ emitted by fossil fuel burning (Friedlingstein et al., 2014; Hashimoto et al., 2015). Moreover, global change experiments and predictions from models agree that R_{soil} is expected to increase in a future climate of elevated CO₂ and warming (Cox, 2001; Davidson and Janssens, 2006; Piao et al., 2009; Pendall et al., 2013; Ryan et al., 2015). Therefore, monitoring R_{soil} is important for quantifying and modeling the global C cycle.

Commonly, R_{soil} is monitored by directly measuring surface soil CO₂ fluxes using various chamber methods (Luo and Zhou, 2010; Risk et al., 2011) or by estimating R_{soil} from soil CO₂ concentrations measured at multiple depths using probe methods (Pendall et al., 2003; Tang et al., 2003; Vargas et al., 2010). The probe methods employ diffusion equations that often rely on the assumption that R_{soil} at the surface is in steady state (SS) with subsurface CO₂ production by roots and micro-organisms (Tang et al., 2003; Lee et al., 2004; Baldocchi et al., 2006; Luo and Zhou, 2010; Vargas et al., 2010; Šimůnek et al., 2012). That is, the SS assumption essentially presumes that CO₂ produced by roots and microbes within the soil profile is instantaneously respired from the soil surface, effectively neglecting delays due to CO₂ transport times. Partitioning R_{soil} (surface flux) into its different components (e.g., sub-surface heterotrophic [microbes] versus autotrophic [root or rhizosphere] respiration) using isotope methods (Hui and Luo, 2004; Ogle and Pendall, 2015), trenching methods (Šimůnek and Suarez, 1993), or soil CO₂ models (Vargas et al., 2010) also relies on the SS assumption. Even simulations of the vertical movement of soil CO₂ through snow have employed a SS diffusion model (Monson et al., 2006). Recent work, however, calls into question whether this SS assumption is valid most of the time or in most systems (Maggi and Riley, 2009; Nickerson and Risk, 2009).

Given the use of the SS assumption in a diverse range of settings, the aim of this study was to determine the meteorological and site specific conditions under which the SS assumption is valid, and the circumstances under which a nonsteady state (NSS) model substantially improves our understanding of subsurface processes that lead to observed R_{soil} . We focused on soil texture because it is a critical factor underlying soil porosity and tortuosity, which, in turn, control soil CO₂ diffusion rates (Bouma and Bryla, 2000). For example, coarse-grained (e.g., high sand content) soils generally facilitate fast CO₂ diffusion rates, especially under low soil moisture conditions associated with high air-filled porosity (Bouma and Bryla, 2000); the opposite is expected for finer-grained (e.g., silt or clay) soils. Thus, we expect coarsegrained soils to generally induce SS conditions for soil CO₂, whereas fine-grained soils would likely produce frequent and longer duration NSS conditions, especially following rain pulses that decrease air-filled pore space, thereby reducing CO_2 diffusivity.

We also focused on the impacts of precipitation variability given that the timing and magnitude of precipitation pulses can have large effects on R_{soil} (Huxman et al., 2004; Schwinning et al., 2004; Sponseller, 2007; Cable et al., 2008; Borken and Matzner, 2009; Ogle et al., 2015). Precipitation indirectly impacts R_{soil} via its influence on soil moisture dynamics, and soil moisture and soil texture affect both diffusivity (a physical process) and CO₂ production (a primarily biological process governed by roots and microbes). For example, as precipitation pulses infiltrate the soil, the CO₂ in the pore spaces gets displaced with water, which may be seen as a transient spike in R_{soil} (e.g., Lee et al., 2004). Such transient spikes, however, may also be attributable to changes in decomposition, microbial growth, and/or C substrate availability in response to wetting (Birch, 1958; Borken et al., 2003; Jarvis et al., 2007; Xiang et al., 2008; Meisner et al., 2013). This transient response may be followed by a depression in $R_{\rm soil}$ since water-filled pores will ultimately slow CO₂ diffusion and transport (Bouma and Bryla, 2000). These linked effects imply that precipitation pulses and their effects on soil moisture are likely to impose NSS soil CO₂ conditions, but the manner in which such pulses impact these processes is temporally dynamic and spatially complex, and therefore difficult to measure directly.

We evaluated the importance of soil texture and precipitation variability on SS versus NSS soil CO₂ behavior via a simulation-based approach. To allow for the possibility of both SS and NSS behavior, we implemented a depth- and time-varying CO₂ transport and production model that built on the groundbreaking work of Fang and Moncrieff (1999), Hui and Luo (2004), Nickerson and Risk (2009), Moyes et al. (2010) and Risk et al. (2012). These processes are captured by a partial differential equation (PDE) model that is grounded in diffusion theory, and solved numerically. Some current NSS models make simplifying assumptions such as assuming depth-invariant CO₂ production rates (e.g., Fang and Moncrieff, 1999), or assuming that production only responds to concurrent environmental conditions (e.g., Nickerson and Risk, 2009). Such simplifications may make it difficult to evaluate physical and biological conditions leading to SS versus NSS behavior.

We addressed the aforementioned shortcomings of existing NSS models with the DETECT (DEconvolution of Temporally varying Ecosystem Carbon componenTs) model, version 1.0 (v1.0), which implemented four improvements. First, we simulated soil CO₂ at 100 different 0.01 m depth increments to ensure numerical accuracy of the solutions (Haberman, 1998). Second, we estimated the soil water content and soil temperature data for all depths and all time points using a separate model (HYDRUS; Šimůnek et al., 2005, 2008). Third, we simulated the production of CO₂ by microbial and root respiration at each depth by linking these processes to existing respiration models that are typically applied to "bulk" soil (Lloyd and Taylor, 1994; Cable et al., 2008; Davidson et al., 2012; Todd-Brown et al., 2012). Fourth, we included antecedent (past) environmental and meteorological conditions as part of the functions that predict soil CO_2 production, due to their importance for predicting soil and ecosystem CO_2 fluxes (Cable et al., 2013; Barron-Gafford et al., 2014; Ryan et al., 2015). For example, soil respiration following a rain event is generally greater if the rain event follows a dry period versus a wet period (Xu et al., 2004; Sponseller, 2007; Cable et al., 2008, 2013; Thomas et al., 2008). Such antecedent effects may underlie the importance of biological versus physical processes in governing the transition between SS and NSS behavior.

After describing the DETECT model, we subsequently use it to explore the effects of soil texture, precipitation pulses, and antecedent conditions on the relative importance of NSS soil CO_2 behavior and to identify the factors giving rise to such behavior. We simulated soil CO_2 concentrations, CO_2 production, and R_{soil} under four different soil textures and three different precipitation regimes. For each scenario, we implemented the DETECT model under the assumption that soil CO₂ production is affected by antecedent moisture and temperature versus the assumption that only concurrent conditions matter. Data from the Wyoming Prairie Heating and CO₂ Enrichment (PHACE) experiment (e.g., Pendall et al., 2013; Carrillo et al., 2014a; Ryan et al., 2015; Zelikova et al., 2015; Mueller et al., 2016) were used to parameterize the model and motivated the selection of the texture and precipitation scenarios. Under the different scenarios, we compared $R_{\rm soil}$ predicted from the DETECT model to that of a simpler SS model, and evaluated the relative impact of SS assumptions on inferring subsurface processes (e.g., CO₂ production by roots and microbes) and surface CO_2 fluxes (i.e., R_{soil}).

2 Methods

2.1 Description of the non steady state DETECT model

The PDE that underlies the DETECT model (v1.0) accounts for time- and depth-varying CO₂ diffusivity and CO₂ production by root and microbial respiration (Fang and Moncrieff, 1999). We use a pair of PDEs, one describing the soil CO₂ derived from root respiration (subscripted with *R*), and the other for CO₂ derived from microbial respiration (*M*) such that for K = R or *M*:

$$\frac{\partial c_K(z,t)}{\partial t} = \frac{\partial}{\partial z} \left(D_{\rm gs}(z,t) \frac{\partial c_K(z,t)}{\partial z} \right) + S_K(z,t). \tag{1}$$

 $c_K(z, t)$ is CO₂ concentration (mg CO₂ m⁻³), $D_{gs}(z, t)$ is the effective diffusivity of CO₂ through the soil (m² s⁻¹), and $S_K(z, t)$ is the source (or production) term (mg CO₂ m⁻³) (Fig. 1b), all of which vary by depth *z* (meters) and time *t* (hours). Note that D_{gs} is assumed to be the same for root-

and microbial-derived CO₂ and is thus not indexed by *K*. In this version of the model, we assumed that CO₂ transport within the soil profile and over time is solely governed by gaseous diffusion, and we ignored other types of CO₂ transport – such as diffusion in the liquid state, convection, and bulk transport via the vertical movement of water – that have been shown to have a negligible contribution (Fang and Moncrieff, 1999; Kayler et al., 2010). Total soil CO₂ and total CO₂ production are given as $c(z,t) = c_M(z,t) + c_R(z,t)$ and $S(z,t) = S_M(z,t) + S_R(z,t)$, respectively. Below we describe the two main components of the PDE model: (1) CO₂ diffusivity, D_{gs} , and (2) the production terms, $S_R(z,t)$ and $S_M(z,t)$. Finally, we note that Eq. (1) is the mass balance equation (see Sect. S3 in the Supplement for more information).

2.1.1 Soil CO₂ diffusivity submodel

The diffusivity of CO₂ within the soil (D_{gs}) depends on the soil structure and water content; we modeled D_{gs} using the Moldrup function (Sala et al., 1992; Moldrup et al., 2004). We chose this formulation because it is more accurate than other common models, such as the Millington and Quirk (1961) and Penman (1940) models (Moldrup et al., 2004). Based on Moldrup et al. (2004), D_{gs} (m² s⁻¹) is defined as

$$D_{gs}(z,t) = D_{g0}(z,t) \cdot \left(2\phi_{g100}(z)^3 + 0.04\phi_{g100}(z)\right) \\ \cdot \left(\frac{\phi_g(z,t)}{\phi_{g100}(z)}\right)^{2+\frac{3}{b(z)}},$$
(2)

where $D_{g0}(z,t) = D_{stp} \cdot \left(\frac{T_{S}(z,t)}{T_0}\right)^{1.75} \cdot \left(\frac{P_0}{P(t)}\right)$ and $D_{stp} =$ $1.39 \times 10^{-5} \text{ m}^2 \text{ s}^{-1}$ is the diffusion coefficient for CO₂ in air at standard temperature $(T_0, 273 \text{ K})$ and pressure (P_0, P_0) 101.325 kPa); $T_S(z,t)$ is the soil temperature (Kelvin) at depth z and time t, and P(t) is the air pressure (kPa) just above the soil surface at time t. The remaining terms in Eq. (2) include $\varphi_g(z,t)$, the air-filled soil porosity, which is related to the total soil porosity ($\varphi_{\rm T}$) and volumetric soil water content (θ) according to $\varphi_g(z,t) = \varphi_T(z) - \theta(z,t)$, and $\varphi_{\rm T}(z)$ is defined as $1 - {\rm BD}(z)/{\rm PD}$, where BD and PD are the bulk density and particle density of the soil, respectively (Davidson et al., 2006); $\varphi_{g100}(z)$ is the air-filled porosity at a soil water potential (Ψ) of $-100 \,\mathrm{cm} \,\mathrm{H_2O}$ (about $-10 \,\mathrm{kPa}$); b(z) is a unitless parameter that is related to the pore size distribution of the soil based on the water retention curve given by $\Psi = \Psi_{\rm e}(\theta/\theta_{\rm sat})^{-b}$, where $\Psi_{\rm e}(z)$ is the air-entry potential - calculated from measurements (Morgan et al., 2011) - and $\theta_{\rm sat}(z)$ is the saturated soil water content (v/v).

