

In silico analysis of carbon ~~stabilisation by plant and soil microbes~~ ~~for different weather scenarios~~ water dynamics in the rhizosphere under drought conditions

Mona Giraud^{*,1,2}, Ahmet Kürşad Sircan^{*,3}, Thilo Streck³, Daniel Leitner¹, Guillaume Lobet⁴, Holger Pagel^{*,1,2}, and Andrea Schnepf^{*,1,2}

*These authors contributed equally to this work

¹Institute of Bio- and Geosciences (IBG-3), Forschungszentrum Jülich, 52428, Jülich, Germany

²Institute of Crop Science and Resource Conservation, University of Bonn, 53115 Bonn, Germany

³University of Hohenheim, Institute of Soil Science and Land Evaluation, Department of Biogeophysics, Germany

⁴Earth and Life Institute, Université catholique de Louvain, Belgium

Correspondence: Mona Giraud (m.giraud@fz-juelich.de)

Abstract. A plant's development is strongly linked to the water and carbon (C) flows in the soil-plant-atmosphere continuum. Ongoing climate shifts will alter the water and C cycles and affect plant phenotypes. Comprehensive models that simulate mechanistically and dynamically the feedback loops between water and C fluxes in the soil-plant system are useful tools to evaluate the sustainability of genotype-environment-management combinations that do not yet exist. In this study, we present the equations and implementation of a rhizosphere-soil model within the CPlantBox framework, a functional-structural plant model that represents plant processes and plant-soil interactions. The multi-scale plant-rhizosphere-soil coupling scheme previously used for CPlantBox was likewise updated, among others to ~~include an implicit time-stepping~~ increase the accuracy and stability of the model outputs. The model was implemented to simulate the effect of dry spells occurring at different plant development stages, and for different soil ~~biokinetic parametrisations~~ kinetic parameterisations of microbial dynamics in soil. We could observe diverging results according to the date of occurrence of the dry spells and soil ~~parametrisations~~ parameterisations. For instance, ~~an earlier dry spell~~ earlier dry spells (from 11th to the 18th day of growth) led to a lower cumulative plant C release, while later dry spells (from 18th to the 25th day of growth) led to higher C input to the soil. For more reactive microbial communities (higher maximum C uptake rate and (de)activation rates), this higher C input caused a strong increase in CO₂ emissions, ~~while, for,~~ For the same weather scenario, we observed ~~a lasting stabilisation of soil C~~ lower microbial CO₂ emissions with less reactive communities. This model can be used to gain insight into C and water flows at the plant scale, and the influence of soil-plant interactions on C cycling in soil.

1 Introduction

Understanding the dynamics driving the water and carbon (C) balances in the soil-plant-atmosphere continuum is key to predict, mitigate, and adapt agroecosystems to climate change (Silva and Lambers, 2021; George et al., 2024). ~~While water plays an~~

20 ~~important role as a resource and as an intra- and inter-domain signalling carrier (De Swaef et al., 2022), C compounds can serve as an essential energy source or a biomass building block (Lynch et al., 2021; George et al., 2024).~~

Plants link the water and C balances in the soil and atmosphere via plant-related processes, such as coupled stomatal regulation-photosynthesis-transpiration ~~-, C and water transport and storage in plant tissues, C usage (growth, maintenance, exudation) root-soil water exchanges and root C release~~ (De Swaef et al., 2022; Lacoïnte and Minchin, 2019; Silva and
25 Lambers, 2021). ~~The plant's capacity to capture, transport and release water (resp. C) across domains (e.g., C fixation leading to rhizodeposit hotspots distribution (Landl et al., 2021a)) or within one domain (e.g., hydraulic lift, (Lynch et al., 2021)) is also strongly defined by~~ the topology of its shoot (de la Fuente Cantó et al., 2020) and root system (Galindo-Castañeda et al., 2023; George et al., 2024; de la Fuente Cantó et al., 2020) ~~e.g., lateral root branching density, number of root tips (Lynch, 2021; Galindo-Castañeda et al., 2023). Likewise, plant anatomical characteristics (e.g. as well as plant anatomical characteristics e.g.~~
30 ~~number and nature of the concentric root cells, number and size of the phloem and xylem vessels) strongly affect the flow within the plant (Lynch et al., 2021) and the plant-soil exchanges (Galindo-Castañeda et al., 2023; Lynch et al., 2021).~~ (Lynch et al., 2021; Galindo-Castañeda et al., 2023; Lynch et al., 2021).

Plant-soil exchanges lead to the creation of the rhizosphere, i.e., root-affected soil with characteristics distinctive from the bulk soil, such as microbial activity and composition, soil structure, organic carbon, or water and nutrient content (George et al., 2024; de la
35 The quantity and quality of the C released by the plant in the rhizosphere influence the transfer of nutrients and energy to the soil (de la Fuente Cantó et al., 2020) and thus microbial activity (Schnepf et al., 2022; Galindo-Castañeda et al., 2023, 2022; de la Fuente Cantó et al., 2020) directly as energy and biomass input and indirectly by changing the properties of the soil (Schnepf et al., 2022; de la Fuente Cantó et al., 2020; George et al., 2024). Root water uptake changes the soil moisture (Ruiz et al., 2021), which eventually triggers drought stress responses of soil microbial communities (Bardgett and Caruso, 2020).
40 ~~Plant-soil feedbacks drive inter- and intra-specific plant interactions (Putten et al., 2013) (e.g., competition for soil resources). Plants can also exert direct feedback effects on themselves (e.g., aboveground-belowground feedback) which will influence their fitness. Those feedbacks depend on soil organic matter (SOM) content and texture, which determine soil water holding capacity and hence moisture availability under drought (Bardgett and Caruso, 2020; George et al., 2024).~~

~~Plant-soil exchanges lead to the creation of the rhizosphere, an area of soil around the root with biotic (rhizodeposition, microbial communities) and abiotic (structure, water content) characteristics distinctive from the bulk soil (George et al., 2024; de la Fuente~~
45 ~~. The hydraulic conductance of the rhizosphere is the primary constraint for plant water transport, especially in water-limited environments (De Swaef et al., 2022; Khare et al., 2022; Roose et al., 2016), leading to abiotic stress and stomatal closure (George et al., 2024) or better drought resilience in case of favorable abiotic conditions (Lynch et al., 2021).~~

The rhizosphere is also the most important hotspot of microbial activity on the planet (George et al., 2024; Lynch et al.,
50 2021). Soil microbes control SOM decomposition and CO₂ release (Pagel et al., 2020) and regulate soil C allocation to the different organic and mineral pools. ~~Rhizosphere microbes are relevant to soil C cycling as the activation-deactivation processes allow for rapid responses to the input of C and energy provided by rhizodeposits (George et al., 2024). Soil microbes are therefore a significant sink of photoassimilates, influencing root C release, root-to-shoot C allocation and overall plant growth (Lynch et al., 2021).~~

55 Due to the tight coupling of water and C cycling between soil, microbes and plant, ~~it becomes relevant to design integrated plant-microbiome traits adapted for specific soils to promote resource capture by plants and ensure the resilience of plant-soil systems to environmental stressors (Galindo-Castañeda et al., 2022; de la Fuente Cantó et al., 2020; Silva and Lambers, 2021) -A thorough understanding of interactions between soil-microbes-plant (Galindo-Castañeda et al., 2022; Bardgett et al., 2013; Roose et al., that can lead to intra- and inter-specific competition (for light, water and nutrients), complementarity (spatio-temporal and niche processes) and facilitation (improving access to resources)~~ a thorough understanding of the interactions between these domains is therefore essential to inform breeding choices (De Swaef et al., 2022; Gaudio et al., 2021).

Mechanistic biogeochemical modelling facilitates the analysis of complex systems (Schnepf et al., 2022; Putten et al., 2013). Especially Functional Structural Plant Models (FSPMs) link the plant structure with its function and simulate processes at the plant level (Schnepf et al., 2022; Ruiz et al., 2021). ~~Coupling an FSPM with a soil model in a multiscale modelling framework leverages the strengths of different models and increase model accuracy (Schnepf et al., 2022; Roose et al., 2016; Gaudio et al., 2021) -Small-scale models provide effective properties and predict emerging behaviour at higher scales, while models at coarser scales provide the initial and boundary conditions to the small scale models (Schnepf et al., 2022)-~~

~~Although many~~ Several studies looked at water and flow of (usually one) nutrient (especially phosphorus) in soil and rhizosphere at the plant scale (Mai et al., 2019; Ruiz et al., 2021; Roose et al., 2016), ~~few focused on both water and C (Ruiz et al., 2021) and accounted for physiochemical processes (Pot et al., 2022). Moreover, many of the spatially explicit rhizosphere models ignore microorganisms (Roose et al., 2016) although the increasing resolution of those models make a detailed characterisation of the soil biochemistry more relevant (Roose et al., 2016; Lynch et al., 2021). FSPMs simulating C (Rees et al., 2020; ?) and water flow were also developed (Lacointe and Minchin, 2019; Giraud et al., 2023) but simulating neither the soil C nor water flows (Lacointe and Minchin, 2019; Rees et al., 2020; ?) or only the water flow (Giraud et al., 2023) -~~ Specifically, in their studies, Rees et al. (2025) and Lacointe and Minchin (2019) simulated the plant C allocation and root C releases, while Landl et al. (2021b) linked carbon exudation with a 3 dimensional root system and a soil model to evaluate the 3D distribution and interactions of the rhizospheres. However, no plant model coupled mechanistic inner plant water and carbon flows with a 3 dimensional soil, representing the rhizosphere and microbial processes, enabling a more precise analysis of the effects of a plant's processes and structure on the rhizosphere C fluxes.

80 ~~This study aimed to set up and implement a fully integrated plant-soil-microbiome model that accounts for root-microbiome interactions directly and enables the in-depth analysis of drought responses of C cycling in plant-soil systems. We present~~ To fill this gap, we set up a multi-scale multi-physics and multi-domain model. The plant domain is represented using an extended version of the FSPM CPlantBox (Giraud et al., 2023). CPlantBox is a 3D functional structural full plant model and was used 85 to model several processes across the plant, single root, ~~ans and~~ and pore scales (Mai et al., 2019; Landl et al., 2021a; Giraud et al., 2023). ~~Root system simulations were also implemented to evaluate the effect of root system structure on the root-soil exchange of water (Khare et al., 2022), C (Landl et al., 2021a), or water and phosphorous (Mai et al., 2019)~~ This FSPM stands out by its unique capacity to represent the inner plant carbon and water flow as well as several carbon- and water-dependent processes, such as carbon assimilation, plant growth, mucilage release and exudation. This set of processes allow for new insights on the

90 effects of different environment on the plant's carbon allocation. We coupled CPlantBox with ~~TraiRhizo (Sircan et al., 2025)~~
the model of Sircan et al. (2025), a 1D axisymmetric soil model ~~based. TraiRhizo offers a unique~~. This soil model offers a
balance between complexity and precision for the C and microbial ~~dynamic~~ dynamics and a simplified representation of the
soil structure (see review of Pot et al. (2022) for an evaluation of the analogous SpatC (Pagel et al., 2020) against other soil
models). ~~For~~ The model of Sircan et al. (2025) was developed parallel to the CPlantBox model, to ensure a good integration
95 within the plant model (e.g., number of C pools represented, interaction with water). It was therefore possible to use the net
plant water and carbon uptake computed by CPlantBox as input for the soil model. Moreover, for this study, ~~TraiRhizo~~ the
model of Sircan et al. (2025) was adapted and re-implemented in DuMu^x (Koch et al., 2021) for a faster and more adaptable
simulation setup. The two models were coupled using an updated version of the multiscale setup of Mai et al. (2019) to obtain
higher accuracy and stability. The resulting framework was used to conduct an in-depth analysis of the root-derived C dynamics
100 in the rhizosphere under drought conditions against baseline scenarios. We evaluated three hypotheses: (1) dry spells will lead
to a lower release of plant carbon and thus to (2) lower microbial growth in the rhizosphere, and (3) these changes in soil
carbon input and loss can lead to both an increase or decrease in short-term soil carbon storage.