2.1.2 CO₂ source (production) terms

Soil CO_2 can be produced in the soil (*S* term in Eq. 1) by five different biological processes: (i) root growth respiration, (ii) root maintenance respiration, (iii) consumption of

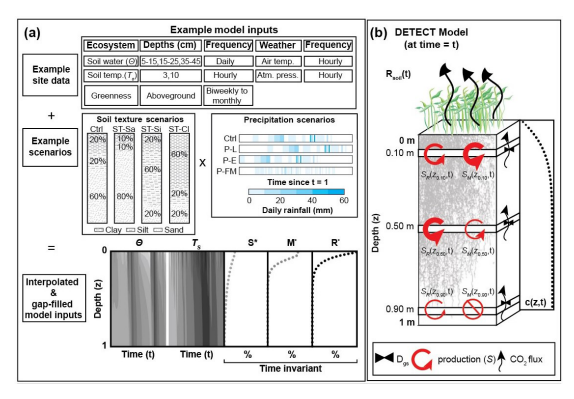


Figure 1. Graphical representation of (a) the required inputs to the DETECT model and the associated scenarios implemented in this study; and (b) the components of the DETECT model at a particular time t, indicating depth-dependent production, CO₂ concentrations, and CO₂ fluxes.

rhizodeposits by root-associated microorganisms and associated microbial respiration, (iv) microbial decomposition of newly produced plant litter that has been incorporated into the soil matrix, and (v) microbial decomposition of older soil organic matter (SOM) (Pendall et al., 2004). Due to the general lack of sufficient data and process understanding to accurately separate all five sources, the DETECT model treats CO_2 production as the sum of two main contributions: CO_2 respired by (1) roots and closely associated microorganisms (the sum of i–iii), giving $S_R(z, t)$, and (2) free-living soil microorganisms (the sum of iv–v), giving $S_{\rm M}(z,t)$. Such simplification based on root and microbial sources is common in models of soil CO₂ transport and production (Šimůnek and Suarez, 1993; Fang and Moncrieff, 1999; Hui and Luo, 2004). Although DETECT v1.0 assumes that root and microbial respiration are independent of one another, they both depend on the same environmental data (e.g., θ and T_S).

CO₂ production by root respiration is represented as the product of three terms: (i) the mass-specific base respiration rate (R_{Rbase}) at a reference soil temperature of $T_S = T_{\text{ref}}$ and at average soil water and antecedent conditions, (ii) root mass expressed as the amount of root carbon, $C_R(z, t)$, and (iii) functions that rescale R_{Rbase} to account for the effect of soil water (θ), temperature (T_S), and their antecedent counterparts. Antecedent temperature is denoted by T_S^{ant} , and for roots antecedent soil water is given by θ_R^{ant} . In general,

 $S_R(z,t)$ is given by

$$S_R(z,t) = R_{\text{Rbase}} \cdot C_R(z,t) \cdot f\left(\theta\left(z,t\right), \theta_R^{\text{ann}}(z,t)\right)$$
$$\cdot g\left(T_S(z,t), T_S^{\text{ann}}(z,t)\right).$$
(3)

The functional form of $C_R(z, t)$ is informed by field data on root biomass C (see Sect. S1 for complete details). The functions f and g are given by

$$f\left(\theta, \theta_{R}^{\text{ant}}\right) = \exp\left(\alpha_{1}\theta(z, t) + \alpha_{2}\theta_{R}^{\text{ant}}(z, t) + \alpha_{3}\theta(z, t) \cdot \theta_{R}^{\text{ant}}(z, t)\right)$$
(4a)

$$g\left(T_{S}, T_{R}^{\text{ant}}\right) = \exp\left(E_{o}(z, t)\right)$$
$$\cdot \left(\frac{1}{T_{\text{ref}} - T_{o}} - \frac{1}{T_{S}(z, t) - T_{o}}\right)$$
(4b)

$$E_o(z,t) = E_o^* + \alpha_4 T_S^{\text{ant}}(z,t).$$
(4c)

 $R_{\text{Rbase}}, \alpha_1, \alpha_2, \alpha_3, \alpha_4, T_o$, and E_o^* are parameters that require numerical values (Table 1; Ryan et al., 2015), θ and T_S are informed by field data, and θ_R^{ant} and T_S^{ant} are computed from the field data (described below). The temperature scaling function, g (Eq. 4b) was motivated by Lloyd and Taylor (1994) and has been successfully used to describe soil and ecosystem respiration (Luo and Zhou, 2010; Cable et al., 2013; Ryan et al., 2015). $E_o(z, t)$ is analogous to an energy of activation term that governs the apparent temperature sensitivity of S_R (Davidson and Janssens, 2006; Cable et al., 2011;

Tucker et al., 2013); we assume E_o responds to antecedent temperature, reflecting a potential thermal acclimation response (Atkin and Tjoelker, 2003; Ryan et al., 2015). T_o is also related to the apparent temperature sensitivity (Cable et al., 2011), and we assume that it is invariant with depth and time (Lloyd and Taylor, 1994; Cable et al., 2013; Barron-Gafford et al., 2014; Ryan et al., 2015). While the functional forms and choice of environmental drivers used for f and g were motivated by previous analyses (Cable et al., 2013; Barron-Gafford et al., 2014), the exact functions and parameter values were based on Ryan et al. (2015) and Cable et al. (2013). Exponential functions are also used for the moisture (f) and temperature (g) scale functions to ensure f > 0and g > 0 (Eq. 4a). The choice of an exponential form of the functions was based on Ryan et al. (2015), with graphical forms of the total CO₂ production based on these functions given in Fig. S10 (Supplement). However, the DETECT model is flexible enough to accommodate alternative functions for f and g. For example, we ran DETECT for the control scenario using a bell-shaped function that described how soil CO₂ production changes with θ (Sect. S4 and Fig. S8, Supplement) as an alternative to Eq. (4a). For this alternative model run, the modeled R_{soil} was very similar to the modeled R_{soil} from the results of this study (Fig. S9, Supplement).

 CO_2 production by microbial respiration and SOM decomposition is represented by a modified version of the Dual Arrhenius and Michaelis–Menten (DAMM) model (Davidson et al., 2012). We exclude the O_2 term, rendering the model relevant to systems that are typically unlimited by O_2 availability, such as the semi-arid site that we focus on, but we accounted for a microbial C pool (C_{MIC}) and a soluble soil-C pool (C_{SOL}) (Todd-Brown et al., 2012) such that

$$S_{\rm M}(z,t) = V_{\rm max}(z,t) \cdot \frac{C_{\rm SOL}(z,t)}{K_{\rm m} + C_{\rm SOL}(z,t)}$$
$$\cdot C_{\rm MIC}(z,t) \cdot (1 - {\rm CUE}). \tag{5}$$

Decomposition is assumed to be an enzymatic process that follows Michaelis–Menten kinetics, where V_{max} is the maximum potential decomposition rate, and K_{m} (the half-saturation constant) is the amount of substrate required for the decomposition rate to reach half of V_{max} . Carbon-use efficiency (CUE) represents the proportion of total C assimilated by microbes that is allocated for microbial growth (Tucker et al., 2013). We excluded a microbial death rate term (Todd-Brown et al., 2012) because we had insufficient data on death rates, and C_{MIC} is only ~ 1% of C_{SOL} at our study site (Carrillo and Pendall, 2018).

In contrast to the original DAMM formulation, we allowed $S_{\rm M}(z,t)$ and $V_{\rm max}(z,t)$ to vary by depth and time, whereas existing applications of the DAMM model are generally applied to "bulk" soil (i.e., do not vary with z). We also modeled $V_{\rm max}$ according to the modified energy of activation function described in Lloyd and Taylor (1994), which essen-

tially parallels Eq. (4b)-(4c):

$$V_{\max}(z,t) = V_{\text{Base}} \cdot f\left(\theta, \theta_M^{\text{ant}}\right) \cdot \exp\left(E_o(z,t)\right)$$
$$\cdot \left(\frac{1}{T_{\text{ref}} - T_o} - \frac{1}{T_S(z,t) - T_o}\right). \tag{6}$$

 V_{Base} is the "base" V_{max} at a reference soil temperature of T_{ref} and at mean values of current θ and antecedent θ and T_S (i.e., mean values of θ_M^{ant} and T_S^{ant}). $E_o(z, t)$ and $f(\theta, \theta_m^{\text{ant}})$ follow the same functional forms and interpretation as described for the root respiration submodel (Eqs. 3 and 4a–c), except that θ_M^{ant} is used instead of θ_R^{ant} , respectively, and different values are specified for the parameters $\alpha_1, \alpha_2, \alpha_3, \alpha_4, T_o$, and E_o^* to reflect microbial respiration The values are given in Table 1, and Sect. 2.4.5 explains how the values were estimated.

Finally, C_{SOL} is modeled as a function of soil organic C content at depth *z*, $C_{SOM}(z)$ based on the fraction, *p*, of $C_{SOM}(z)$ that is soluble and the diffusivity of the substrate in liquid, D_{liq} (Davidson et al., 2012). The equation for C_{SOL} is given by

$$C_{SOL}(z,t) = C_{SOM}(z) \cdot p \cdot \theta(z,t)^3 \cdot D_{liq}.$$
(7)

The values of *p* and D_{liq} were taken from laboratory analysis (see Sect. 2.4.5) and Davidson et al. (2012), respectively. We assumed that $C_{SOM}(z)$ and $C_{MIC}(z)$ (see Eq. 5) are constant over time given the relatively short simulation periods explored here (a single growing season); but the model could easily be modified to allow for time-varying C_{SOM} and C_{MIC} . Here, $C_{SOM}(z)$ and $C_{MIC}(z)$ are simple, empirical functions that were informed by data (see Sect. S1 for details). Moreover, while assumption of time invariant $C_{SOM}(z)$ and $C_{MIC}(z)$ is an implicit SS assumption about biological factors affecting soil CO₂ dynamics, this assumption allows us to isolate the importance of NSS conditions that are primarily due to physical CO₂ transport characteristics.

2.1.3 Soil respiration

The efflux of CO₂ from the soil surface (soil respiration, R_{soil}) is computed as

$$R_{\text{soil}}(t) = \frac{D_{\text{gs}}(z=0.01,t)}{\Delta z} \left(c(z=0.01,t) - c_{\text{atm}}(t) \right).$$
(8)

 $D_{\rm gs}(z=0.01,t)$ is the diffusivity of CO₂ in the soil and c(z=0.01,t) is the total CO₂ concentration (microbial- and root-derived), respectively, at z=0.01 m depth and time t; $c_{\rm atm}(t)$ is the CO₂ concentration in the atmosphere above the soil surface; and Δz is the depth increment that the model solves for soil CO₂ concentration (here, $\Delta z = 0.01$ m).

2.2 Numerical implementation of the DETECT model

The numerical solution to the NSS version of the DETECT model v1.0, as described in Eqs. (1)–(8), requires an initial

Table 1. Summary of scalar parameters used in the non-steady-state (DETECT) model, arranged into four groups: parameters unique to the microbial respiration submodel for $S_M(z, t)$ (group 1); parameters unique to the root respiration submodel for $S_R(z, t)$ (group 2); parameters that are shared between the $S_M(z, t)$ and $S_R(z, t)$ submodels (group 3); parameters used to calculate soil CO₂ diffusivity, D_{gs} (group 4). See Sect. 2.4.5 for details about how the parameters were estimated.