In the first part of the paper, we present the equations of the model, with emphasis on the soil processes. We then present
the scheme for the coupling of the different domains and modules. In the second section of the paper, we implement the model
105 to evaluate how short dry spells at different stages of the plant's development affect the soil C balance and, in particular, the
C ~~stabilisation~~ fluxes according to the ~~biokinetic~~ kinetic characteristics of the microbial community. In the last section of the
paper, we discuss the output of the coupled model against experimental and *in silico* results, and we compare our framework
against other existing simulation tools.

2 Material and methods

110 2.1 Overview of the multiscale framework

2.1.1 Individual models

For the plant domain, we use the FSPM CPlantBox, as presented by Giraud et al. (2023). In CPlantBox, the plant is represented
as a series of connected 1-dimensional segments, making a branched network of nodes in a 3-dimensional space (Fig. 1(a)).
CPlantBox simulates the plant water flow using the analytical solution of (Meunier et al., 2017), and the ~~coupled the resulting~~
115 stomatal regulation (Leuning, 1995) and FvCB-photosynthesis (Farquhar et al., 1980) models. The plant C balance is evaluated
with a numerical solver (Lacointe and Minchin, 2019), which yields the semi-mechanistic plant growth rate.

For the soil domain, Sircan et al. (2025) developed a 1-dimensional axisymmetric soil model that computes the C transport and
microbial dynamics according to a set C release rate from a root segment. Their model was re-implemented within the existing
soil framework of the FSPM (Mai et al., 2019) and adapted to take into account variations of water content. It was expanded
120 to represent ~~the as well a~~ 3-dimensional macroseale soil (Fig. 2) (Khare et al., 2022; Mai et al., 2019; Koch et al., 2021).
~~The macroseale model is made of 3-dimensional sub-control volumes (i.e., voxels), while the microseale model is comprised~~

of axisymmetric 1D sub-control volumes (i.e., segments). A list of the symbols of the soil models, their units and meanings can be found in appendix A. ~~The explicitly represented soil~~ In the model, microbes are represented as two functional groups: the copiotrophs, characterized by rapid growth and high carbon uptake but lower carbon use efficiency; and the oligotrophs, characterized by slower growth and higher carbon use efficiency under carbon-limited conditions. Both microbial groups can be either active or dormant and emit CO₂ due to microbial respiration. See Table G1 for a complete list of the differing parameters between those two groups. The remaining represented soil organic C pools are: low molecular weight organic compounds (dissolved and sorbed) ~~, and high molecular weight organic compounds, copiotrophic and oligotrophic microorganisms with active and dormant metabolic states, emitted CO₂ due to microbial respiration.~~ The high molecular weight compounds are not affected by advection. In theory, the two inert soil organic pools (low- and high-molecular weight organic carbon) can be used to represent the fast- and slow-cycling soil organic carbon pools (Sircan et al., 2025). However, because of the calibration used for this study, both carbon types were used to represent the two faster decomposing carbon pools, while the remaining slower decomposing pool was not explicitly represented (see section 2.4.4). In the following section, we give an overview of the spatial and temporal coupling of these models within a multiscale framework.

The hydraulic conductance of the soil–root interface is the primary constraint for plant water transport when the soil is dry. Achieving realistic results requires, therefore, high spatial resolution near the soil-root interface, leading to computationally expensive simulations (Roose et al., 2016; Khare et al., 2022). To balance computing speed and model accuracy, we therefore coupled the models presented in this section in a multiscale framework where the spatial resolution is set according to the distance from the soil-root interface.

2.1.2 Spatial coupling

Figs. 1 and 2 give a graphical representation of the spatial coupling method, adapted from the approach presented by Mai et al. (2019) and Khare et al. (2022). Each root segment below-ground is allocated to a ~~macro~~scale-3D voxel, selected according to the location of the segment centre when it was first created. For each root segment, we initialise a ~~micro~~scale-1D model that computes the water flow, C transport, and microbial dynamics based on the plant-soil water and C exchanges prescribed by the root segment. Other than in Sircan et al. (2025), in which the plant-soil exchange was prescribed as boundary conditions at the root-soil interface, the net releases of water and C in this coupled model come from the simulation outputs of the FSPM, driven by inner plant flows and processes, such as photosynthesis, transpiration, and growth (Giraud et al., 2023).

The volume of the ~~macro~~scale-3D voxel is divided between the ~~micro~~scale domains-1D soil models it contains. Therefore, the maximum possible volume of a single ~~micro~~scale-represented implicitly by a single 1D soil domain is the volume of the ~~its~~ containing voxel. The total perirhizal volume around a root system (~~bulk soil~~) corresponds to the total volume of all the ~~micro~~scale-1D domains. It is also equal to the sum of the volumes of all ~~macro~~scale-3D voxels containing at least one root segment. Therefore, the ~~bulk soil~~ total perirhizal volume depends on the root system structure and on the user-defined resolution at the ~~macro~~scale-3D soil rather than an *a priori* evaluation of the rhizosphere volume. The microbial reactions and adsorption are computed at the voxel scale for voxels that do not contain root segments. The biochemical reactions are computed at the ~~micro~~scale-1D scale for voxels containing at least one segment. The resulting variation rates are represented at the ~~macro~~scale

in the 3D soil as a sink term. The flow of water and transport of solutes between the voxels is computed at the macroscale in the 3D soil and given as sink term to the microscale segments 1D soil models. Moreover, we scale the size of the microscale segments 1D sub-control volumes, meaning that the segments near the root surface are smaller than the ones further away. The macroscale 3D soil domain is divided into subdomains whose concentrations and flows are computed in parallel while the microscale 1D domains are distributed between processes run in parallel. The plant processes are not simulated in parallel.

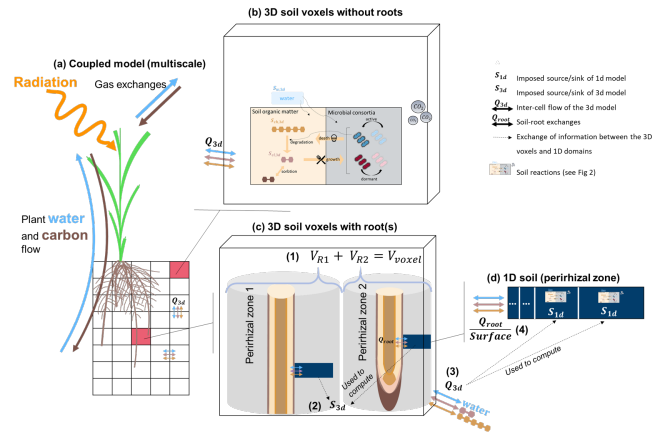


Figure 1. Representation of the multiscale coupling with (a) the whole multiscale model, (b) example of a 3D macroscale soil voxels voxel without roots, (c) example of a 3D soil voxel with two roots, (d) example of a 1D microscale model representing the whole perirhizal zone around a root segment. The 3D soil model (black grid) is made up of 3D voxels (transparent rectangles with black borders). The 1D model is represented using the dark blue line in the 3D voxels. It lies between a root segment and the end of the implicitly-represented perirhizal zone (grey hollow cylinder). (1) The sum of the volume of the perirhizal zones of the roots segments in a voxel is equal to the volume of the voxel. (2) The changes in content in the 1D models are used to define set sinks for the 3D voxels that contain at least one root. (3) The 3D flow entering or leaving a 3D voxel with roots is sent to the 1D models via sink terms in their sub-control volume. (4) The root releases and uptake are set as inner Neumann boundary conditions of the 1D soil models. The carbon-related reactions are computed in the voxels without roots and in the sub-control volumes of the 1D soils. More information on the reactions and the 1D models are given in Fig. 2.

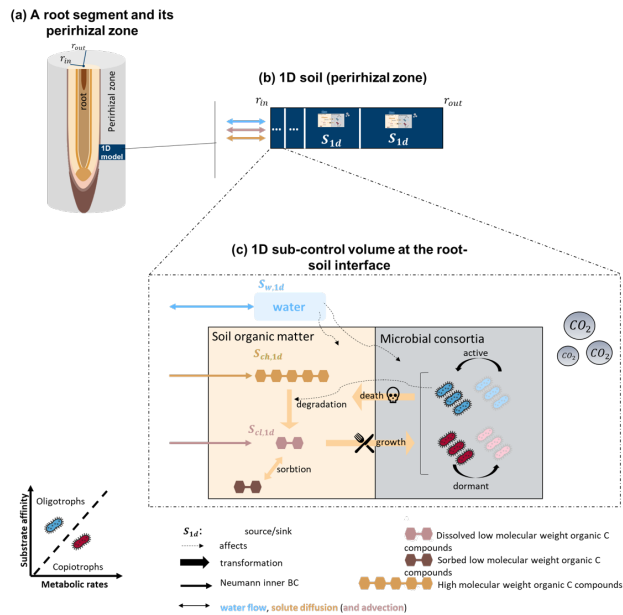


Figure 2. Representation of the updated microscale model Representation of the updated 1D soil model (Sircan et al., 2025) reimplemented within the multiscale framework (Mai et al., 2019; Koch et al., 2021). The model represents the soil domain around the root (perirhizal zone). Eight carbon pools are represented: carbon from organic compounds with a low molecular weight (dissolved and sorbed), organic compounds with a high molecular weight, active and dormant copiotrophs and oligotrophs, emitted microbial CO_2 . Water flow and content are also represented. Water affects the microbes (by impeding water stress) and the inert soil organic matter by lowering the dissolved carbon concentration. The graphic was adapted from Sircan et al. (2025). The same reactions are also computed in 3D soil voxels that do not contain any roots (see Fig. 1).

Fig. 3 gives a representation of the hard-coded direct interactions between the atmospheric conditions and the C and water pools represented in the plant and soil. For instance, for a set high soil water content, high transpiration leads to a lower plant water potential and, thus, to a lower growth. In contrast, growth has a direct positive effect on the microbial development, as it stops the onset of water stress (see Eqns.D10, D29, D30). However, high soil water content also leads to a larger root system and leaf surface, allowing for higher transpiration lower concentration of dissolved organic carbon. This lower dissolved carbon concentration was a negative effect on the microbial development by lowering the growth rate and making the microbes go into dormancy (see Eqns.D27, D28). It is the interactions between those processes that drive the water and C balances of the overall system. It is important to note that we represent the direct effects of temperature on plant processes but not on the microbial activity.

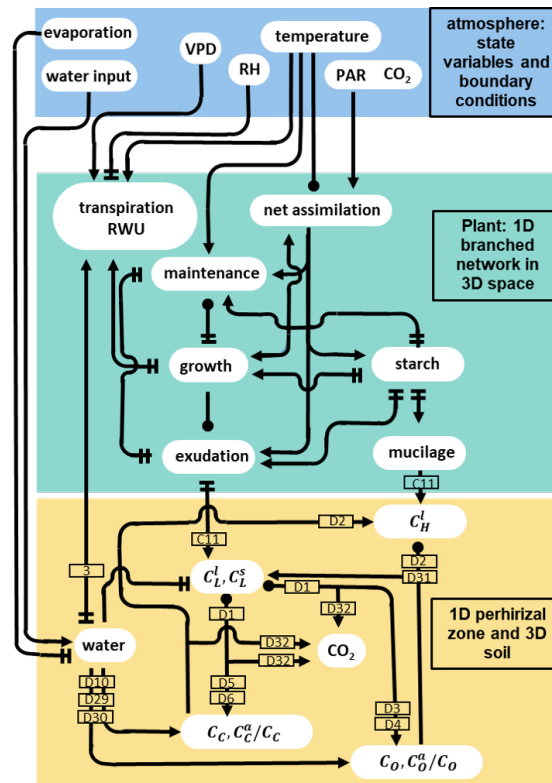


Figure 3. Graphical representation of the direct interactions represented between the different carbon and water pools plant model (green rectangle), soil models (yellow rectangle) and the atmospheric conditions given as input (blue rectangle). The lines ending with an arrow represent a positive effect, two perpendicular lines represent a negative effect, and a circle represents both a positive and negative effect (the net effect depends on the simulated conditions). The numbers in the rectangles give the indexes of the equations describing the corresponding process. The equations affecting the plant domain are defined in Giraud et al. (2023). The atmospheric conditions are represented by user-defined state variables and boundary conditions, e.g. vapour pressure deficit (VPD), relative humidity (RH), and incoming photosynthetically active radiations (PAR). Soil carbon pool data include the concentration of carbon from low molecular weight organic compounds in the soil and water phase (resp. C_L^s , C_L^l), high molecular weight organic compounds in the water phase $C(C_H^l, C_H)$, total or active copiotrophs (resp. C_C , C_C^a) and oligotrophs (resp. C_O , C_O^a).

170 2.1.3 Temporal coupling

The modules of the plant and soil models are implemented sequentially. The order of the implementation is presented in Fig. 4. Each "outer" loop (encompassing all the modules) represents one time-step. Within each time step, two nested fixed-point iterations are implemented:

1. the "first fixed point iteration" contains the interactions between the [a] soil water flow and solute transport (for the **microscale and macroscale-1D and 3D** domains) and [b] plant water flow and stomatal regulation.

175

2.2 Governing Equations

These sections present the method used to simulate water and carbon turnover within the plant and the soil. Each section is divided between the plant-specific and soil-specific equations. For the soil part, the common equations for both the **macro- and microscale-3D and 1D soil models** are presented, followed by equations specific to each domain. The soil volume (volume of the solid and pores) is written in cm^3 *scv* (sub-control volume). The amounts of solutes and microbes are given in units of equivalent carbon (*mol C*). As the equations describing the microbial reactions remain similar to those set up by Sircan et al. (2025), these ordinary differential equations are included in appendix D.

2.2.1 Plant water flow

The plant water model is described in Giraud et al. (2023). Briefly, we compute the plant xylem water flow as:

$$205 \quad -K_{ax,x} \frac{\partial^2 \psi_{h,x}}{\partial \xi^2} = -p_{seg} \times q_{w,plant-out} \text{ for } \Omega \setminus \partial\Omega, \quad (1)$$

$$\frac{\partial \psi_{h,x}}{\partial \xi} = 0 \text{ for } \partial\Omega \text{ (all organ end-nodes)} \quad (2)$$

$K_{ax,x}$ ($cm^4 hPa^{-1} d^{-1}$) is the xylem intrinsic axial conductance and ξ (cm) is the local axial coordinate of the plant. p_{seg} (cm) is the perimeter of the plant exchange zone, Ω the domain considered (here the plant), and $\partial\Omega$, the domain's boundary (here: the tip or end-node of each organ). We assume a no-flux boundary condition here as the plant-exterior water exchanges occur along the segments, therefore the equation given above is valid for the whole domain, except the boundaries $\Omega \setminus \partial\Omega$.

The net water sink is given by the lateral water flow ($q_{w,plant-out}$ in $cm d^{-1}$), which corresponds to the xylem-soil ($q_{w,root-soil}$) or xylem-atmosphere ($q_{w,leaf-air}$) water exchange:

$$q_{w,plant-out} = \begin{cases} q_{w,root-soil} = \min(K_{lat,x}, K_{sri})(\psi_{h,x} - \psi_{m,1DS}) & , \text{ if root} \\ q_{w,leaf-air} = K_{lat,min}(\psi_{h,x} - \psi_{h,ox}) & , \text{ if leaf blade} \end{cases} \quad (3)$$

With $K_{lat,x}$ ($cm hPa^{-1} d^{-1}$) the lateral hydraulic conductivity of the root cortex (for roots) or of the vascular bundle-sheath (for leaves). K_{sri} ($cm hPa^{-1} d^{-1}$) is the equivalent lateral hydraulic conductivity and $\psi_{m,1DS}$ (hPa) is the soil matric potential at the inner boundary of the corresponding 1D **microscale**-soil domain (root-soil interface). $\psi_{h,ox}$ (hPa) is the water potential in the outer-xylem compartment, computed by the FvCB-stomatal regulation module. Note that, as changes in plant water storage are neglected, root water uptake equals transpiration at any given time step: $\sum_i^{segs} p_i q_{w,root-soil,i} = \sum_i^{nr} p_i q_{w,leaf-air,i}$, with p_i (cm) the perimeter of the segment i within the set *segs*.

220

K_{sri} is computed from the soil and plant data:

$$K_{sri} = \begin{cases} \frac{K_{sat} \kappa_m(S_{sat})}{\Delta r_{root-1DS}} \text{ if } q_{w,root-soil} \leq 0 \\ \frac{K_{sat}}{\Delta r_{root-1DS}} \text{ else} \end{cases}, \quad (4)$$

$$K_{sat} = \frac{\kappa}{\mu_w} \quad (5)$$

With K_{sat} ($cm\ hPa^{-1}\ d^{-1}$) the soil saturated hydraulic conductivity, μ_w ($hPa\ d$) the viscosity of the soil water solution, assumed equal to that of pure water at $20^\circ C$. κ (cm^2) is the intrinsic permeability of the soil and κ_m ($-$) the relative conductivity loss, dependent on the soil water saturation (S_{sat} , $-$). $\Delta r_{root-1DS}$ (cm) is the distance in radial coordinate between the root surface and the centre of the inner segment of the [microscale-1D soil](#) domain. Net root water release ($q_{w,root-soil} > 0$) is only limited by K_{sat} in the soil, while net root water uptake ($q_{w,root-soil} < 0$) is also limited by κ_m .

2.2.2 Soil water flow

The variation in volumetric soil water content (θ in $cm^3\ water/cm^3\ scv$) is defined according to the Richards equations (Richards, 1931):

$$\frac{\partial \theta}{\partial t} = \nabla \cdot (K(S_{sat})(\nabla \psi_{m,XDS} - \nabla \psi_{g,XDS})) + S_\theta \quad (6)$$

With $K(S_{sat})$ ($cm^2\ hPa^{-1}\ d^{-1}$) the unsaturated hydraulic conductivity (hereafter written as K for simplicity), $\nabla \psi_{m,XDS}$ and $\nabla \psi_{g,XDS}$ (hPa/cm) are the gradients of matric and gravitational water potentials in either the [macroscale-3D](#) (ψ_{3DS}) or [microscale-1D](#) (ψ_{1DS}) soil domain, respectively.

S_θ ($cm^3\ water\ cm^{-3}\ scv\ d^{-1}$) is the net source of water in the domain. ∇ is the nabla operator, either for the 1D axisymmetric ([radial derivative](#) $\frac{\partial}{\partial r}$) or for the 3D domain ($\frac{\partial}{\partial x} + \frac{\partial}{\partial y} + \frac{\partial}{\partial z}$) ([Schey, 1973](#)). We moreover have:

$$\theta = \phi S_{sat} \quad (7)$$

with ϕ ($cm^3\ pore/cm^3\ scv$) the soil porosity (assumed constant). [For](#) simplicity, we assume that $\phi = \theta_s$, with θ_s ($cm^3\ water/cm^3\ scv$) the θ value at saturation. θ_r ($cm^3\ water/cm^3\ scv$) is the lowest possible θ value (residual water content). The hydraulic conductivity is defined using the pore size distribution model of Mualem (1976) and the water retention function is computed according to van Genuchten (1980):

$$S_{sat} = \left(\frac{1}{1 + (\alpha \psi_{c,XDS})^n} \right)^{1-1/n} \quad (8)$$

$$\psi_{c,XDS} = \psi_{n,XDS} - \psi_{m,XDS} \quad (9)$$

$$K = \kappa_m(S_{sat})K_{sat} \quad (10)$$

$$\kappa_m(S_{sat}) = S_{sat}^{l_{soil}} \times [1 - (1 - (S_{sat})^{n/(n-1)})^{1-1/n}]^2 \quad (11)$$

ψ_n (hPa) is the reference pressure, [set to, for simplicity, at](#) $1000\ hPa$, and ψ_c (hPa) is the capillary pressure. α (cm^{-1}) is related to the inverse of the air entry suction, n ($-$) is a shape parameter, l_{soil} ($-$) is the soil tortuosity. The [implementation implementations](#) of Eq. (6) for the [macroscale-3D](#) soil and [microscale-1D](#) soil are given respectively in appendices B1 and B2.

2.2.3 Plant carbon transport

The plant sucrose model used for this study is based on the one presented by Giraud et al. (2023). The model was adapted to simulate mucilage release. The variation of sucrose-carbon concentration in the sieve tube (C_L^{st} in $mol C cm^{-3} st d^{-1}$, with $cm^3 st$ the sieve tube volume) is computed from Eq. (12):

$$255 \quad \frac{\partial C_L^{st}}{\partial t} + \frac{\partial}{\partial \xi} \left(C_L^{st} \times K_{ax,st} (RT_{seg} \frac{\partial C_L^{st}}{\partial \xi} + \frac{\partial \psi_{h,x}}{\partial \xi} + \frac{\partial \psi_{g,x}}{\partial \xi}) \right) - S_{exud,st} - S_{starch,st} - S_{other,st} = 0$$

$$\text{for } \Omega \setminus \partial\Omega, t > t_0, \quad (12)$$

$$K_{ax,st} (RT_{seg} \frac{\partial C_L^{st}}{\partial \xi} + \frac{\partial \psi_{h,x}}{\partial \xi} + \frac{\partial \psi_{g,x}}{\partial \xi}) = 0 \quad \text{for } \partial\Omega \quad (13)$$

$$260 \quad C_L^{st} = C_{L,0}^{st} \quad \text{for } t = t_0, \quad (14)$$

With ξ (cm) the local axial coordinate of the plant, R ($hPa cm^3 K^{-1} mol^{-1}$) the ideal gas constant, $K_{ax,st}$ ($cm^4 hPa^{-1} d^{-1}$) the sieve tube axial intrinsic conductance, T_{seg} (K) the temperature of the plant segment. $S_{exud,st}$ ($mol cm^{-3} st d^{-1}$) is the loss of C_L^{st} by exudation of low molecular weight organic carbon compounds, S_{starch} ($mol cm^{-3} st d^{-1}$) is the loss or gain of C_L^{st} by starch synthesis or degradation, $S_{other,st}$ ($mol cm^{-3} st d^{-1}$) represents other net sinks of C_L^{st} -like gain by assimilation and inflow from the mesophyll compartment.