Symbol	Description	Value	Units	Eq(s).
Group 1 -	- root submodel parameters			
R*	Total root biomass C in a 1 m deep by 1 cm ² soil column	111.5	$\mathrm{mg}\mathrm{C}\mathrm{cm}^{-2}$	3
<i>R</i> _{RBase}	Root mass-base respiration rate at 10 °C and mean environmental conditions	6×10^{-5}	${ m mg}{ m C}{ m cm}^{-3}{ m h}^{-1}$	3
$\alpha_{1(R)}$	The effect of soil water content (θ) on root respiration	11.65	unitless	3, 4a
$\alpha_{2(R)}$	The effect of antecedent θ (θ_R^{ant}) on root respiration	20.7	unitless	3, 4b
$\alpha_{3(R)}$	The interactive effect of θ and θ_R^{ant} on root respiration	-164.2	unitless	3, 4c
Group 2 -	- microbial submodel parameters			
S*	Total soil organic C in a 1 m deep by 1 cm ² soil column	711.6	${ m mg}{ m C}{ m cm}^{-2}$	5
M*	Total microbial biomass C in a 1 m deep by 1 cm ² column of soil	12.3	$mgCcm^{-2}$	5
V _{Base}	Value of V_{max} at 10 °C and mean environmental conditions	0.0015	$mgCcm^{-3}h^{-1}$	5,6
$\alpha_{1(M)}$	The effect of θ on microbial respiration	14.05	unitless	5,6
$\alpha_{2(M)}$	The effect of antecedent θ (θ_M^{ant}) on microbial respiration	11.05	unitless	5,6
$\alpha_{3(M)}$	The interactive effect of θ and θ_M^{ant} on microbial respiration	-87.6	unitless	5,6
Km	Michaelis–Menten half saturation constant	10^{-5}	${ m mg}{ m C}{ m cm}^{-3}{ m h}^{-1}$	5
CUE	Microbial carbon-use efficiency	0.8	${ m mg}{ m C}{ m mg}^{-1}{ m C}^{-1}$	5
р	Fraction of soil organic C that is soluble	0.004	_	7
D _{liq}	Diffusivity of soil C substrate in liquid	3.17	unitless	7
Group 3 -	- shared parameters between root/microbial submodels			
$E_{o}*$	Temperature sensitivity parameter, somewhat analogous to an energy of activation	324.6	Kelvin	4c
T_o	Temperature sensitivity-related parameter	227.5	Kelvin	4c
α_4	The effect of antecedent soil temperature (T_S^{ant}) on root and microbial respiration	-4.7	unitless	4c
Group 4 -	- soil CO ₂ diffusivity submodel parameters			
$\alpha_{3(R)}$	Absolute value of the slope of the line relating $log(\Psi)$ versus $log(\theta)$	4.547	unitless	2
BD	Soil bulk density	1.12	$\rm gcm^{-3}$	2
φ_{g100} PD	Air-filled porosity at soil water potential of -100 cm H_20 ($\sim 10 \text{ kPa}$) Particle density	18.16	%	2

(9a)

(9b)

condition (IC) and two boundary conditions (BCs), which we specified as

 $c(z, t = 0) = c_0(z)$

Upper BC : $c(z = 0, t) = c_{atm}(t)$

Lower BC:
$$\frac{\partial c(z=1,t)}{\partial z} = 0.$$
 (9c)

The function $c_0(z)$ is determined and parameterized in two stages: (1) observed soil CO₂ concentration data at three depths from the start of the 2007 growing season were used to parametrize a simple function that described the change in CO₂ concentration for all depths; (2) the DETECT model was run forward for the growing season of 2007, then the modeled CO₂ concentrations for all depths on the final day of the 2007 growing season (31 September 2007) was used as the initial condition for running the DETECT model for 2008. See Sect. S2 in the Supplement for specific details. We set $c_{\text{atm}}(t)$ equivalent to 356 ppm for all t, which was the average near-surface, ambient atmospheric CO_2 concentration measured at the PHACE site in the 2008 growing season. Following the methods of Haberman (1998), we adopted a zero-flux lower BC (Eq. 9c) due to the lack of data at or near a depth of 1 m.

Prior to solving the non-linear PDE (Eq. 1), we expanded the RHS of Eq. (1) using the "product rule for differentiation". We then numerically solved the non-linear PDE (Eq. 1) by employing a forward Euler discretization with a centered difference method for the depth derivative at a depth increment of $\Delta z = 0.01$ m. To ensure numerical stability, we calculate model outputs at a numerical time step of $\Delta t = \frac{dt}{Ndt}$, where dt is the time step at which the predicted outputs are stored (6 h), and Ndt is the number of numerical time steps. Ndt is computed based on the fastest (largest) diffusion coefficient at each time step such that $Ndt = \frac{dt \times \max(D_{gs})}{0.5 \times (\Delta z)^2}$, where max(D_{gs}) is the maximum D_{gs} across all depth increments at time t (Haberman, 1998). We solved Eq. (1) separately for

both root- and microbial-derived CO_2 concentrations, such that for K = R or M:

$$\frac{c_{K}(z,t+\Delta t)-c_{K}(z,t)}{\Delta t} = D_{gs}(z,t)$$

$$\left(\frac{c_{K}(z+\Delta z,t)-2c_{K}(z,t)+c_{K}(z-\Delta z,t)}{(\Delta t)^{2}}\right)$$

$$+\left(\frac{D_{gs}(z+\Delta z,t)-D_{gs}(z-\Delta z,t)}{2\Delta z}\right)$$

$$\left(\frac{c_{K}(z-\Delta z,t)-c_{K}(z+\Delta z,t)}{2\Delta z}\right)$$

$$+S_{K}(z,t).$$
(10)

We rearranged Eq. (10) to solve for $c_K(z, t + \Delta t)$, which was iterated forward for all time steps and depth increments; total CO₂ concentration at each time step and depth is calculated as $c(z, t + \Delta t) = c_R(z, t + \Delta t) + c_M(z, t + \Delta t)$. For clarity, we emphasize that Eq. (10) is the discretized version of Eq. (1), which we require in order to numerically solve Eq. (1) (Haberman, 1998). We programmed the DETECT model v.10 and the numerical solution method in Matlab (Mathworks, 2016).

2.3 Steady-state (SS) solution to the DETECT model

A primary goal of this work was to test if soil CO₂ and associated R_{soil} predicted from the non-steady-state (NSS) model (DETECT) could be distinguished from that of the steadystate (SS) solution. The SS version of Eq. (1), which we refer to as the SS-DETECT model, can be solved analytically as an ordinary differential equation (ODE) by setting the $\partial c/\partial z$ term to zero (Amundson et al., 1998). As with the NSS model, we found the SS solution to Eq. (1) separately for root- and microbial-derived CO₂ concentrations, $c_R^*(z,t)$ and $c_M^*(z,t)$, respectively. Using the upper and lower boundary conditions described for the NSS model (Eq. 9b and c), the analytical SS solutions at time t and depth z are derived by Amundson et al. (1998) and Cerling (1984). The solution is given by

$$c_K^*(z,t) = \frac{S_K^*(t)}{D_{\rm gs}(z,t)} \left(z - \frac{z^2}{2} \right) + c_{\rm atm}(t) \tag{11}$$

$$S_K^*(t) = \frac{1}{100} \sum_{z=0.01}^{1 \text{ m}} S_K(z, t), \qquad (12)$$

where K = R and K = M refers to the soil CO₂ from root (R) and microbial (M) sources, respectively. $S_K^*(t)$ is the depth-averaged source term for microbial or root production (averaging over 100 different 0.01 m increments). The soil CO₂ diffusivity term, $D_{gs}(z, t)$, and upper boundary condition, $c_{atm}(t)$, are the same as previously defined (Eqs. 2 and 9b, respectively; Amundson et al., 1998).

2.4 Application of the DETECT and SS-DETECT models to the PHACE site

In this subsection, we provide an overview of the study site, including the PHACE experiment, and relevant data sources from PHACE that we used to drive the DETECT and SS-DETECT models. We also summarize how we calibrated the models in the context of the PHACE site, and we highlight data that we used to informally validate the general behavior of the models. We conclude by describing the simulation experiments that we conducted to test the effects of soil texture and precipitation variability on the importance of NSS versus SS soil CO_2 conditions.

2.4.1 Field site and PHACE experiment

The Prairie Heating and CO₂ Enrichment (PHACE) field experiment is located in south-central Wyoming (latitude $41^{\circ}50'$ N, longitude $104^{\circ}42'$ W, elevation 1930 m). The site is a mixed-grass prairie with a semi-arid climate characterized by long winters (mean January temperature = -2.5° C) and warm summers (mean July temperature = 17.5° C), with mean annual precipitation of 384 mm (Morgan et al., 2011). The vegetation is predominantly composed of two C₃ grasses, western wheatgrass (*Pascopyrum smithii* (Rydb.) *A. Löve*) and needle-and-thread grass (*Hesperostipa comata* (Trin. & Rupr.)), and a C₄ perennial grass, blue grama (*Bouteloua gracilis* (H.B.K.) Lag). The soil is a fine-loamy, mixed, mesic Aridic Argiustoll, and biological crusts are not present (Bachman et al., 2010).

2.4.2 Environmental driving data

We simulated the transport and production of soil CO₂ for each 0.01 m depth increment, from the surface (0 m) to a depth of 1 m, across all 732 time steps (i.e., four time steps per day [every 6 h] for 183 days from April to September). To do this, we required soil environmental data consisting of water content (θ) and temperature (T_S) and meteorological data including precipitation, air temperature, and air pressure. The θ and T_S data that were used to drive the DETECT model were created using the HYDRUS software (see Sect. 2.4.3), calibrated against actual measurements of θ and T_S . For the meteorological data, actual measurements from the PHACE site were used.

The PHACE experiment involved an incomplete factorial of CO_2 , warming, and irrigation (six treatment levels total), with five replicate plots per treatment level, resulting in a total of 30 instrumented plots. One of the five plots from the control treatment – ambient CO_2 , temperature (no heating), and precipitation (no supplemental irrigation) – was chosen at random and had a system installed to measure soil CO_2 concentrations continuously for three different soil depths (3, 10, and 20 cm). This plot, therefore, provided the data for driving the DETECT and SS-DETECT models. Data that

we used were collected during the growing season (March–September) of 2008; θ was measured hourly at three depths (5–15, 15–25, and 35–45 cm; EnvironSMART probe, Sentek Sensor Technologies, Stepney, Australia) and we used daily averages to drive the models. T_S was measured hourly at two depths (3 and 10 cm) using type-T thermocouples. Hourly precipitation (mm), air temperature (°C), relative humidity (%), and surface barometric air pressure (kPa) were recorded by an automated weather station at the site.

2.4.3 High-resolution environmental data

To accommodate the 0.01 m depth increments specified for the DETECT model, we used the coarse-resolution field data (above) to create finer-resolution driving data. For example, temporal gap-filling of the θ , T_S , and micrometeorological datasets was required due to gaps that occurred during a small number of days (< 1, 6, and 2.5 %, respectively) as a result of instrument failure. We used data from other nearby plots to estimate the values of the missing data, but we also used cubic spline interpolation where gaps remained. Details of these gap-filing methods can be found in Ryan et al. (2015).

We used HYDRUS-1D v4.16.0090 to simulate θ and T_S in 0.01 m increments from a depth of 0.01 to 1 m (Chou et al., 2008; Šimůnek et al., 2008; Piao et al., 2009) based on precipitation data at the site. HYDRUS simulates the movement of water by solving the Richards' equation for water movement (Richards, 1931; Chou et al., 2008; Sitch et al., 2008) and heat transport via Fickian based advection-dispersion equations. Soil hydraulic and heat transport parameters were estimated in HYDRUS using the inverse mode to solve for parameter values based on the PHACE θ (5–10, 15–25, and 35–45 cm) and T_S (3 and 10 cm) data (Šimůnek et al., 2005, 2008). HYDRUS was then run in forward mode based on the tuned soil hydraulic parameters to estimate θ and T_S at all 100 different 0.01 m depth increments at six-hourly time intervals. For consistency, HYDRUS-derived θ and T_S were used as the environmental input data to the DETECT models, even at the depths for which PHACE data were available.