We assume that the exudation of low molecular weight organic carbon occurs via a passive diffusion (Rakshit et al., 2021; Ma et al., 2022), represented by a mass transfer approach:

$$S_{exud,st} = q_{exud} \frac{2\pi r_{in} L}{V_{st}} \quad (15)$$

$$q_{exud} = K_{lat,st} (C_L^{st} - C_L^l) \quad (16)$$

270 With $K_{lat,st}$ ($cm d^{-1}$) is the lateral sieve tube conductivity for sucrose, V_{st} ($cm^3 st$) is the sieve tube volume. We assume that exudation occurs mainly near the root tip (Rakshit et al., 2021; Badri and Vivanco, 2009; Neumann and Römheld, 2009). Therefore, $K_{lat,st} > 0$ only for the last segment of growing roots. C_L^l ($mol C/cm^3 water$) corresponds to the concentration of dissolved low molecular weight organic carbon in the soil water at the root-soil interface. It is obtained from the inner segment of the [microscale-1D](#) soil domain assigned to the root segment. We assume that the mucilage is made from starch (*starch*, in $mol C/cm^3$) before being released via an active process (Rougier, 1981; Nguyen, 2009; Kutschera-Mitter et al., 1998). The production and release processes are represented in the following equations:

$$\frac{\partial starch}{\partial t} = S_{starch,st} - S_{mucil,st} \quad (17)$$

$$S_{starch,st} = k_{targ,starch} (C_L^{st} - C_{L,targ}^{st}) \quad (18)$$

$$S_{mucil,st} = q_{mucil} \frac{2\pi r_{in} L_i}{V_{st}} \quad (19)$$

$$280 \quad q_{mucil} = k_{mucil} starch \quad (20)$$

With $k_{targ, starch}$ (1/d) and k_{mucil} (cm/d) parameters controlling respectively the synthesis and decay rate of starch and the release of mucilage. $C_{L,targ,st}^l$ defines the target C_L^{st} and thus the threshold between starch synthesis and desynthesis. As in Hirschberg et al. (1998), the release of mucilage is represented via a simple first-order kinetic. We then obtain S_{mucil} ($mol\ cm^{-3}\ st\ d^{-1}$), the release rate of mucilage. $k_{mucil} > 0$ only at the tip (in our model: the last segment) of growing roots.

285 Both q_{exud} and q_{mucil} ($mol\ cm^{-2}\ d^{-1}$) can then be used by the [microscale-1D](#) soil domains (see Eq. (C13), Eq. (C17)).

2.2.4 Soil carbon transport

The changes in solute concentration are computed using the advection-diffusion-reaction equation—see Ahusborde et al. (2015) and Helmig (1997, Eq 3.100):

$$\frac{\partial \theta C_X^l}{\partial t} - \nabla \cdot (C_X^l \mathbf{u}_X) - \nabla \cdot (D_X(\theta, \phi) \nabla C_X^l) - S_X (-S_{1DS-3DS,X}) = 0 \quad (21)$$

$$290 \quad S_{ads,X} + S_{microbe,X} = S_X \quad (22)$$

C_X^l (mol/cm^3 water) is either the concentration of carbon from dissolved low molecular weight organic compounds (C_L^l) or high molecular weight organic compounds (C_H^l) in the soil liquid solution. S_X ($mol\ C\ cm^{-3}\ scv\ d^{-1}$) is the net source of C_X^l , and it encompasses the adsorption of C_X^l ($S_{ads,X}$, $mol\ C\ cm^{-3}\ scv\ d^{-1}$, see section 2.2.5) and the changes caused by microbial reactions ($S_{microbe,X}$, $mol\ C\ cm^{-3}\ scv\ d^{-1}$, see appendix D and Eq. (C1)). $S_{1DS-3DS,X}$ is an additional source term only used [at the microscale by the 1D soil models](#) and presented in Eq. (C7). \mathbf{u}_X is the advective flux.

295

$D_X(\theta, \phi)$, the effective dispersion in the porous medium (caused by molecular diffusion and hydrodynamic dispersion), is obtained from Millington and Quirk (1961), as implemented by Koch et al. (2021):

$$D_X(\theta, \phi) = D_{Xw} \frac{\theta^{\frac{10}{3}}}{\phi^2} \quad (23)$$

With D_{Xw} (cm^2/d) the diffusion coefficient of the carbon compound in water. Low molecular weight organic C compounds can be modelled as true solutes and are thus subject to convection. High molecular weight carbon compounds correspond to the plant mucilage releases, a gel-like substance when hydrated (Ma et al., 2022; Kutschera-Mitter et al., 1998). To simulate the rheology of mucilage slime, we adopt a simplified approach with a null advective flux (\mathbf{u}_H) and a lower diffusion coefficient ($D_{H,w} < D_{L,w}$). [The For simplicity, the](#) implementation of Eq. (21) for the [macroscale-3D](#) soil and [microscale-1D](#) soil are given respectively in appendices C1 and C2.

300

305 2.2.5 Adsorption

We describe in this section the adsorption of low molecular weight organic C compounds (C_L) from the liquid phase (C_L^l) onto the solid phase (C_L^s). As Sircan et al. (2025), we describe the adsorption as linear. However, instead of equilibrium sorption, we used linear first-order kinetics. ~~This was necessary in order to obtain the same result at the micro- and the macroscale. Moreover, not~~ [Not](#) assuming instantaneous equilibrium [allows-allowed](#) us to set lower time steps during the simulations. We further assume that only the low molecular weight organic substances can be sorbed. Adsorption was implemented according

310

to Ahrens et al. (2020):

$$\frac{\partial C_L^s}{\partial t} = k_{ads} C_L^l (C_{L,max}^s - C_L^s) - k_{des} C_L^s \quad (24)$$

$$C_{L,max}^s = C_{L,max}^l = \rho_b \frac{k_{clsmax} k_{clay+<20\mu msilt}}{M_{mass,C}} \quad (25)$$

with k_{ads} ($23265.14 \text{ cm}^3 \text{ scv mol}^{-1} \text{ C d}^{-1}$) and k_{des} (4.47 d^{-1}) the adsorption and desorption rates, defined according to the values given by Ahrens et al. (2020). $C_{L,max}^s$ ($\text{mol C cm}^3 \text{ scv}$) is the maximal concentration of adsorbed C_L^s , ρ_b ($1.51 \text{ g mineral soil/cm}^3 \text{ scv}$) is the soil bulk density, $M_{mass,C}$ ($12.011 \text{ g C mol}^{-1} \text{ C}$) is the molar mass of carbon, $k_{clay+<20\mu msilt}$ (0.45–, Wang et al. (2018)) is the fraction of clay and silt smaller than $20 \mu\text{m}$. k_{cssmax} (–) is an empirical factor defined according to the ~~make up~~ make-up of the clay minerals in the soil. According to Jiang et al. (2014), the Selhausen soil (used to calibrate the model) is dominated by illites, a 2 : 1 mineral. Consequently, $k_{cssmax} = 0.079$ (Ahrens et al., 2020). This yields $C_{L,max}^s = 4.5e^{-3} \text{ mol C cm}^{-3} \text{ scv}$). From the adsorption dynamic, we get the adsorption net sink for dissolved low molecular weight organic carbon: $S_{ads,L} = -\frac{\partial C_L^s}{\partial t}$, used in Eq. (22). As high molecular weight organic carbon compounds represent the mucilage gel, no adsorption is represented: $S_{ads,H} = 0$.

2.2.6 Plant growth and perirhizal zone

We simulate dynamic plants that grow according to input parameters (representing plants' genetically determined characteristics) and to their water and sucrose status (see Giraud et al. (2023)). This leads to changes in root segment lengths and to the creation of new root segments. The method used to define the shape, water and solute content of new or growing microscale-1D soil domains is presented below.

Currently, each root segment (and associated microscale-1D soil domain) is allocated to the soil voxel containing its centre when the segment is first created, even if one or both segment extremities are out of the voxel. In order for the output to be realistic, it is therefore important for the maximum root segment size to be lower or near the length of a soil voxel. Moreover, for the current setup, root segments will remain in the same soil voxel even if the centre of an existing segment (because of segment elongation) moves to another voxel. We assume that these simplifications will have a limited influence on the results if the resolution is high enough. After simulating plant growth for the time step N , we obtain the new lengths (L , cm) and radius (r_{in} , cm) of the roots, which are, respectively, the lengths and inner boundary of the microscale-1D soil domains. From these shape data, we recompute the distance to the outer radial boundary of the each 1D domains domain (r_{out} , cm) according to the volume of the soil voxels ($V_{3DS,scv}$) and the number of microscale-1D soil domains contained within (nr). For the microscale 1D soil domain k in the voxel i :

$$r_{out,k}^N = \sqrt{\frac{r_{in,k}^2 L_k^N}{\sum_j^{nr_{i,N}} r_{in,j}^2 L_j^N} \frac{V_{3DS}}{\pi L_k^N} + r_{in,k}^2} \quad (26)$$

Consequently, creation of new roots or changes in L and r_{in} of existing roots in a voxel can lead to changes in the perirhizal zone volume allocated to an individual microscale-1D soil domain. From the soil point of view, a higher density of roots in the voxel leads to smaller perirhizal zones for each of the root segments contained in it. The description of the method used to

update the amounts of water and carbon in the [microscale-1D soil](#) models as they are created and change shape is presented in the appendix E. In brief, the gradient of existing 1D domains with decreasing volume is preserved. We then compute the volume of newly freed soil for each voxel. From the amount of water and carbon lost by the 1D models, we obtain the mean water content and carbon concentration in the freed space. This volume (and corresponding water and solute contents) is distributed
 345 between the new and expanding [microscale-1D soil](#) domains to respect the results of Eq. (26).

2.3 Model application: The effect of dry spells on soil carbon and microbial dynamics near a growing plant

The implementation setup for the coupled model follows [the one that](#) used in Giraud et al. (2023), with Van Genuchten parameters measured from one testing site of the Research Center Jülich, located in Selhausen (50.8659 N, 6.4471 E). We
 350 simulated the growth of a C3 monocot under a control scenario (hereafter called *baseline*) and compared it with dry spells at two plant development stages. Both dry spells were represented by a period of one week without rainfall: 11 to 18 days and 18 to 25 days after sowing, hereafter called *earlyDry* and *lateDry*. The [periods of the dry spells were taken from \(Giraud et al., 2023\)](#). [The start of the earlier spell was set late enough to assume that the seed no longer affected the plant's carbon balance. The end of the late spell was set early enough to finish before the end of the plant's vegetative growth, as the development of flowers, seeds and fruits is not yet implemented within CPlantBox.](#) The virtual plants were obtained by running the plant model up
 355 to day 10 (after sowing) using a mean empirical growth rate (neither dependent on the carbon- nor water-flow) and without simulation of the water and carbon fluxes. From the 10th day onward, we simulated the carbon and water fluxes and ~~of~~ the carbon- and water-limited growth, as well as the soil carbon dynamic. The plants all grew under the *baseline* conditions with a static soil (constant mean $\psi_{m,soil} = -100$ cm at hydraulic equilibrium), except during the dry spells: day of growth 11 to 18
 360 for *earlyDry*, and 18 to 25 for *lateDry*). At the start of the dry spells, the mean soil water potential was lowered to -450 cm and warmer and drier atmospheric conditions were simulated ([see Table 1](#)). After the dry spells, the *baseline* environmental and static soil water conditions were used. We used a no-flux boundary condition for the [macroscale-3D](#) soil. Fig. 5 summarises the timeline of the three scenarios.

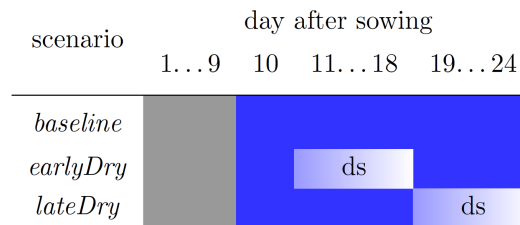


Figure 5. Timeline of the simulations. The grey (resp. blue) cells indicate the empirical (resp. semi-mechanistic) growth period and "ds" the implementation of the dynamic soil. The blue gradient during the dynamic soil simulation represents the decreasing soil water content. The sharp shift from dark to light (resp. light to dark) blue at the beginning (resp. end) of the dry spells represents the implementation of the *dry spell* (resp. *baseline*) atmospheric and initial (resp. constant) soil variables. Figure taken from Giraud et al. (2023)

Table 1. Environmental variables. Taken from Giraud et al. (2023).

	min-max of sinusoidal function ^a			$\theta_{init,soil}^b$
	PAR	$T_{atm}^c (K)$	$h_{atm}^d (\%)$	$(cm^3 cm^{-3})$
<i>baseline</i>	0-940	288.95-295.15	0.6-0.88	0.4
<i>dry spell</i>		293.85-303.42	0.44-0.74	0.28

^a T_{atm} , PAR and h_{atm} go from their minimum to their maximum value via a sinusoidal function with a period of 1 day to represent the day-night cycle. All the atmospheric variables correspond to values 2 m aboveground.

^b $\theta_{init,soil}$ is the value of the constant soil water content of *baseline*; and it corresponds to the initial value of the dynamic soil water content for the dry spell periods.

^c T_{atm} is the air temperature. It affects only the plant processes and not the microbial activity.

^drelative air humidity.

the *baseline* variables were used for all simulations before and after the dry spells

For the plant model, the calibration of Giraud et al. (2023) was adapted to better follow our assumptions regarding the plant mucilage and carbon dynamic (e.g., preferential carbon usage for growth over exudation), and to address the divergences between the outputs of Giraud et al. (2023) and experimental ~~observation~~ observations (light dry spells caused carbon rather than water scarcity). Changes were also made to the lateral conductivity ($k_{lat,x}$, $cm hPa^{-1} d^{-1}$) in the mature root zone: the constant value was computed according to Bramley et al. (2007, Fig. 4.15) and set to 1.5% of $k_{lat,x}$ within the immature root zone, see appendix H for a more detailed description.

370

The selection of the ~~biokinetic~~ kinetic parameter sets is given in appendix I. Briefly, Sircan et al. (2025) computed 1650 parameter sets respecting parameter and process constraints defined according to a literature review. Those sets were divided into three groups according to the resulting ~~C_L^i radial profile for the microscale~~ radial concentration profile of the low-weight dissolved organic molecules-C for the 1D soil domains. 33 sets were selected randomly from each group and a simulation was run for four days. We then selected three parameter sets giving diverging results. Table 2 presents the characteristics of the selected sets. The first set, thereafter called *highCO2*, led to a high CO₂ production in spite of a low microbial development. The second set, thereafter called *highMB*, led to the highest microbial development. The third set, thereafter called *lowMUuptake*, led to the highest soil solute concentration (low microbial solute uptake).

375

Table 2. Qualitative summary of the characteristics of the three selected ~~biokinetic~~ kinetic parameter sets (*highCO2*, *highMB*, *lowMUptake*). O: oligotrophs, C: copiotrophs, (de)act.: activation and deactivation, VS: compared against. The evaluation has the following order (representing the low-to-high gradient): $- < . < + < ++$

denomination	fitness		act. VS deact.		(de)act. VS growth + decay		O VS C		CO ₂ losses
	O	C	O	C	O	C	(de)act.	fitness	
<i>highCO2</i>	+	-	.	.	+	+	.	+	++
<i>highMB</i>	+	.	+	+	++	.	-	+	+
<i>lowMUptake</i>	-	.	-	-	.	.	.	-	-

2.4 Analysis of the outputs

380 The data were treated and visualised with Python3.10 (Van Rossum and Drake Jr, 1995).

2.4.1 Scale definition for model evaluation

To take advantage of the multiscale setup, we evaluated the model outputs at four different scales: ~~macro~~scale, ~~bulk soil~~, ~~perirhizal_{trunc}~~, ~~micro~~scale, ~~3D scale~~, ~~aggregated perirhizal scale~~, ~~1D scale~~. We define the ~~macro~~scale ~~3D scale~~ as the soil voxels (Fig. 1(b)) that contain at least one root at the end of the simulations in any of the scenarios. This soil space is therefore constant and equal for all ~~the~~ simulations. We define the ~~bulk soil as the macro~~scale ~~aggregated perirhizal volume as the 3D~~ data of voxels containing at least one root at each time step, potentially different for each simulation. ~~For the perirhizal_{trunc} analysis, we aggregated the mean concentrations for each microscale domain truncated at a specified distance from the root surface, denoted as $\Delta r_{analysis}$. The value of $\Delta r_{analysis}$ was pragmatically set to 0.06 cm. It corresponds to the minimum distance between the inner and outer boundaries of the microscale domains obtained during the simulation. By setting a~~ consistent distance between the inner and outer boundaries of the truncated domains, we obtained a partial uniformity between the domains that reduced variations that could arise from differences in soil allocation to each domain (see Eq. (26)). Moreover, using these truncated ~~At the~~ 1D models allowed us to evaluate a soil area nearer to the root surface, for comparison with the bulk soil scale. ~~At the microscale, we evaluated the data for each~~ ~~scale, we evaluated the data for each~~ segment of the 1D models (Fig. 1(e)) ~~individually~~ models (Fig. 1(c)) individually. Therefore, the 1D scale covers the same area as the aggregated ~~perirhizal volume but represents a higher resolution: one data point for each sub-control volume of the 1D-models, against one data point for each sub-control volume of the selected 3D soil voxel. A graphical representation of the different scales is given in Fig. J1.~~

385

390

395

2.4.2 Vertical concentration profiles

To evaluate the vertical concentration profiles, we computed the total soil volume, water volume, and C content (for each C pool) for each depth at the bulk-soil-aggregated perirhizal scale. This allowed us to determine the corresponding mean C concentration and water content around the root systems. Additionally, we present the minimum and maximum C concentrations (for each C pool) and water content values at each soil depth to capture the lateral variability.

2.4.3 ~~Complementary cumulative volume distributions~~

~~We created complementary cumulative volume distribution plots of the concentration at the perirhizal_{trunc} scale. In these plots, the x-axis represents different concentration values. The y-axis shows the absolute or relative volume of the selected perirhizal zone (up to 0.06 cm from the root surface) with concentrations equal to or greater than the corresponding x-axis value. This representation provides insight into the spatial variability and distribution of concentrations within the specified zone. Additionally, these graphs allow us to evaluate how much of the perirhizal zone volume exceeds certain concentration levels, such as hotspot thresholds.~~

2.4.3 Relative change of soil carbon distribution

To evaluate the change in plant-soil C turnover for the different weather scenarios and biokinetic-kinetic parameter sets, we categorised C into four pools: solutes (dissolved and sorbed low molecular weight organic C and high molecular weight organic C), microbes (including total oligotrophic and copiotrophic C), active microbes (including active oligotrophic and copiotrophic C), and emitted CO₂ (microbial respiration). We evaluated the C content at the macroseale-3D scale for each of these pools at each time step and normalised the values for *earlyDry* and *lateDry* under each biokinetic-kinetic parameter set according to the corresponding baseline scenario: $(C_{X,spell} - C_{X,baseline})/C_{X,baseline}$, with *X* the name of the C pool.

2.4.4 Soil organic carbon hotspots

One important effect of plant C releases is the resulting changes in soil organic C (SOC). Using the study of Poeplau and Don (2023), we set three SOC thresholds defined according to the $SOC/SOC_{expected}$ ratio. These ratios correspond to the thresholds between soil quality classes: 0.65 (degraded to moderate soil quality), 0.83 (moderate to good soil quality), 1.16 (good to very good soil quality). These thresholds are used to distinguish hot spots. Since the studied scale in our study is much smaller than the one in the study of Poeplau and Don (2023), we only use their threshold value, but refrain from the qualitative interpretation. $SOC_{expected}$ ($mol\ C\ cm^3\ scv$) is the average expected SOC according to the soil clay content. Following the method of Poeplau and Don (2023) and the Selhausen clay content data of Wang et al. (2018), we set $SOC_{expected} = \frac{\rho_b}{1000M_C}(0.0288k_{clay} + 13.674) = 2.36e^{-3}\ mol\ C\ cm^3\ scv$, with k_{clay} the soil clay content in $g\ kg^{-1}$, ρ_b ($g/cm^3\ scv$) the soil bulk density, M_C ($g\ C/mol\ C$) the C molar mass. The ~~1/1000~~ $1e^{-3}$ factor is used to go from $g\ kg^{-1}$ to $g\ g^{-1}$.

A first evaluation of the outputs showed that our simulated values were almost always in the lowest SOC class and significantly below the mean Selhausen SOC reported by Wang et al. (2018). For the hotspot analysis, we therefore added an SOC_{slow} class representing the very slow decomposing SOC, assumed to be constant over the study period. We set $SOC_{slow} = SOC_{theoric} - SOC_{simulated,init}$, with $SOC_{simulated,init}$ the total SOC explicitly simulated in our model (solute and microbial pools) at the beginning of the simulation, and $SOC_{theoric}$ the value from Wang et al. (2018). We then set the variable $SOC = SOC_{simulated} + SOC_{slow}$, computed ~~at the microseale~~by the 1D soils. This is the value used to compare with the SOC thresholds given by Poeplau and Don (2023). We use a stacked area graph to represent the SOC hotspots, as soil above the higher thresholds is also above the lower thresholds. As presented by Landl et al. (2021a), we computed the volume-normalised hotspot volume ~~to obtain~~. The relative hotspot volume was obtained by dividing the volume of each hotspot by the volume of all the 3D voxels that contained at least one root. Using a relative value allowed us to compare hotspot volume across root system architecture and age. The volume-normalised hotspot can be used as a metric for the efficiency of the rhizodeposition (Landl et al., 2021a).