2.4.4 Antecedent soil water and soil temperature conditions

We explicitly evaluated the impact of antecedent (past) θ and T_S conditions on CO₂ production by roots and microbes, motivated by prior work that estimated the relative importance of antecedent conditions and their time scales of influence on soil and ecosystem CO₂ efflux (Cable et al., 2013; Barron-Gafford et al., 2014; Ogle et al., 2015; Ryan et al., 2015). Antecedent soil water content and antecedent soil temperature $-\theta_K^{ant}(z, t)$ and $T_S^{ant}(z, t)$, respectively, for K = R (roots) and M (microbes) were computed as weighted averages of the HYDRUS-produced $\theta(z, t)$ and $T_S(z, t)$ data, respectively. These calculations were done external to the DETECT model, and the antecedent variables were supplied as driving variables to DETECT. For example, for each 0.01 m increment (z) and time period (t), antecedent soil water associated with microbial CO₂ production was calculated as

$$\theta_M^{\text{ant}}(z,t) = \sum_{j=1}^J w(j) \cdot \theta(z,t-j).$$
(13)

The w's are the antecedent importance weights, which sum to 1 from i = 1 (previous time period) to i = J (J previous time periods). The weights were informed by results from an analysis of ecosystem respiration at the PHACE site (Ryan et al., 2015). For microbes, J = 4 days and w = (0.75, 0.25, 0.25, 0.25)0, 0), indicating the strong importance of θ conditions occurring the previous day (j = 1) (Oikawa et al., 2014). Similar equations were used to compute $\theta_R^{\text{ant}}(z, t)$ and $T_S^{\text{ant}}(z, t)$, each with their own set of weights (*w*'s) and time scales (*J*'s). For example, the time step and J for θ differ among microbes (2 days) and roots (3 weeks); for roots, $\theta_R^{\text{ant}}(z,t)$ was computed as a weighted average of past, average weekly values of θ , with j denoting weeks into the past, for J = 4 weeks, and w = (0.2, 0.6, 0.2, 0), indicating a strong lag response to θ conditions occurring two weeks ago (Cable et al., 2013; Ryan et al., 2015). For antecedent soil temperature, we assumed that each of the past four days were equally important by setting the w vectors to (0.25, 0.25, 0.25, 0.25), for both microbes and roots (Ryan et al., 2015). The specification of J and the w's are independent of the DETECT model formulation and can be varied by the user. For clarity we summarize these weight parameters in Table 2.

2.4.5 Overview of parameterization approach using PHACE data

In general, our aim was to specify realistic values for the parameters in the DETECT model. We did not formally "fit" the DETECT model to data, but rather, we simply determined reasonable values based on simple analyses of relevant PHACE data sets, results published for the PHACE site, or results from other relevant studies. The full list of parameters is given in Table 1, and below we describe the logic behind specifying the values in Table 1.

The depth-distributions of root biomass C (C_R, Eq. 3), soil microbial biomass C (C_{MIC}, Eq. 5), and soil organic C (C_{SOM}, Eq. 7) are expressed in terms of a total C content in a 1 m deep soil column (R^* , M^* , and S^* , respectively; mg C cm⁻²), multiplied by the proportion of that C that occurs at depth z ($f_R(z)$, $f_M(z)$, and $f_S(z)$, respectively). See Sect. S1 (Supplement) for details. Regarding the data, soil organic C (Fig. S5, Supplement) was determined by combustion of acidified, root-free soil collected from 0–5, 5–15, 15– 30, 30–45, 45–75, and 75–100 cm depths, using a Costech Elemental Analyzer. Microbial biomass C was determined by the chloroform fumigation and extraction in 0.05 M K₂SO₄ (Carrillo et al., 2014b). Extracts were analyzed for total C

Table 2. Summary of quantities in the non-steady-state (DETECT) model that vary by depth only (z), or by depth (z) and time (t). Those in group 1 represent input variables (derived prior to the running of the DETECT model), while group 2 contains the modeled quantities (used as part of the operation of the DETECT model). Equation (S1) can be found in Sect. S1 in the Supplement.

Symbol	Description	Units	Eq(s)
Group 1			
$f_R(z)$	A function describing the distribution by depth of root carbon.	unitless	S
$C_R(z,t)$	The amount of root carbon.	${ m mg}{ m C}{ m cm}^{-3}{ m h}^{-1}$	3, S
$f_S(z)$	A function describing the distribution by depth of carbon from soil organic matter (SOM)	unitless	S
$C_{SOM}(z)$	The amount of carbon from SOM.	${ m mg}{ m C}{ m cm}^{-3}{ m h}^{-1}$	7, S
$f_M(z)$	A function describing the distribution by depth of microbial carbon	unitless	S
$C_{MIC}(z)$	The amount of microbial carbon.	${ m mg}{ m C}{ m cm}^{-3}{ m h}^{-1}$	3, S
$\theta(z,t)$	Soil water content	${ m m}^{3}{ m m}^{-3}$	3, 6,
$\theta_R^{\mathrm{ant}}(z,t)$	Antecedent soil water content (used in S_R function) calculated as a weighted average of soil water content from the previous four days. The weights are $w = (0.75, 0.25, 0.0)$.	$m^{3} m^{-3}$	
$\theta_M^{\mathrm{ant}}(z,t)$	Antecedent soil water content (used in $S_{\rm M}$ function) calculated as a weighted average of soil water content from the previous four days. The weights are $w = (0.2, 0.6, 0.2, 0)$.	$\mathrm{m}^3\mathrm{m}^{-3}$	
$T_S(z,t)$	Soil temperature	Kelvin	3,
$T_S^{\mathrm{ant}}(z,t)$	Antecedent soil temperature calculated as a weighted average of soil temperature from the previous four weeks. The weights are $w = (0.25, 0.25, 0.25, 0.25)$.	Kelvin	3,
Group 2			
c(z,t)	Total soil CO ₂ .	$mg CO_2 m^{-3}$	
$c_R(z,t)$	Soil CO ₂ derived from root sources.	$\mathrm{mg}\mathrm{CO}_{2}\mathrm{m}^{-3}$	
$S_R(z,t)$	Source term describing the production of soil CO ₂ from root respiration.	$mg CO_2 m^{-3}$	
$c_{\mathbf{M}}(z,t)$	Soil CO ₂ derived from microbial sources.	$mg CO_2 m^{-3}$	
$S_{\mathbf{M}}(z,t)$	Source term describing the production of soil CO ₂ from microbial respiration.	$\mathrm{mg}\mathrm{CO}_{2}\mathrm{m}^{-3}$	
$D_{\rm gs}(z,t)$	Diffusivity of soil CO ₂	$m^{2} s^{-1}$	1,
$\varphi_{\rm g}(z,t)$	Air-filled soil porosity.	$m^{3} m^{-3}$	1,
$C_{SOL}(z,t)$	The amount of soluble carbon from SOM.	${ m mg}{ m C}{ m cm}^{-3}{ m h}^{-1}$	5,
$V_{\max}(z,t)$	Maximum potential decomposition rate (microbial carbon).	$mg C cm^{-3} h^{-1}$	
$E_o(z,t)$	Analogous to energy of activation.	Kelvin	4

on a total organic carbon analyzer (Shimadzu TOC-VCPN; Shimadzu Scientific Instruments, Wood Dale, IL, USA) after treating with 1 M H3PO4 (1 μ L per 10 mL of extract) to remove any carbonates. Root biomass C was estimated from ash-free root biomass and elemental analysis (Carrillo et al., 2014a; Mueller et al., 2016). The solubility parameter, *p*, was estimated as the ratio of C_{SOL} to C_{SOM} using measurements of these two quantities which were based on unfumigated extracts obtained for microbial biomass estimations as above (C_{SOL}) and on total *C* concentration in soil (C_{SOM}).

The values used for the base microbial respiration rates and the half-saturation constant (V_{Base} [Eq. 6] and K_{m} [Eq. 5]; Table 1) were estimated by fitting the microbial respiration submodel, but without the C_{MIC} or CUE terms (Eq. 5), to microbial respiration data from the PHACE control plots (Fig. S7, Supplement). The C_{MIC} and CUE terms were not included in this earlier version of S_{M} submodel – which was used for model calibration purposes – because we did not have measurements of these two variables at the time. We estimated V_{Base} and K_{m} using a Markov chain Monte Carlo approach, identical to the approach used in Ryan et al. (2015). In the absence of root respiration data, we assumed that base root respiration (R_{Rbase} [Eq. 3]; Table 1) was proportional to the microbial base rate term (Hanson et al., 2000). The parameters denoting the effects of current soil moisture (e.g., α_1 ; Eq. 4a), antecedent moisture (α_2), and the interaction between current and antecedent moisture (α_3) on root and microbial respiration were derived from Ryan et al. (2015), based on an analysis of ecosystem respiration $(R_{\rm eco})$ data from PHACE. However, we adjusted the values (Table 1) by trial and error to reflect the expectation that the effects of current soil moisture should be stronger for microbial compared to root respiration because microbes tend to respond more rapidly to precipitation pulses (Risk et al., 2008), whereas root respiration is likely to show a delayed response which depends more strongly on past moisture conditions (Cable et al., 2008, 2013). Of the remaining two parameters describing $S_{\rm M}$ (Eqs. 5–6; Table 1), the value of CUE was based on results from a soil incubation study conducted at a nearby site (Tucker et al., 2013), whilst our value for D_{liq} was taken from Davidson et al. (2012). Three parameters (E_o^* , T_o , and α_4 ; Eq. 4a–b) were shared between the S_R and S_M submodels, and their values were also obtained from Ryan et al. (2015). Finally, the parameters used for CO₂ diffusivity (*b*, BD, and φ_{g100} ; Eq. 2) were based on published, site-specific data (Morgan et al., 2011).

2.4.6 Informal model validation with soil respiration measurements

We evaluated the accuracy of the DETECT model by comparing (1) predicted R_{soil} (Eq. 8) against plot-level measurements of ecosystem respiration (R_{eco}) (see below) and (2) predicted soil CO₂ concentrations, c(z, t), versus observed concentrations; all observed data were from the PHACE study. Since we did not rigorously parameterize the DETECT model with PHACE data, we were simply looking for reasonable, qualitative agreement between the modeled variables and the observations (e.g., similar order of magnitude, comparable temporal trends). Observed R_{eco} was measured on control plots every two to four weeks during the target growing season, using a canopy gas exchange chamber, and instantaneous fluxes were scaled to daily rates using a linear, empirical function (Jasoni et al., 2005; Bachman et al., 2010). We assumed that R_{soil} was similar to R_{eco} given that above ground biomass was < 20% of total plant biomass (Mueller et al., 2016). Measurements of microbial respiration were obtained by applying glyphosate herbicide to small subplots in May 2008, limiting ecosystem CO₂ efflux to microbial sources (Pendall et al., 2013), Non-steady state soil chambers were used to estimate the resulting surface soil fluxes every two weeks around midday (Oleson et al., 2013; Ogle et al., 2016). Soil CO₂ concentrations were also measured with non-dispersive infrared sensors (Vaisala GM222, Finland) installed at 3, 10, and 20 cm below the soil surface, averaged on an hourly basis (Risk et al., 2008; Vargas et al., 2011; Brennan, 2013). Observations of soil [CO₂] for control plots were compared against predictions of c(z, t)at z = 0.03, 0.1, and 0.2 m and at the corresponding times.

2.5 Simulation experiments

We evaluated the impact of three potentially important factors that could affect the frequency of NSS (Eqs. 1 and 9a–c) relative to SS (Eq. 11) conditions: (1) soil texture, (2) precipitation patterns, and (3) importance of antecedent conditions. In the control (Ctrl) scenario, we calculated the source terms and diffusion terms (S_K and D_{gs} in Eqs. 1 and 2) based on soil environmental (θ and T_S), soil texture (sandy clay loam: 60 % sand, 20 % silt, 20 % clay), and meteorological data (e.g., precipitation) measured at the PHACE site in 2008. We varied soil texture, relative to that of the site, by altering the relative amounts of sand, silt, and clay, giving three levels (Table 3): 80 % sand, 10 % silt, and 10 % clay (sandy loam, scenario denoted as ST-Sa); 20 % sand, 60 % silt, and 20 % clay (silt loam, ST-Si); 20 % sand, 20 % silt, and 60 % clay (clay, ST-Cl). The control (Ctrl) scenario was also paired with the observed daily precipitation data for 2008. We explored three additional precipitation scenarios, under the control soil texture, by shifting the daily precipitation to occur one month earlier, or one month later, or by using precipitation data from 2009 (scenarios P-E, P-L and P-FM, respectively; Table 3). For P-FM, we chose 2009 because it had approximately the same total precipitation between April and September as 2008 (340 and 348 mm for 2008 and 2009, respectively), but it fell as more frequent events of smaller magnitudes. For each texture and precipitation scenario, HYDRUS was used to compute the corresponding T_S and θ at the required depth and time intervals. Specifically, the different soil texture and precipitation regimes were used as inputs for the HYDRUS software when generating T_S and θ for all 100 depths and all 732 time points. Hence, the differences in soil texture and differences in precipitation regimes were implemented by using different input files for the HYDRUS-generated θ and T_S data.