440 2.4.5 Radial concentration profiles

Finally, for the ~~microseale~~1D scale, we presented C concentration data for all segments of the 1D domains on one scatter plot per weather scenario and parameter set, giving, on the x-axis, the radial coordinate of the segment centers according to the root surface.

2.4.6 Complementary results

445 Some additional results, namely the complementary cumulative volume distributions, can give an overview of the status of the perirhizal zones of each simulation. However, as they do not affect the main conclusion of this work, these results and related analysis were moved to the appendices K3, L and M.

3 Results

3.1 Three-dimensional simulation of microbial carbon dynamics in the soil-plant system

450 ~~The integrated plant-soil-microbiome model extends the FSPM model of Giraud et al. (2023)) by implementing microbial carbon (C) dynamics in soil. Consequently, the extended FSPM model now provides 3D simulations of plant architecture, water and C dynamics in the plant and the soil, and microbial dynamics in the soil, including the rhizosphere. While this study focuses on C cycling and microbial dynamics at the root-soil interface, more detailed simulation results regarding water and C flow in a plant for the three considered weather scenarios are given in Giraud et al. (2023).~~ Fig. 6 shows a typical simulation after 24.5d of plant growth (midday) for a biokinetic-kinetic parameterisation that results in high microbial CO_2 production (*highCO2*) under the *baseline* scenario. The simulation highlights ~~an increasing a~~a concentration of dissolved low molecular

weight organic C (C_L^l) and active copiotrophs (C_C^a) that increase with time in response to root exudation in the upper part of the root system, near its centre.

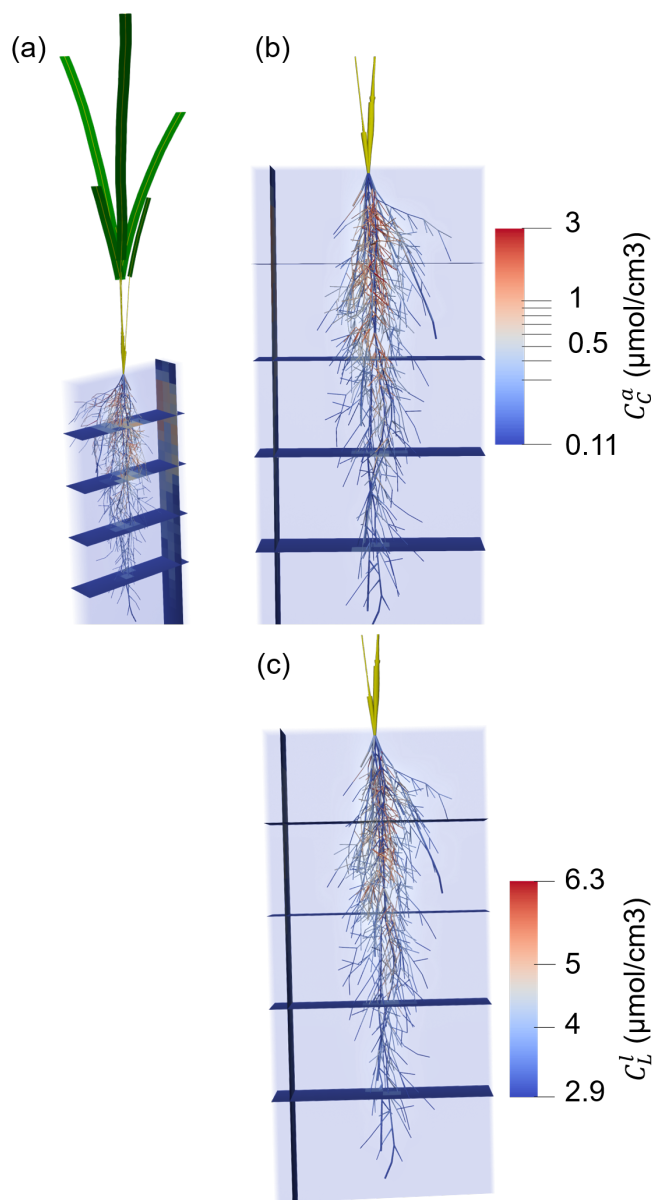


Figure 6. 3D representation of the plant and soil domains for microbial biokinetic-kinetic parameterisation *highCO2* under the *baseline* scenario at 24.5d of growth, with (A) the whole plant, (B, C) zoom on the root system. (A, B) present the concentration of active copiotrophic biomass C (C_C^a) and (C) that of dissolved low molecular weight organic C (C_L^l). The soil colours give the concentration in the bulk soil-aggregated perirhizal volume. The root colours give the concentration at the soil-root interface. Unified colours were given to the aboveground roots, shoot segments and leaf blades.

3.2 Plant processes

460 Fig. 7.A presents the plant xylem mean water potential (ψ_x , cm) for each ~~biokinetic~~kinetic parameterisation and weather scenario. ~~The simulation results reflect the diurnal dynamics of ψ_x in response to the truncated sinusoidal curve used to describe the atmospheric conditions (light input, air relative humidity and temperature). These boundary conditions affected the plant stomatal regulation and, thus, the C assimilation and transpiration rate. We observed an increase of ψ_x over time for *baseline* as the leaf:root ratio decreased. With drought, ψ_x rapidly decreased at the beginning of the dry spells and recovered after its end in~~
465 ~~response to changes in soil water content and mean air relative humidity during the dry spells (see (Giraud et al., 2023, Table 1)). Towards the end of the simulation with *earlyDry*, ψ_x converged to the values of *baseline*, indicating a full recovery of the plant water uptake capacity. As a consequence of the lower leaf:root ratio, ψ_x was lower at the beginning of the dry spells for *earlyDry* (11th day) than for *lateDry* (18th day). However, in comparison to *earlyDry*, the higher plant transpiration under *lateDry* with higher total leaf and root surface area led to a quicker decrease in soil water content. This mechanism induced lower ψ_x in the~~
470 ~~*lateDry* compared to the *earlyDry* scenario from the third day after the start of the dry spells onward (13th and 20th day). while~~
Fig. 7.B shows the the cumulative amount of C used for growth (and growth respiration) ~~during the simulation. The diurnal cycle of ψ_x affected the plant growth. During the day, the transpiration-induced lower ψ_x limited plant growth compared to the night, where increased ψ_x permitted faster, albeit still water-limited, C assimilation and biomass production. The required C was taken from the starch reserves built up during the day. We can also observe trends over the simulation. Unlike younger~~
475 ~~plants, older plants could maintain growth during the diurnal period outside of the dry spells. During the dry spells, growth occurred only at night and at a severely reduced rate. On the last three nights of the late dry spell, we observed no night-growth due to lower plant night ψ_x . Plant maintenance respiration increased during the spells due to the higher air temperatures. In the *earlyDry* scenario, plant growth was severely limited compared to *baseline*, resulting in a lower plant maintenance respiration rate after the dry spell ended (18th day). For *lateDry*, we saw a lower maintenance respiration rate from the fourth day of~~
480 ~~the spell. The total cumulative respiration (growth and maintenance related) at the end of the simulation was highest for the *baseline* scenario. It was about 29% lower for *earlyDry* and 18% lower for *lateDry*. The biokinetic parameterisation had no significant effect on the plant water processes or growth and maintenance. As the focus on this study is the water and carbon dynamic within the perirhizal zone, the plant specific results are not included in the main body of the text. However, the outputs~~
plant processes differ from those of Giraud et al. (2023). The plant results are therefore presented and discussed in section K3.
485 Figs 7.C and D present the cumulative amount of C used for (respectivelyrespectively) passive exudation and active mucilage release during the simulation. The plant water balance affected the exudation rate and mucilage release rate much more strongly
than the kinetic parameterisation of the soil microbes. During the simulation, exudation occurred only under limited C use for growth in response to low ψ_x plant water potential. Although the mucilage release was not always maintained at night (e.g., at the end of the *earlyDry* simulation), this process was less affected by the diurnal cycle than the exudation. This led to qualitative
490 changes in the type of C released by the plant throughout the day. In comparison to *baseline* the early dry spell led to a lower total exudation at the end of the simulation: -32%, -35%, and -37% for *highCO2*, *highMB*, and *lowMUuptake*, respectively. Conversely, the late dry spell resulted in higher exudation compared to *baseline*: +29% for *highCO2* and *highMB*, +26% for

lowMUptake. Thus, the dry spells triggered both positive and negative effects on exudation. On the one hand, more C became available for exudation with decreasing plant growth. On the other hand, the lower ψ_x xylem water potential led to a closing
495 of the stomata, causing a lower assimilation rate per unit of leaf area.

Lower plant growth also led to lower leaf and total root surface area compared to *baseline*, which limited exudation. The effect of the simulated spells on total exudation was, therefore, dependent on the stage of plant growth at the start of the spell. After the end of the early dry spell (days 18-23), the daily plant exudation for *earlyDry* was higher than for *baseline*, partially reducing the exudation gap between the two. The total mucilage release under *earlyDry* at the end of the simulation
500 was close to that under *baseline*: -11%, -2%, and +6% for parameter sets *highCO2*, *highMB*, and *lowMUptake*, respectively. In contrast, mucilage release reached +139% for *highCO2* and *highMB* and +170% for *lowMUptake* under *lateDry*. ~~We found an effect of the biokinetic parameterisation on exudation and mucilage release.~~ The *highCO2* and *highMB* parameterisation, associated with high microbial C usage rates resulted in lower concentrations of dissolved low molecular weight organic C (C_L^1 $mol\ C/cm^{-3}$). This lower C_L^1 concentration of low-weight dissolved organic molecules-C increased passive exudation.
505 The higher exudation was then compensated by the plant starch reserves (in the mesophyll and around the plant transport tissues) and did not affect the plant growth or maintenance. As the mucilage release rate is a function of the plant starch content, higher exudation induced lower mucilage release.

These differences in exudation and mucilage release appeared only after the onset of the dry spells. The release rates between the biokinetic kinetic parameterisations for *earlyDry* became similar again from the third day after the end of the dry spell
510 onward. The highest exudation rate per unit of root surface for a single segment across all simulations was obtained at 11 day 5hrs with *baseline* and *lateDry* ($0.153\ mmol\ C\ cm^{-2}\ d$), ~~at 10 day 21hrs for *highCO2* and *lowMUptake* with *earlyDry* ($0.145\ mmol\ C\ cm^{-2}\ d$), and at 12 day 8hrs for *highMB* with *earlyDry* ($0.148\ mmol\ C\ cm^{-2}\ d$).~~

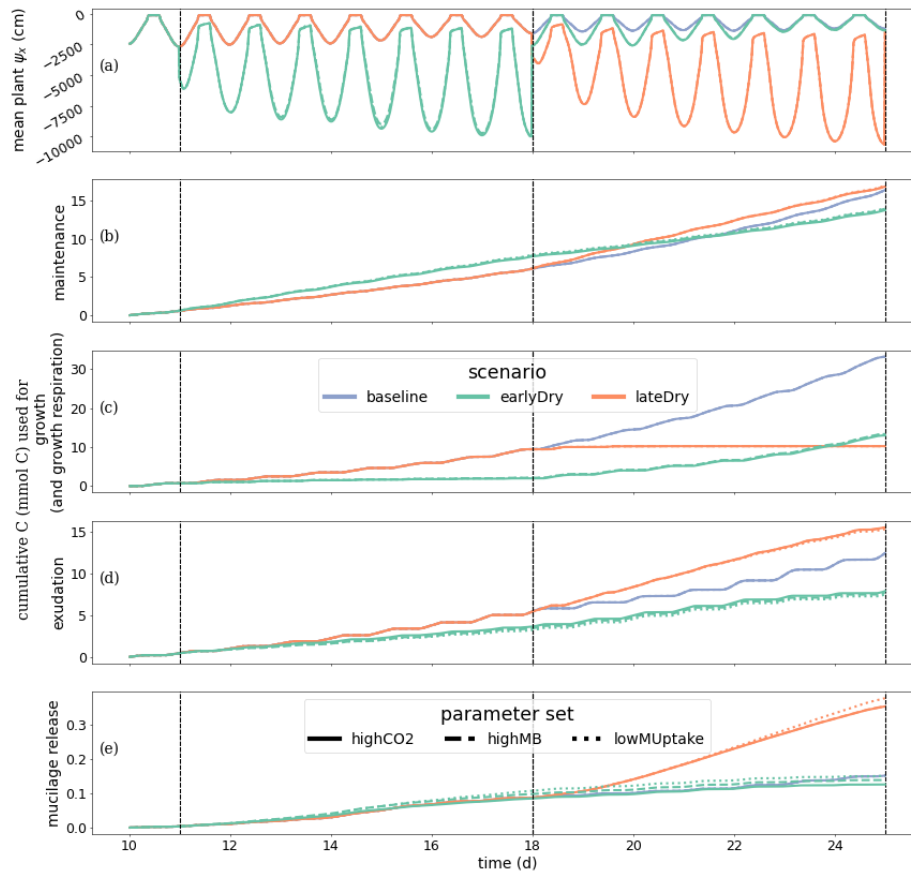


Figure 7. Dynamics of Plant water- and carbon-related metrics: (A) the mean plant xylem water potential (ψ), (B) cumulative carbon growth and growth respiration, (C) exudation, (D) mucilage release. Line colours indicate the weather scenarios: baseline (blue) against an early (green) and late (red) dry spell. Line types represent different **biokinetic-kinetic** parameter sets. The vertical dotted lines show the beginning and end of the early (day 11 to 18) and late (day 18 to 25) dry spells. Please note that the scales of the y-axis differ in each subplot.

3.3 Vertical concentration profiles

515 **Fig. ?? presents Figs. 8 and 9 present** the vertical concentration profiles of organic C, microbial pools and produced CO_2 from microbial respiration at the end of the simulation for the **bulk-soil-aggregated perirhizal** scale. The root system structure affected the concentration profiles. The peaks of organic C between 0 cm and 13 cm of depth corresponded with the location of the highest root **length-tip** density (Fig. ??(a), (b), (c) 8 and Fig. K1). A second peak of maximum C concentration was

observed at the bottom of the profile close to the tips of primary roots. ~~That second peak is especially strong for lateDry due to lower root growth, i.e., very few root segments reached 20 cm, such that the mean C concentration equals the maximum concentration. Conversely~~ Moreover, the higher lateral dispersion of the roots in the upper soil layers led to a stronger lateral variability in the concentration profile (larger min-max ranges). ~~Stronger plant growth under baseline resulted in perirhizal zones and rhizodeposition at lower depths such that the C accumulation at the root tips was much less pronounced compared to the dry spell scenarios.~~

525 The ~~biokinetic-kinetic~~ parameterisation strongly influenced the C concentration profiles (Fig. ??(e)-(j)9). With *highCO2*, solute and microbial C concentrations ($C_L^l, C_L^s, C_H^l, C_C, C_O$) ~~were lowest~~ were lowest: $3.5 \pm 0.49 C_L^l, 77.96 \pm 9.25 C_L^s, 595.72 \pm 6.99 C_H, 10.17 \pm 0.79 C_C, 9.31 \pm 0.02 C_O$, all in $\mu\text{mol cm}^{-3}$. The *highMB* parameterisation also led to low solute concentrations but triggered increased microbial C due to higher microbial C consumption: $2.48 \pm 0.20 C_L^l, 55.96 \pm 2.98 C_L^s, 698.46 \pm 7.76 C_H, 22.50 \pm 8.44 C_C, 9.49 \pm 0.04 C_O$, all in $\mu\text{mol cm}^{-3}$. With *lowMUptake*, solute concentrations were highest because of low microbial C uptake: $8.18 \pm 3.67 C_L^l, 174.06 \pm 67.33 C_L^s, 651.78 \pm 8.24 C_H, 12.19 \pm 4.23 C_C, 9.39 \pm 0.08 C_O$, all in $\mu\text{mol cm}^{-3}$. These results are consistent with the observations obtained during the model conditioning (see appendix I).

~~Although the~~ The weather scenarios had a less pronounced effect on the concentration profiles than the ~~biokinetic parameterisation, the dry spells induced distinct effects. Under kinetic parameterisation. We obtained~~ $4.51 \pm 2.62 C_L^l, 98.87 \pm 54.34 C_L^s, 645.90 \pm 42.18 C_H, 14.26 \pm 6.77 C_C, 9.41 \pm 0.10 C_O$ under *baseline*, $4.37 \pm 2.56 C_L^l, 96.33 \pm 53.12 C_L^s, 647.53 \pm 42.37 C_H, 14.16 \pm 6.42 C_C, 9.38 \pm 0.07 C_O$ under *earlyDry*, ~~the concentration profiles were~~ and $5.47 \pm 4.68 C_L^l, 116.66 \pm 87.57 C_L^s, 655.15 \pm 43.45 C_H, 17.15 \pm 9.87 C_C, 9.40 \pm 0.08 C_O$ under *lateDry*, all in $\mu\text{mol cm}^{-3}$. For each kinetic parameter set, *earlyDry* led to concentration profiles close to that of *baseline* (e.g., C_C , Fig. ??(e)), or between *baseline* and *copiotrophs-C* (in $\mu\text{mol cm}^{-3}$): respectively 10.03 ± 0.69 and 9.98 ± 0.59 for *lateDry* (e.g., C_H^l , Fig. ??(b)). This shows the resilience soil

540 ~~C cycling to the early dry spell.~~ *highCO2*, 20.64 ± 6.86 and 20.88 ± 6.86 for *highMB*, 11.74 ± 3.32 and 11.91 ± 4.75 for *lowMUptake* (Fig. 9). In particular, despite the lower total exudation, the concentration of C_L^l low-weight dissolved organic molecules-C for *earlyDry* is close to that of the *baseline* due to the increased exudation rate during the second half of the simulation (Fig. ??(a)) Figs. 7, 8): respectively (in $\mu\text{mol cm}^{-3}$): 3.36 ± 0.32 and 3.62 ± 0.63 for *highCO2*, 2.50 ± 0.14 and 2.54 ± 0.25 for *highMB*, 7.24 ± 2.59 and 7.36 ± 2.72 for *lowMUptake*. Under *lateDry*, we observed

545 higher C_H^l concentrations of high molecular weight organic compounds-C: $+1 \pm 0\%$ compared with the *baseline*. This can be attributed to higher plant mucilage release for this scenario (Fig. ??(b)). The highest C_C^a 8). *lateDry* led to an increase in the mean concentration of activated oligotrophs-C compared with *baseline*: $+81 \pm 64\%$ with *highCO2*, $+82 \pm 74\%$ with *highMB*, $+251 \pm 276\%$ with *lowMUptake*. However, the overall highest concentration values of activated oligotrophs-C and C_O^a values ~~copiotrophs-C~~ at the last time step ~~are were~~ obtained with *baseline* (Fig. ??(g), (h)9). The differences in the

550 ~~microbes' activation~~ high amount of activated microbial biomass can partly be explained by the water content remaining at the high default value (-100 cm) during the whole simulation under *baseline*. C uptake by copiotrophs for *highCO2* and *highMB* was higher (high C_C copiotrophs-C, and microbially respired C as CO_2 but lower C_L^l low-weight dissolved

organic molecules-C concentration) under *lateDry* for *highCO2* and *highMB* (Fig. ??(d),(e)). This results can be linked to the higher total exudation and lower water content, making the Cressources more readily available to the microbes) – respectively for *highCO2* and *highMB*: $+6.4 \pm 2.1$ and $+30.0 \pm 12.0$ copiotrophs-C, $+85.3 \pm 34.3$ and $+58.1 \pm 24.6$ microbially respired C, -6.5 ± 6.2 and -6.3 ± 4.2 low-weight dissolved organic molecules-C compared with *baseline*. However, for *lowMUuptake*, the higher C uptake by copiotrophs occurred under *baseline*. Thus, water was relatively more limiting than C supply for copiotrophic growth with *lowMUuptake* (compared with *highCO2* and *highMB*) because of a lower maximum uptake rate. For that same reason, maximum C_O for each parameter set is very slightly higher ($< +0.7\%$) under *baseline* and under *lateDry* led to $+45.0 \pm 17.7$ low-weight dissolved organic molecules-C compared with *baseline*.

Due to the short simulation time, C_O oligotrophs-C (Fig. ??(f)) showed no significant growth at the bulk soil scale ($< 3\%$ of growth), but aggregated perirhizal scale ($< 3\%$ of growth and $< 0.5\%$ of difference between the weather scenarios). Nevertheless, the dry spells weakly affected the proportion of active and dormant oligotrophs (Fig. ??(h),(j)).

Mean bulk soil concentrations of the soil carbon with depth (a) dissolved and (e) sorbed (C_L^s) low-molecular-weight organic molecules, (b) high-molecular-weight organic molecules (C_H^l), (d) produced CO_2 , (e, g, i) total (C_C), active (C_C^a) and dormant (C_C^d) copiotrophs, (f, h, j) total (C_O), active (C_O^a) and dormant (C_O^d) oligotrophs. Line colours indicate the weather scenarios: *baseline* (blue) against an early (green) and late (red) dry spell. Line types represent different biokinetic parameter sets. The semi-transparent bands give the range between the minimum and maximum concentration in soil voxels with at least one root segment at each depth.

3.4 Complementary cumulative volume distributions

Fig. L1 shows the perirhizal_{trunc} volume with a concentration inferior or equal to the value on the x-axis. The description and motivation of the evaluation method for this figure are given in section 2.4.3. A video presenting this graphic for each time step is available in the supplementary documents ("Video S1"). Root exudation and mucilage release triggered concentration peaks of solutes and microbial C in the perirhizal_{trunc} volume (Fig. L1). Particularly copiotrophs were strongly stimulated reaching high abundances up to $580 \mu\text{mol}/\text{cm}^3$ soil for *lateDry* with *highMB*. Both dry spells led to strongly reduced total perirhizal_{trunc} volumes at the end of the simulation of 42% for *earlyDry* and 34% for *lateDry* in relation to *baseline*. As a consequence, maximum C_H^l and C_C^l concentrations were lowest with compared with *baseline* because the released plant C triggering copiotrophic growth was distributed in a larger volume.