All of the above scenarios assumed that antecedent conditions were not important, which was achieved by setting all antecedent effects parameters (α_2 , α_3 , and α_4 ; Table 1) equal to zero. We contrasted these scenarios against ones that included antecedent conditions (thus, computed θ_K^{ant} and T_S^{ant} in Eqs. 3 and 6) in the calculation of soil CO₂ production by roots (K = R) and microbes (K = M); all such scenario names were appended with "ant" (Table 3, Fig. 1a). For each scenario summarized in Table 3, we evaluated the potential for NSS conditions by comparing the predicted R_{soil} produced by the DETECT model versus the SS-DETECT model.

3 Results

3.1 Control scenarios

Soil CO₂ was in steady state (SS) during most of the growing season under the control soil texture (sandy clay loam) and precipitation conditions that assumed no antecedent affects (Ctrl scenario). For example, soil respiration (R_{soil}) predicted by the DETECT model was approximately equal to R_{soil} predicted by the SS-DETECT model during times of no or little precipitation (Fig. 2a, days < 218 or > 230). Conversely, R_{soil} predicted by the SS-DETECT model was temporarily greater and more variable than that predicted by the DETECT model immediately following a large precipitation event (Fig. 2a, days 218–229). However, the total cumulative R_{soil} between days 92 to 274 – hereafter "total growing season R_{soil} " – under SS (497 g C m⁻²) versus NSS (498 g C m⁻²) assumptions was approximately equal (a difference of ~ 0.2 %).

The differences between the R_{soil} from DETECT and SS-DETECT using the antecedent parametrization of the

Table 3. The scenario code, description, and summary of results associated with each model scenario; the 14 scenarios below were applied to both the DETECT and SS-DETECT models. The scenarios involved a non-factorial combination of different soil texture, precipitation regimes, and inclusion/exclusion of antecedent effects on the root and microbial CO_2 production rates.

Scenario	Description	Primary result(s)	
Scenarios th	at assume no antecedent effects		
Ctrl (control)	Uses soil texture (sandy clay loam: 60 % sand, 20 % clay) and precipitation (for 2008) data from the PHACE site; CO ₂ production only responds to concurrent environmental conditions.	R_{soil} was very similar under SS and NSS soil CO ₂ assumptions.	
Soil texture	scenarios		
ST-Sa	Same as Ctrl, but the soil texture is set to sandy loam (80% sand, 10% clay).	For ST-Cl, R_{soil} was greater in magnitude and more different under SS vs. NSS conditions,	
ST-Si	Same as Ctrl, but the soil texture is set to silt loam (20 $\%$ sand, 20 $\%$ clay).	due to NSS conditions producing greater R_{soi} after a major precipitation event. The results are	
ST-Cl	Same as Ctrl, but the soil texture is set to clay (20 $\%$ sand, 60 $\%$ clay).	similar, but muted, for the ST-Si scenario.	
Precipitation	a scenarios		
P-E	Same as Ctrl, but daily precipitation was shifted to occur one month earlier.	Varying the timing or magnitude of precipita- tion pulses had little effect on the magnitude of	
P-L	Same as Ctrl, but daily precipitation was shifted to occur one month later.	R_{soil} or on the difference between SS and NSS predictions of R_{soil} .	
P-FM	Same as Ctrl, but daily precipitation was based on data from 2009, which is characterized by more frequent, smaller events.		
Scenarios th	at incorporate antecedent effects on CO2 production rates		
Ctrl-ant	All scenarios parallel those described above, except both current	$R_{\rm soil}$ was generally greater in magnitude under	
ST-Sa-ant	and antecedent conditions (past soil water and past soil temper-	both SS and NSS conditions, especially for ST-	
ST-Si-ant	ature) are used in the calculation of the source terms (i.e., root	Si-ant and ST-Cl-ant (relative to ST-Si and ST-	
ST-Cl-ant	and microbial CO ₂ production rates).	Cl).	
P-E-ant			
P-L-ant P-FM-ant			

source terms of the models (Ctrl-ant scenario; Fig. 2b) were generally consistent with the results from the Ctrl scenario (Fig. 2a). However, the magnitude of R_{soil} predicted by both the DETECT and SS-DETECT models was up to $9 \text{ g C m}^{-2} \text{ day}^{-1}$ greater during days following the major rain event (i.e., during days 230-243) when antecedent conditions were considered. Moreover, the incorporation of antecedent effects led to a longer delay between the timing of the major rain event and the maximum R_{soil} , which occurred \sim five days later than when only current conditions were considered (Fig. 2a vs. 2b). As a result, total growing season $R_{\rm soil}$ was ~ 15 % higher under the Ctrl-ant scenario (e.g., 571 g C m^{-2} under NSS assumptions, Fig. 2b) compared to the Ctrl scenario (e.g., 498 g C m^{-2} under NSS, Fig. 2a). This increase in predicted R_{soil} under the Ctrl-ant scenario for days 230-243 was primarily driven by greater root respiration (Fig. 2a vs. 2b).

3.2 Effects of soil texture

Varying soil texture resulted in the greatest difference in daily R_{soil} between the DETECT and SS-DETECT models; however, integrated over the growing season, these differences were very small (Fig. 3a, b, c). In particular, total growing season $R_{\rm soil}$ predicted by SS-DETECT was ~ 1.5 % less than predicted by DETECT for soils consisting primarily of sand and silt (ST-Sa and ST-Si scenarios; Fig. 3a, b), but was ~ 3.3 % less for a clay dominated soil (ST-Cl scenario; Fig. 3c red versus grey bars). These differences in R_{soil} under NSS versus SS assumptions were approximately the same for the scenarios involving antecedent effects (Fig. 3d, e, f). Despite the minor differences at the growing season scale, notable differences emerged at the daily scale. For example, with the largest precipitation event of the year and the 10 days that followed (days 218–228), the median absolute difference between daily R_{soil} from the SS-DETECT and DETECT models was 22-24 % for the ST-Sa and ST-Si scenarios regardless of whether or not antecedent variables are

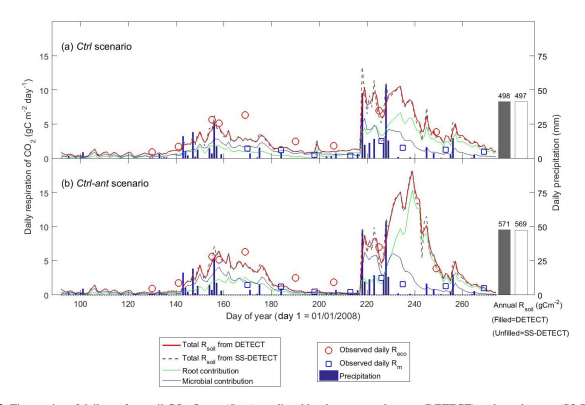


Figure 2. Time series of daily surface soil CO₂ fluxes (R_{soil}) predicted by the non-steady-state (DETECT) and steady-state (SS-DETECT) models over the growing season (1 April–30 September), based on the control scenarios (**a**) without (Ctrl) and (**b**) with (Ctrl-ant) antecedent effects (see Table 2). Only R_{soil} is simulated using the SS-DETECT model, whereas R_{soil} and its root and microbial contributions are simulated using the DETECT model. The predicted fluxes are overlaid with observed ecosystem respiration (R_{eco} ; R_{soil} + aboveground plant respiration) and microbial respiration (R_m ; based on plots where vegetation was removed).

included (Figs. 3 and S3a, b). These differences increased to 31-35% for the two clay soil texture scenarios (ST-Cl and ST-Cl-ant).

Soil texture also affected the magnitude of predicted R_{soil} compared to the control scenarios, both with and without antecedent effects (Ctrl-ant and Ctrl, respectively). In particular, we found that total growing season R_{soil} , whether from the DETECT or the SS-DETECT model, was $\sim 30\%$ and ~ 60 % higher for the ST-Si and ST-Cl scenarios relative to the Ctrl scenario (Figs. 3b, c, 4a). The change in R_{soil} was negligible, however, when the sand content was increased from 60 % (Ctrl) to 80 % (ST-Sa) for both models (Figs. 3a, 4a). The antecedent versions of the fine-textured scenarios (ST-Si-ant and ST-Cl-ant) resulted in \sim 45 and \sim 95% increases in total growing season R_{soil} , respectively, compared to the Ctrl-ant scenario (Figs. 3e, f, 4b). As with the Ctrl-ant scenario (Sect. 3.1), greater root respiration following the end of the second precipitation period between days 230 and 245, primarily drove the larger percentage increases for the SL-Siant and SL-Cl-ant scenarios compared to the non-antecedent versions.

3.3 Effects of precipitation regimes

Although varying the timing, frequency, or magnitude of precipitation led to little difference between R_{soil} as predicted by the DETECT and SS-DETECT models (Fig. S2), these precipitation regimes did affect the magnitude of R_{soil} predicted by both models. For example, total growing season $R_{\rm soil}$ predicted under the alternative precipitation scenarios was lower relative to the Ctrl scenario. This decrease was relatively small (5-10%) for the non-antecedent versions of the models (Fig. 4c), but was comparatively larger (15-22%) for the antecedent versions (Fig. 4d). This reduction appears to be driven by the amount of time over which daily $R_{\rm soil}$ responded to the second precipitation period, which occurred around day 220, 190, and 250 in the Ctrl, P-E, and P-L scenarios, respectively. Following this precipitation event, daily R_{soil} achieved values around $10 \,\mathrm{g}\,\mathrm{C}\,\mathrm{m}^{-2}\,\mathrm{day}^{-1}$ for about 20 days in the Ctrl scenario (Fig. 2a, days 220-240), but for only about five days in the P-E and P-L scenarios (Fig. S2a, b, after days 190 and 250, respectively). Increasing the frequency of precipitation while retaining approximately the same annual amount (i.e., scenario P-FM) resulted in daily R_{soil} being consistently less than that of the

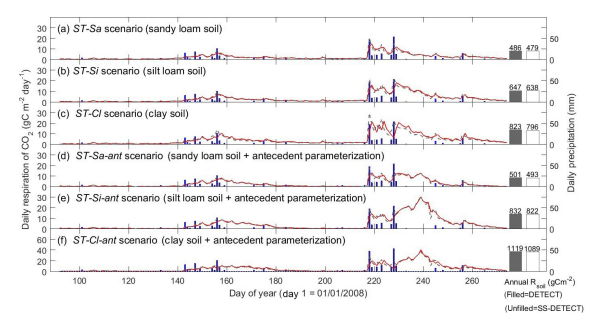


Figure 3. Time series of daily surface soil respiration (R_{soil}) predicted from the non-steady-state (NSS) DETECT model (red solid lines) and the steady-state (SS-DETECT) model (grey dashed lines), for different soil texture scenarios. The first three scenarios are the same as the control (Ctrl), except they assume a different soil texture: (a) more sandy soil, (b) more silty soil, or (c) more clayey soil. Panels (d), (e), and (f) show the R_{soil} predictions from the same soil texture scenarios as in (a)–(c), but also include the antecedent effects of soil moisture and temperature. See Table 2 for descriptions of each scenario. R_{soil} predictions are overlaid with daily precipitation.

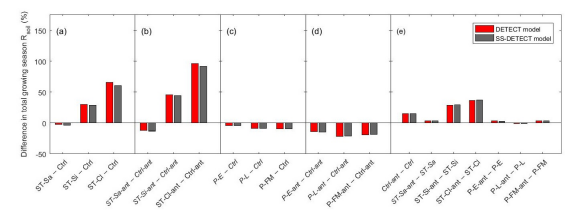


Figure 4. Differences of total growing season (April–September) soil respiration (R_{soil}) as predicted by the non-steady-state (DETECT) and steady-state (SS-DETECT) models, for different pairs of scenarios. Comparisons are grouped so that they quantify the effects of (**a**) soil texture without antecedent effects, (**b**) soil texture with antecedent effects, (**c**) precipitation without antecedent effects, (**d**) precipitation with antecedent effects, and (**e**) antecedent effects. See Table 2 for descriptions of each scenario.

Ctrl scenario, which led to a reduction in total growing season R_{soil} in the P-FM scenario (Fig. S2c).

3.4 Effects of antecedent responses

When antecedent soil water content and soil temperature were included in the DETECT model we found that predicted R_{soil} was 15 % greater for the control scenario and 29–37 % greater for the fine textured soil scenarios, compared to the corresponding scenarios that did not include antecedent con-

ditions (Fig. 4e). When the sand content was 80 % or for any of the different precipitation regimes, there was a negligible difference between R_{soil} predicted by the antecedent versus non-antecedent parametrizations of DETECT.

Daily R_{soil} predicted by the DETECT model based on the Ctrl and Ctrl-ant scenarios agreed well with observed ecosystem respiration (R_{eco}), but R_{eco} was slightly higher than predicted R_{soil} (Fig. 2a, b), which was expected since $R_{eco} = R_{soil}$ + aboveground autotrophic respiration. For the most part, this data-model agreement was similar whether

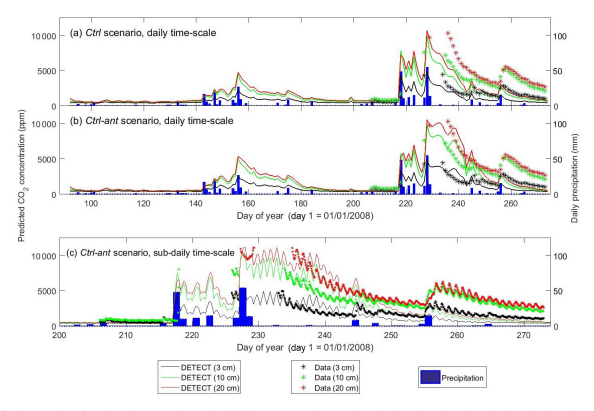


Figure 5. Time series of predicted versus observed soil CO_2 concentrations at 3 cm depth, 10 cm depth, and 20 cm depth, where the predictions are based on the non-steady-state (NSS) DETECT model. Predicted [CO_2] is shown for the daily time scale for the control scenarios (**a**) without (Ctrl) and (**b**) with (Ctrl-ant) antecedent effects, and for (**c**) the subdaily (every 6 h) time scale for the Ctrl-ant scenario. Units are in parts per million (ppm).

the antecedent model terms were included (Fig. 2b) or not (Fig. 2a). Unfortunately, R_{eco} data were not available during the time period (days 230–250) associated with the greatest disagreement between the Ctrl and Ctrl-ant scenarios. During this period, frequent hourly measurements of soil [CO₂] were in better agreement with predicted soil CO₂ from the Ctrl-ant scenario compared to the Ctrl scenario (Figs. 5a, b, S4a, b). After day ~ 250, based on the DETECT model, both scenarios (Ctrl and Ctrl-ant) under-predicted the observed soil [CO₂] by ~ 50 % (Fig. 5).

4 Discussion

The DETECT and SS-DETECT models provide a framework for evaluating the circumstances under which steady-state (SS) assumptions of soil CO₂ production and surface soil respiration (R_{soil}) are valid, and to identify the major physical (i.e., soil texture, soil moisture) and/or biological (i.e., root and microbial respiration responses) factors that lead to nonsteady-state (NSS) conditions.

4.1 Steady-state versus non-steady-state conditions

At the seasonal scale, there was reasonable agreement between total growing season R_{soil} predicted under the assumption of SS versus NSS conditions, but the strength of this agreement depended on soil texture (see Sect. 4.2). At the daily scale, R_{soil} predicted by the DETECT model deviated from values expected under the assumption of SS conditions for 11 days or 4% of the days during the April-September growing season (Fig. 2, days 218-228). These discrepancies, attributed to NSS conditions, were generally limited to periods following large rain events. For applications that assume SS conditions, such as isotopic partitioning studies (Hui and Luo, 2004; Ogle and Pendall, 2015), the SS assumption seemed reasonable during periods of minimal or no precipitation, representative of times during which soil water content changes very little or gradually. For sites or time periods characterized by pulsed precipitation patterns, our results suggested that NSS conditions would be more likely over longer periods of time.

4.2 Effect of varying soil texture

Our results indicated that soil texture exerts the strongest control over the prevalence of NSS soil CO₂ conditions. For

a predominantly (e.g., 60%) sandy or silty soil, soil CO₂ transport and efflux generally aligned with the SS assumption (Figs. 2, 3a–b). This was consistent with previous work that used SS models to predict R_{soil} for similar soil types (Baldocchi et al., 2006; Vargas et al., 2010).

For very fine-texture soil dominated by clay, however, SS assumptions were far less appropriate. The larger difference - relative to the Ctrl scenario - in R_{soil} predicted under SS versus NSS conditions for fine-texture (i.e., 60 % clay) soil was apparent at both the growing season scale and the daily scale following a large precipitation event (Figs. 3, S3a, b). In general, the DETECT model predicted that R_{soil} should be higher in clay compared to sandy soil after precipitation events, a result supported by field experiments (Cable et al., 2008), but this texture effect is muted under assumptions of SS. Moreover, recovery of R_{soil} to SS rates after a large rain event took \sim 30 days in the clay soil (Fig. 3c, days 218 to 248) compared to ~ 10 days for the other coarser soil texture scenarios (Figs. 2, 3a–b, days 218 to \sim 230). These effects of soil texture on the prevalence of NSS conditions can be attributed to soil physical properties and their effects on airfilled porosity and CO₂ diffusivity. Fine textured soils have smaller pores and tend to retain water for longer (Bouma and Bryla, 2000), which has the effect of decreasing soil CO_2 diffusivity (Fig. 6). Thus, under moist conditions that follow a rain event, it may take about $15 \min$ for a CO₂ molecule produced at 0.5 m to diffuse to the surface in a clay soil compared to only 1–2 min for a sandy soil. This means that the increase in CO₂ concentration near the soil's surface will be almost immediate under a coarsely textured soil (Fig. 6a), but slightly delayed under a finely texture soil. Finally, finetextured soils have slower infiltration rates (Hillel, 1998), delaying the exposure of more deeply distributed roots and microbes to increased moisture availability. While this effect may not directly impact the SS assumption, it would lead to greater time lags between precipitation pulses and R_{soil} peaks.

These findings have important implications for studies that rely on the SS assumption to predict subsurface soil CO₂ production. The SS assumption may be sufficient for systems defined by coarse-textured soils, but it may lead to erroneous conclusions if applied to fine-textured soils, especially at the very short-term scale (e.g., diurnal R_{soil}) during times of precipitation. Our simulation experiments made the simplifying assumption that soil texture is constant with depth, but in many ecosystems, texture may vary greatly with depth (Ogle et al., 2004). An important next step is to extend the simulations to explore the impacts of depth-varying soil texture on SS versus NSS conditions. The DETECT model can easily accommodate such modifications; allowing soil texture to vary by depth would have a direct effect on soil water content, which is simulated outside of DETECT using HYDRUS (Chou et al., 2008; Šimůnek et al., 2008; Piao et al., 2009), and can accommodate such depth variation.

4.3 Effect of varying the timing or frequency of precipitation

Unlike soil texture, varying the timing, frequency, and magnitude of precipitation resulted in predicted R_{soil} that was almost identical under SS and NSS assumptions, both at the growing season and daily time scales (Fig. S2). We had anticipated that such changes in the precipitation regime would impact SS conditions via impacts on soil air-filled porosity and potentially by changing the covariance between soil water and soil temperature, both of which affect soil CO₂ diffusivity (e.g., see Eq. 2). We did not explore, however, the effect of decreasing the frequency while simultaneously increasing the magnitude of individual pulses. We hypothesize that this latter scenario could produce more exaggerated or extended NSS conditions given that large rain events would infiltrate deeper, reducing CO2 diffusivity across greater soil depths, thus slowing the transport of more deeply derived CO_2 . Increasing the number of small events, as done in the P-FM scenario, would generally confine water inputs to shallow layers, from which CO₂ has shorter distances to travel to reach the surface, creating less opportunity for R_{soil} to exhibit NSS behavior.

4.4 Effect of antecedent conditions

The inclusion or exclusion of antecedent soil moisture and temperature effects on CO₂ production rates had little to no impact on the balance between SS versus NSS behavior of $R_{\rm soil}$. However, incorporating antecedent effects generally increased the magnitude of R_{soil} . This was for two reasons. Firstly, microbial respiration was stimulated more during the initial onset of the main precipitation period when antecedent effects were considered (Fig. 2b vs. Fig. 2a, day 218, blue line). This is expected because the instantaneous response of microbes to a rain event is expected to be greater following a dry period compared to during a wet period (Xu et al., 2004; Sponseller, 2007; Cable et al., 2008, 2013; Thomas et al., 2008). These dynamics are incorporated in the antecedent version of the models when the parameter corresponding to the interaction between current and antecedent soil water content is negative (e.g., α_3 , Table 1). Secondly, root respiration was greatly enhanced following the end of this period of precipitation (Fig. 2b vs. Fig. 2a, days \sim 230– 250, green line), despite there being little precipitation after day 230 (Fig. 2b). This likely occurred because our DETECT model assumed that soil water over relatively longer time periods (past 1-2 weeks, Eq. 12) affects current root respiration rates. This partly reflects the mechanism that roots are able to take up more soil water that has infiltrated to deeper depths (Cable et al., 2013). The microbes, however, are coupled to past conditions over comparatively short time periods (a couple of days).

The importance and benefit of including antecedent terms for modeling soil respiration or ecosystem respiration has

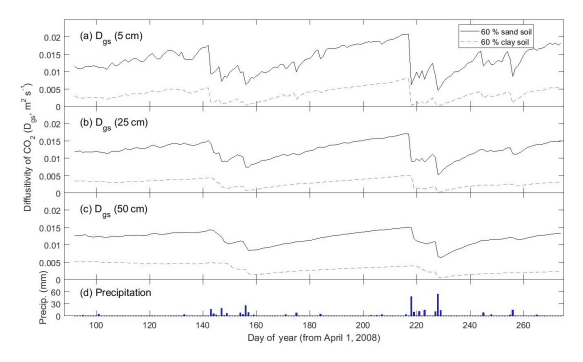


Figure 6. Time series of how the modeled diffusivity of CO_2 (D_{gs}) at three different depths (5, 25, and 50 cm) varies between a predominantly sandy soil (solid line) and a predominantly clay soil (dashed line). Predictions are from the non-steady state (DETECT) model for the Ctrl (60% sand) and ST-Cl (60% clay) scenarios; see Table 2 for a description of the scenarios.

been well documented (Cable et al., 2013; Barron-Gafford et al., 2014; Ryan et al., 2015). Thus, we encourage future studies to include influences of past conditions when modeling subsurface and surface CO_2 fluxes. Fortunately, our simulation experiments suggest that the lagged responses of microbial and root respiration to soil moisture and temperature do not have a notable impact on the SS assumption.

4.5 Comparison of modeled soil CO₂ with data

The good agreement between modeled and observed soil CO₂ concentrations – particularly when including antecedent effects - was very encouraging because the DETECT model was not rigorously tuned or calibrated to fit data on soil $[CO_2]$ or ecosystem CO₂ fluxes (R_{eco}) (Figs. 5, S4a,b). However, discrepancies remained between the predicted and observed CO₂ fluxes, particularly after rain events. These discrepancies could be an artifact of the input data used to calculate CO₂ production (i.e., the source term). Some parameter values were drawn from the literature and others were estimated by fitting a non-linear regression model to data. For example, the parameters describing the current and antecedent soil water content effects (α 's) were obtained by fitting a non-linear model to R_{eco} data (Ryan et al., 2015). While measured R_{eco} represents both root respiration and microbial respiration contributions, it also reflects aboveground respiration, which is not currently treated in the DETECT model. Moreover, we made further assumptions about how the R_{eco} parameter estimates translate to component processes (root and microbial responses), and we relied on literature information about how microbes and roots respond to precipitation events (e.g., the timing, magnitude, and lags). Future studies could rigorously fit the DETECT model to field data, such as observations of R_{soil} , soil CO₂ concentrations, and ¹³C isotope fluxes. Using a Bayesian methodology to do this would allow one to incorporate multiple data sets to inform all parameters in DETECT.

4.6 Non-steady state model of soil CO₂ transport and production

An important contribution of this study was the development of a non-steady state (NSS) model of soil CO₂ transport and production (the DETECT model version 1.0), which is particularly useful for systems that may frequently experience NSS conditions. Other comparable NSS models exist (e.g., Šimůnek and Suarez, 1993; Fang and Moncrieff, 1999; Hui and Luo, 2004), but they generally treat the production (source) terms - root/rhizosphere respiration and microbial decomposition of soil organic matter - simplistically, and accompanying model code is not available. Our DETECT v1.0 model includes more detailed submodels for the production terms, inspired by recent studies (e.g., Lloyd and Taylor, 1994; Pendall et al., 2003; Davidson et al., 2012; Todd-Brown et al., 2012; Carrillo et al., 2014a); in contrast to these studies, which essentially described models for "bulk" soil, we applied the CO₂ production models to every depth increment. Additionally, we have provided model code, implemented in Matlab (see Code Availability section), with the goal of making the DETECT model, and ability to accommodate NSS conditions, more accessible to potential users.

Future versions of DETECT could include other characteristics of soil CO₂ production and transport not included in v1.0. These include (1) a transport process that simulates the physical displacement of CO₂ in the soil following a precipitation event; (2) alternative options for some of the functions used, for example, there are a number of ways of estimating soluble soil C from soil organic C and soil water content (Eq. 7); (3) an estimation of the parameters and their associated uncertainties using formal methods (e.g., MCMC) that rely on measurements of C stocks and C fluxes; (4) quantification of the uncertainty of the model outputs (soil CO₂ concentration, soil respiration) by propagation of uncertainty from the parameters; (5) coupling DETECT with a dynamic soil C model in order for the C_{SOM} pools to be dynamic rather than prescribed independently of DETECT.

5 Conclusions

Determining the conditions under which steady-state (SS) assumptions are appropriate for modeling soil CO₂ production, transport, and efflux is crucial for accurately modeling the contribution of soils to the carbon cycle. We found that soil texture exerted the greatest control over whether SS assumptions are appropriate. When the soil at a site is coarse (60%)or more sand), SS assumptions appeared to be appropriate, and one could apply a simpler, more computationally efficient SS model, such as SS-DETECT (see also Amundson et al., 1998). As the soil texture becomes increasingly finer, SS assumptions start to break down, especially following large precipitation events that can greatly impact soil water content and associated soil air-filled porosity, thus affecting CO₂ diffusivity. Under such conditions, the more complex and computationally demanding NSS model (DETECT) is preferred. We found that precipitation regime characteristics and/or the inclusion of antecedent soil moisture and temperature conditions had little singular effect on whether SS or NSS assumptions were appropriate. However, while these factors do not directly impact SS versus NSS behavior, they were found to be important for accurately modeling the soil carbon cycle because they notably impacted the magnitude of the soil CO₂ efflux.

Code availability. All of the Matlab script files for running the DE-TECT model can be found in Ryan and Ogle (2017b). These Matlab script files are set up so that the model runs at the PHACE field site. The above weblink also provides a user manual which gives instructions for running DETECT at either the PHACE site or at a user specified field site. We also provide Matlab script files for creating a time series of predicted versus observed soil respiration (Fig. 2) and a time series of predicted versus observed soil CO₂ (Fig. 5). These can be found via http://doi.org/10.5281/zenodo.927313. Following publication, these Matlab files and the data files (see next section) will be available to download from the Ogle lab website via http://jan.ucc.nau.edu/ogle-lab/.

Data availability. Measurement data made at the PHACE field site, which are required as inputs for the DETECT model, can be found via Ryan and Ogle (2017a).

The Supplement related to this article is available online at https://doi.org/10.5194/gmd-11-1909-2018-supplement.

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

Acknowledgements. We thank Dan LeCain, David Smith, and Erik Hardy for implementing and managing the PHACE experiment, and Jack Morgan for project leadership. This material is based upon work supported by the US Department of Agriculture Agricultural Research Service Climate Change, Soils & Emissions Program; USDA-CSREES Soil Processes Program (#2008-35107-18655); US Department of Energy Office of Science (BER), through the Terrestrial Ecosystem Science program (#DE-SC0006973); the Western Regional Center of the National Institute for Climatic Change Research; and by the National Science Foundation (DEB#1021559). Any opinions, findings, and conclusions or recommendations expressed in this material are those of the author(s) and do not necessarily reflect the views of the National Science Foundation.

Edited by: Carlos Sierra

Reviewed by: Fernando Moyano and one anonymous referee

References

- Amundson, R., Stern, L., Baisden, T., and Wang, Y.: The isotopic composition of soil and soil-respired CO₂, Geoderma, 82, 83– 114, 1998.
- Atkin, O. K. and Tjoelker, M. G.: Thermal acclimation and the dynamic response of plant respiration to temperature, Trends Plant Sci., 8, 343–351, 2003.
- Bachman, S., Heisler-White, J. L., Pendall, E., Williams, D. G., Morgan, J. A., and Newcomb, J.: Elevated carbon dioxide alters impacts of precipitation pulses on ecosystem photosynthesis and respiration in a semi-arid grassland, Oecologia, 162, 791–802, 2010.
- Baldocchi, D., Tang, J., and Xu, L.: How switches and lags in biophysical regulators affect spatial-temporal variation of soil respiration in an oak-grass savanna, J. Geophys. Res., 111, G02008, https://doi.org/10.1029/2005JG000063, 2006.
- Barron-Gafford, G. A., Cable, J. M., Bentley, L. P., Scott, R. L., Huxman, T. E., Jenerette, G. D., and Ogle, K.: Quantifying the timescales over which exogenous and endogenous conditions affect soil respiration, New Phytol., 202, 442–454, 2014.

- Birch, H.: The effect of soil drying on humus decomposition and nitrogen availability, Plant Soil, 10, 9–31, 1958.
- Borken, W. and Matzner, E.: Reappraisal of drying and wetting effects on C and N mineralization and fluxes in soils, Glob. Change Biol., 15, 808–824, 2009.
- Borken, W., Davidson, E., Savage, K., Gaudinski, J., and Trumbore, S. E.: Drying and wetting effects on carbon dioxide release from organic horizons, Soil Sci. Soc. Am. J., 67, 1888–1896, 2003.
- Bouma, T. J. and Bryla, D. R.: On the assessment of root and soil respiration for soils of different textures: interactions with soil moisture contents and soil CO₂ concentrations, Plant Soil, 227, 215–221, 2000.
- Brennan, A.: Vegetation and Climate Change Alter Ecosystem Carbon Losses at the Prairie Heating and CO₂ Enrichment Experiment in Wyoming, Department of Botany, University of Wyoming, 2013.
- Cable, J. M., Ogle, K., Williams, D. G., Weltzin, J. F., and Huxman, T. E.: Soil texture drives responses of soil respiration to precipitation pulses in the Sonoran Desert: implications for climate change, Ecosystems, 11, 961–979, 2008.
- Cable, J. M., Ogle, K., Lucas, R. W., Huxman, T. E., Loik, M. E., Smith, S. D., Tissue, D. T., Ewers, B. E., Pendall, E., and Welker, J. M.: The temperature responses of soil respiration in deserts: a seven desert synthesis, Biogeochemistry, 103, 71–90, 2011.
- Cable, J. M., Ogle, K., Barron-Gafford, G. A., Bentley, L. P., Cable, W. L., Scott, R. L., Williams, D. G., and Huxman, T. E.: Antecedent conditions influence soil respiration differences in shrub and grass patches, Ecosystems, 16, 1230–1247, 2013.
- Carrillo, Y. and Pendall, E.: Shifted role of bacteria in soil carbon cycling under future climate conditions: implications for soil carbon stocks, in review, 2018.
- Carrillo, Y., Dijkstra, F. A., LeCain, D., Morgan, J. A., Blumenthal, D., Waldron, S., and Pendall, E.: Disentangling root responses to climate change in a semiarid grassland, Oecologia, 175, 699– 711, 2014a.
- Carrillo, Y., Dijkstra, F. A., Pendall, E., LeCain, D., and Tucker, C.: Plant rhizosphere influence on microbial C metabolism: the role of elevated CO₂, N availability and root stoichiometry, Biogeochemistry, 117, 229–240, 2014b.
- Cerling, T. E.: The stable isotopic composition of modern soil carbonate and its relationship to climate, Earth Planet. Sc. Lett., 71, 229–240, 1984.
- Chou, W. W., Silver, W. L., Jackson, R. D., Thompson, A. W., and Allen-Diaz, B.: The sensitivity of annual grassland carbon cycling to the quantity and timing of rainfall, Glob. Change Biol., 14, 1382–1394, 2008.
- Cox, P. M.: Description of the TRIFFID dynamic global vegetation model, 2001.
- Davidson, E. A. and Janssens, I. A.: Temperature sensitivity of soil carbon decomposition and feedbacks to climate change, Nature, 440, 165–173, 2006.
- Davidson, E. A., Savage, K. E., Trumbore, S. E., and Borken, W.: Vertical partitioning of CO₂ production within a temperate forest soil, Glob. Change Biol., 12, 944–956, 2006.
- Davidson, E. A., Samanta, S., Caramori, S. S., and Savage, K.: The Dual Arrhenius and Michaelis–Menten kinetics model for decomposition of soil organic matter at hourly to seasonal time scales, Glob. Change Biol., 18, 371–384, 2012.

- Fang, C. and Moncrieff, J. B.: A model for soil CO₂ production and transport 1, Model development, Agr. Forest Meteorol., 95, 225–236, 1999.
- Friedlingstein, P., Andrew, R. M., Rogelj, J., Peters, G., Canadell, J. G., Knutti, R., Luderer, G., Raupach, M. R., Schaeffer, M., and Van Vuuren, D. P.: Persistent growth of CO₂ emissions and implications for reaching climate targets, Nat. Geosci., 7, 709– 715, 2014.
- Haberman, R.: Elementary applied partial differential equations, 3rd Edn., Prentice Hall Englewood Cliffs, NJ, 1998.
- Hanson, P., Edwards, N., Garten, C., and Andrews, J.: Separating root and soil microbial contributions to soil respiration: a review of methods and observations, Biogeochemistry, 48, 115– 146, 2000.
- Hashimoto, S., Carvalhais, N., Ito, A., Migliavacca, M., Nishina, K., and Reichstein, M.: Global spatiotemporal distribution of soil respiration modeled using a global database, Biogeosciences, 12, 4121–4132, https://doi.org/10.5194/bg-12-4121-2015, 2015.
- Hillel, D.: Environmental soil physics: Fundamentals, applications, and environmental considerations, Academic press, 1998.
- Hui, D. and Luo, Y.: Evaluation of soil CO₂ production and transport in Duke Forest using a process-based modeling approach, Global Biogeochem. Cy., 18, GB4029, https://doi.org/10.1029/2004GB002297, 2004.
- Huxman, T. E., Snyder, K. A., Tissue, D., Leffler, A. J., Ogle, K., Pockman, W. T., Sandquist, D. R., Potts, D. L., and Schwinning, S.: Precipitation pulses and carbon fluxes in semiarid and arid ecosystems, Oecologia, 141, 254–268, 2004.
- Jarvis, P., Rey, A., Petsikos, C., Wingate, L., Rayment, M., Pereira, J., Banza, J., David, J., Miglietta, F., and Borghetti, M.: Drying and wetting of Mediterranean soils stimulates decomposition and carbon dioxide emission: the "Birch effect", Tree Physiol., 27, 929–940, 2007.
- Jasoni, R. L., Smith, S. D., and Arnone, J. A.: Net ecosystem CO₂ exchange in Mojave Desert shrublands during the eighth year of exposure to elevated CO₂, Glob. Change Biol., 11, 749–756, 2005.
- Kayler, Z. E., Sulzman, E. W., Rugh, W. D., Mix, A. C., and Bond, B. J.: Characterizing the impact of diffusive and advective soil gas transport on the measurement and interpretation of the isotopic signal of soil respiration, Soil Biol. Biochem., 42, 435–444, 2010.
- Lee, X., Wu, H. J., Sigler, J., Oishi, C., and Siccama, T.: Rapid and transient response of soil respiration to rain, Glob. Change Biol., 10, 1017–1026, 2004.
- Lloyd, J. and Taylor, J.: On the temperature dependence of soil respiration, Funct. Ecol., 8, 315–323, 1994.
- Luo, Y. and Zhou, X.: Soil respiration and the environment, Academic press, 2010.
- Maggi, F. and Riley, W. J.: Transient competitive complexation in biological kinetic isotope fractionation explains nonsteady isotopic effects: Theory and application to denitrification in soils, J. Geophys. Res., 114, G04012, https://doi.org/10.1029/2008JG000878, 2009
- Mathworks: The MathWorks, Inc. Natick, Massachusetts, USA MATLAB and Statistics Toolbox Release, 2016.
- Meisner, A., Bååth, E., and Rousk, J.: Microbial growth responses upon rewetting soil dried for four days or one year, Soil Biol. Biochem., 66, 188–192, 2013.

- Millington, R. J. and Quirk, J. P.: Permeability of porous solids, T. Faraday Soc., 57, 1200–1207, 1961.
- Moldrup, P., Olesen, T., Yoshikawa, S., Komatsu, T., and Rolston, D. E.: Three-porosity model for predicting the gas diffusion coefficient in undisturbed soil, Soil Sci. Soc. Am. J., 68, 750–759, 2004.
- Morgan, J. A., LeCain, D. R., Pendall, E., Blumenthal, D. M., Kimball, B. A., Carrillo, Y., Williams, D. G., Heisler-White, J., Dijkstra, F. A., and West, M.: C4 grasses prosper as carbon dioxide eliminates desiccation in warmed semi-arid grassland, Nature, 476, 202–205, 2011.
- Moyes, A. B., Gaines, S. J., Siegwolf, R. T., and Bowling, D. R.: Diffusive fractionation complicates isotopic partitioning of autotrophic and heterotrophic sources of soil respiration, Plant Cell Environ., 33, 1804–1819, 2010.
- Mueller, K. E., Blumenthal, D. M., Pendall, E., Carrillo, Y., Dijkstra, F. A., Williams, D. G., Follett, R. F., and Morgan, J. A.: Impacts of warming and elevated CO₂ on a semi-arid grassland are non-additive, shift with precipitation, and reverse over time, Ecol. Lett., 19, 956–966, 2016.
- Nickerson, N. and Risk, D.: Physical controls on the isotopic composition of soil-respired CO₂, J. Geophys. Res., 114, G01013, https://doi.org/10.1029/2008JG000766, 2009.
- Ogle, K. and Pendall, E.: Isotope partitioning of soil respiration: A Bayesian solution to accommodate multiple sources of variability, J. Geophys. Res.-Biogeo., 120, 221–236, 2015.
- Ogle, K., Wolpert, R. L., and Reynolds, J. F.: Reconstructing plant root area and water uptake profiles, Ecology, 85, 1967–1978, 2004.
- Ogle, K., Barber, J. J., Barron-Gafford, G. A., Bentley, L. P., Young, J. M., Huxman, T. E., Loik, M. E., and Tissue, D. T.: Quantifying ecological memory in plant and ecosystem processes, Ecol. Lett., 18, 221–235, 2015.
- Ogle, K., Ryan, E., Dijkstra, F. A., and Pendall, E.: Quantifying and reducing uncertainties in estimated soil CO₂ fluxes with hierarchical data-model integration, J. Geophys. Res.-Biogeo., 121, 2935–2948, 2016.
- Oikawa, P., Grantz, D., Chatterjee, A., Eberwein, J., Allsman, L., and Jenerette, G.: Unifying soil respiration pulses, inhibition, and temperature hysteresis through dynamics of labile soil carbon and O₂, J. Geophys. Res.-Biogeo., 119, 521–536, 2014.
- Oleson, K., Lawrence, D., Bonan, G., Drewniak, B., Huang, M., Koven, C., Levis, S., Li, F., Riley, W., and Subin, Z.: Technical description of version 4.5 of the Community Land Model (CLM), Ncar Tech. Note NCAR/TN-503+ STR, National Center for Atmospheric Research, Boulder, CO, 422 pp., https://doi.org/10.5065/D6RR1W7M, 2013.
- Pendall, E., Del Grosso, S., King, J., LeCain, D., Milchunas, D., Morgan, J., Mosier, A., Ojima, D., Parton, W., and Tans, P.: Elevated atmospheric CO₂ effects and soil water feedbacks on soil respiration components in a Colorado grassland, Global Biogeochem. Cy., 17, 1046, https://doi.org/10.1029/2001GB001821, 2003.
- Pendall, E., Bridgham, S., Hanson, P. J., Hungate, B., Kicklighter, D. W., Johnson, D. W., Law, B. E., Luo, Y., Megonigal, J. P., and Olsrud, M.: Below-ground process responses to elevated CO₂ and temperature: A discussion of observations, measurement methods, and models, New Phytol., 162, 311–322, 2004.

- Pendall, E., Heisler-White, J. L., Williams, D. G., Dijkstra, F. A., Carrillo, Y., Morgan, J. A., and LeCain, D. R.: Warming reduces carbon losses from grassland exposed to elevated atmospheric carbon dioxide, PloS ONE, 8, e71921, https://doi.org/10.1371/journal.pone.0071921, 2013.
- Penman, H. L.: Gas and vapor movements in soil: The diffusion Soil of vapors through porous solids, J. Agric. Sci., 30, 437–462, 1940.
- Piao, S., Ciais, P., Friedlingstein, P., de Noblet-Ducoudre, N., Cadule, P., Viovy, N., and Wang, T.: Spatiotemporal patterns of terrestrial carbon cycle during the 20th century, Global Biogeochem. Cy., 23, GB4026, https://doi.org/10.1029/2008GB003339, 2009.
- Richards, L. A.: Capillary conduction of liquids through porous mediums, Physics, 1, 318–333, 1931.
- Risk, D., Kellman, L., and Beltrami, H.: A new method for in situ soil gas diffusivity measurement and applications in the monitoring of subsurface CO₂ production, J. Geophys. Res.-Biogeo., 113, G02018, https://doi.org/10.1029/2007JG000445, 2008.
- Risk, D., Nickerson, N., Creelman, C., McArthur, G., and Owens, J.: Forced Diffusion soil flux: A new technique for continuous monitoring of soil gas efflux, Agr. Forest Meteorol., 151, 1622– 1631, 2011.
- Risk, D., Nickerson, N., Phillips, C., Kellman, L., and Moroni, M.: Drought alters respired δ^{13} CO₂ from autotrophic, but not heterotrophic soil respiration, Soil Biol. Biochem., 50, 26–32, 2012.
- Ryan, E. and Ogle, K.: Inputs folder contents for DETECT model v1.0 [Data set], Zenodo, https://doi.org/10.5281/zenodo.926064, 2017a.
- Ryan, E. and Ogle, K.: Source code for running DETECT model v1.0. Zenodo, https://doi.org/10.5281/zenodo.927501, 22 September 2017b.
- Ryan, E. M., Ogle, K., Zelikova, T. J., LeCain, D. R., Williams, D. G., Morgan, J. A., and Pendall, E.: Antecedent moisture and temperature conditions modulate the response of ecosystem respiration to elevated CO₂ and warming, Glob. Change Biol., 21, 2588–2602, 2015.
- Sala, O., Lauenroth, W., and Parton, W.: Long-Term Soil Water Dynamics in the Shortgrass Steppe, Ecology, 73, 1175–1181, 1992.
- Schwinning, S., Sala, O. E., Loik, M. E., and Ehleringer, J. R.: Thresholds, memory, and seasonality: understanding pulse dynamics in arid/semi-arid ecosystems, Oecologia, 141, 191–193, 2004.
- Šimůnek, J. and Suarez, D. L.: Modeling of carbon dioxide transport and production in soil: 1. Model development, Water Resour. Res., 29, 487–497, 1993.
- Šimůnek, J., Van Genuchten, M. T., and Sejna, M.: The HYDRUS-1D software package for simulating the one-dimensional movement of water, heat, and multiple solutes in variably-saturated media, University of California-Riverside Research Reports, 3, 1–240, 2005.
- Šimůnek, J., van Genuchten, M. T., and Šejna, M.: Development and applications of the HYDRUS and STANMOD software packages and related codes, Vadose Zone J., 7, 587–600, 2008.
- Šimůnek, J., Van Genuchten, M. T., and Šejna, M.: HYDRUS: Model use, calibration, and validation, Trans. Asabe, 55, 1261– 1274, 2012.
- Sitch, S., Huntingford, C., Gedney, N., Levy, P., Lomas, M., Piao, S., Betts, R., Ciais, P., Cox, P., and Friedlingstein, P.: Evalua-

tion of the terrestrial carbon cycle, future plant geography and climate-carbon cycle feedbacks using five Dynamic Global Vegetation Models (DGVMs), Glob. Change Biol., 14, 2015–2039, 2008.

- Sponseller, R. A.: Precipitation pulses and soil CO₂ flux in a Sonoran Desert ecosystem, Glob. Change Biol., 13, 426–436, 2007.
- Tang, J., Baldocchi, D. D., Qi, Y., and Xu, L.: Assessing soil CO₂ efflux using continuous measurements of CO₂ profiles in soils with small solid-state sensors, Agr. Forest Meteorol., 118, 207–220, 2003.
- Thomas, A. D., Hoon, S. R., and Linton, P. E.: Carbon dioxide fluxes from cyanobacteria crusted soils in the Kalahari, Appl. Soil Ecol., 39, 254–263, 2008.
- Todd-Brown, K. E., Hopkins, F. M., Kivlin, S. N., Talbot, J. M., and Allison, S. D.: A framework for representing microbial decomposition in coupled climate models, Biogeochemistry, 109, 19–33, 2012.
- Tucker, C. L., Bell, J., Pendall, E., and Ogle, K.: Does declining carbon-use efficiency explain thermal acclimation of soil respiration with warming?, Glob. Change Biol., 19, 252–263, 2013.

- Vargas, R., Baldocchi, D. D., Allen, M. F., Bahn, M., Black, T. A., Collins, S. L., Yuste, J. C., Hirano, T., Jassal, R. S., and Pumpanen, J.: Looking deeper into the soil: biophysical controls and seasonal lags of soil CO₂ production and efflux, Ecol. Appl., 20, 1569–1582, 2010.
- Vargas, R., Carbone, M. S., Reichstein, M., and Baldocchi, D. D.: Frontiers and challenges in soil respiration research: from measurements to model-data integration, Biogeochemistry, 102, 1– 13, 2011.
- Xiang, S.-R., Doyle, A., Holden, P. A., and Schimel, J. P.: Drying and rewetting effects on C and N mineralization and microbial activity in surface and subsurface California grassland soils, Soil Biol. Biochem., 40, 2281–2289, 2008.
- Xu, L., Baldocchi, D. D., and Tang, J.: How soil moisture, rain pulses, and growth alter the response of ecosystem respiration to temperature, Global Biogeochem. Cy., GB4002, https://doi.org/10.1029/2004GB002281, 2004.
- Zelikova, T. J., Williams, D. G., Hoenigman, R., Blumenthal, D. M., Morgan, J. A., and Pendall, E.: Seasonality of soil moisture mediates responses of ecosystem phenology to elevated CO₂ and warming in a semi-arid grassland, J. Ecol., 103, 1119–1130, 2015.