While *lateDry*, we observed $+19.0 \pm 37.6$ active oligotrophs-C with *highCO2* and *highMB* caused the highest CO_2 , C_C and C_C^d peaks at the bulk soil scale (see respectively Fig. ??(d),(e),(i)), for the perirhizal_{trunc} zone the highest values for these parameter sets were obtained with *earlyDry* (Fig. L1(e),(d),(f)). The simulated copiotroph growth during the dry spell occurred earlier under *earlyDry*, creating a larger pool of C_C^d ready to be re-activated and grow during periods of high exudation. As the growth rate is a function of the current C_C^a , high C_C^d early in the simulation favoured the overall growth in those hotspots.

Parameterizing the model with *highMB* under *lateDry* caused an especially high C_C over most of the perirhizal_{trunc} volume with an increase of active and a strong build-up of dormant copiotrophs (Fig. L1(d),(e),(f)). Conversely, although -42.9 ± 15.4 with *highMB* under *lateDry* also led to a slight increase in the total and dormant oligotrophic concentrations

(Fig. L1(g), (i)), it resulted in a decrease in the amount of active oligotrophs (Fig. L1(h)). This simulated pattern indicates a pronounced competition between oligotrophs and copiotrophs for this scenario-parameter set combination. Indeed, the C_O^a values started decreasing strongly during the second half of the late dry spell, when the exudation rate became also lower, $+28.6 \pm 45.5$ with *lowMUptake*.



Figure 8. Perirhizal zone volume according to the minimal concentration of organic carbon in the truncated perirhizal zones, for Mean concentrations of the soil carbon with depth within the aggregated perirhizal volume

(a) carbon of low-weight dissolved organic molecules (C_L^l), and (b) carbon of high-weight organic molecules in the water phase (C_H^l), (c) emitted CO_2 , (d) total (C_C) (e) active (C_C^a) (f) dormant sorbed ($C_C^d C_L^s$) copiotrophs in the soil phase low-molecular-weight organic molecules-C, (g) total (C_O) (h) active (C_O^a) (i) high-molecular-weight organic molecules-C (j) dormant (C_O^d) oligotrophs in the soil phase produced CO_2 . The line colors give Line colours indicate the weather scenario scenarios: baseline (blue) against an early (green) and late (red) dry spell. The line types represent the different biokinetic parameterisation kinetic parameter sets. Subplots d, e, f and g, h, i have The semi-transparent bands give the same scale on range between the x-axis minimum and maximum concentration in soil voxels with at least one root segment at each depth.



Figure 9. Mean concentrations of the soil carbon with depth within the aggregated perirhizal volume

(a, c, e) total (C_C), active (C_C^a) and dormant (C_C^d) copiotrophs, (b, d, f) total (C_O), active (C_O^a) and dormant (C_O^d) oligotrophs. Line colours indicate the weather scenarios: baseline (blue) against an early (green) and late (red) dry spell. Line types represent different kinetic parameter sets. The semi-transparent bands give the range between the minimum and maximum concentration in soil voxels with at least one root segment at each depth.

3.4 Carbon pool distribution

We wished to evaluate the changes in C turnover within the plant-soil system, according to the weather scenarios and for the different biokinetic kinetic parameter sets. Consequently, Fig. 10 gives the relative changes of C content in different C pools, compared with the *baseline* scenarios. The relative increases of activated microbes-C and solutes-C (Figs. 10.B(a) and 10(ed)) correspond to the night periods, when less C was exudated-exuded with *baseline* compared with the other scenarios. These periods also corresponded to higher soil matric potentials, diminishing the negative effects of the low soil θ on microbial activation during the dry spells. This relatively higher activated microbes-C pool was observed for the *highMB* and *highCO2* and *highMB* parameter sets with *lateDry* (up to +50.0% for *highCO2* and +88.3% for *highMB*), although the values were lower than for *baseline* at the last time step. The *lateDry* scenario led consequently to a redistribution of C in the microbial and emitted CO_2 pools. Therefore, in spite of the higher total exudation, the solutes-C pool is lower compared with the corresponding *baseline* scenarios (-2.7% for *highCO2* and -0.64% for *highMB*). On the contrary, for *lowMUptake*, the *lateDry* scenario

led to a higher accumulation of C within the solutes ~~pools~~ ~~pools~~ ~~+4.43%~~ (Fig. 10(d)). We can also note that, while we have converging relative solutes-C values between *earlyDry* and *lateDry* for *highMB* and *highCO2* and *highMB* by the end of the simulations, the values for *lowMUptake* are diverging. For all *earlyDry* scenarios, ~~we observe that less C enters~~ ~~less C entered~~ the soil domain, leading to relatively lower values compared with *baseline* for all ~~pools~~ ~~C pools~~ (Figs. 10(a-d)) by the end of the simulations. We also found a small relative uptick of activated microbes-C for the *highMB* and *highCO2* and *highMB* parameter sets after the spell (up to ~~-2.3%~~ and ~~+21.6%~~, Fig. 10(ba)), linked with the increased soil water content and plant exudation (Fig. 7). ~~The same graphic is presented for the bulk soil in appendix M1. Fig. 10(e) shows the ratio of cumulative microbial-to-plant emitted CO₂ (see Figs. 7(b,c) and 10(c)). This ratio was strongly affected by the microbial parameterisation. At the end of the simulation, the ratio was 0.89 ± 0.2 for highCO2 and highMB, against 24.6 ± 1.8 for lowMUptake. Under lateDry, we observed an increase of the microbial-to-plant emitted CO₂ in spite of the higher plant maintenance respiration. This is higher maintenance respiration was, however, over-compensated by the lower growth respiration.~~

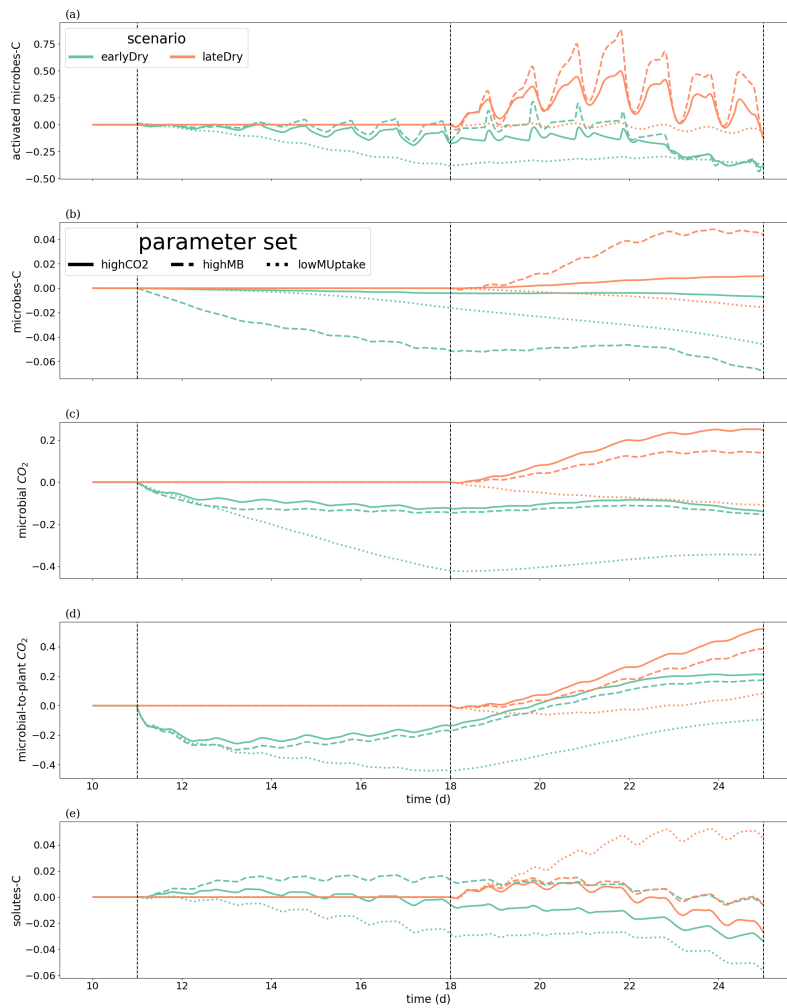


Figure 10. Relative change of root-released carbon use in response to drought scenarios according to the biokinetic parameters at the macroscale. Relative change of carbon allocation in response to drought scenarios according to the kinetic parameters at the 3D scale. The panels present the difference of C between each drought treatment and the baseline in relation to the baseline C usage for (a) microbes-C, (b) activated microbes-C, (c) CO_2 emitted by microbes, (d) soil solutes-C, as well as (e) the difference in microbial-to-plant emitted CO_2 . The line types represent the different biokinetic-kinetic parameterisation. The dotted lines show the beginning and end of the early (day 11 to 18) and late (day 18 to 25) dry spells. The line types represent the different kinetic parameterisation. Please note that the scales of the y-axis differ in each subplot.

3.5 SOC hotspots

615 To evaluate the root systems' exudation efficiency, Fig. 11 presents the relative volumes of the SOC hotspots (see definition in 2.4.4) in the bulk-soil aggregated perirhizal volume. We observed distinct characteristics over the whole simulation for each biokinetic-kinetic parameterisation. For *highCO2*, the absolute hotspot volume remained low and decreased with growth

(after an initial increase during the dry spells). For *highMB*, the total absolute hotspot volume ($SOC \geq 0.65SOC_{expected}$) peaked in the first half of the simulation before becoming stable under *baseline*. With this ~~biokinetic-parametrisation~~kinetic parameterisation, we observed the portion the relative hotspot volume within the highest class ($SOC \geq 1.16SOC_{expected}$): 8%, 32%, and 29% of the ~~hotspot-hotspot~~ volume belonged to the highest class under, respectively, *baseline*, *earlyDry*, and *lateDry*. Because of the low microbial activity, we observed the highest relative SOC hotspot volume for *lowMUptake* for all weather ~~scenario: $2.9e^{-2}$, $2.8e^{-2}$~~ scenarios: $2.9e^{-2}$, $2.8e^{-2}$, and 0.11 under *baseline*, *earlyDry*, and *lateDry*.

Outside of the dry spell, diurnal ~~eyeing-cycling~~ cycling in the hotspot volume followed the plant C releases during the day and soil C mineralisation without replenishment at night. This dynamic is especially strong with *highCO2* as the hotspot volume is (almost) back to 0 at the end of the day periods. The dry spells buffered the diurnal cycle for all scenarios as the exudation became more stable. Moreover, the biggest hotspot volume and intensity were obtained with the *lateDry* spell, following the higher exudation and mucilage release. *earlyDry* did not lead to a higher total hotspot volume (with $SOC \geq 0.65SOC_{expected}$) for the *highMB* and *lowMUptake* ~~parametrisation-parameterisation~~ at day 25 of growth. However, a bigger portion of the SOC ~~hotspot-hotspots~~ belonged to the higher hotspot classes ($\geq 0.83SOC_{expected}$ or $\geq 1.16SOC_{expected}$).

For *highCO2* and *highMB*, we observed an increase in hotspot volume during the first half of the dry spells, followed by a decrease during the second half. This can be partly explained by the decrease in daily plant C releases during the second half of the spell. The decreasing SOC hotspot volume in the second half of the spells was also due to the activation and growth of the microbial communities and C utilization after the first strong influx of plant C releases, followed by partial dormancy. Upon the next influx of C (the following day), the microbes reactivated and used the C more quickly. For *lowMUptake*, characterised by its lower microbial activity, we observed a continuous increase in hotspot volume throughout the spells.

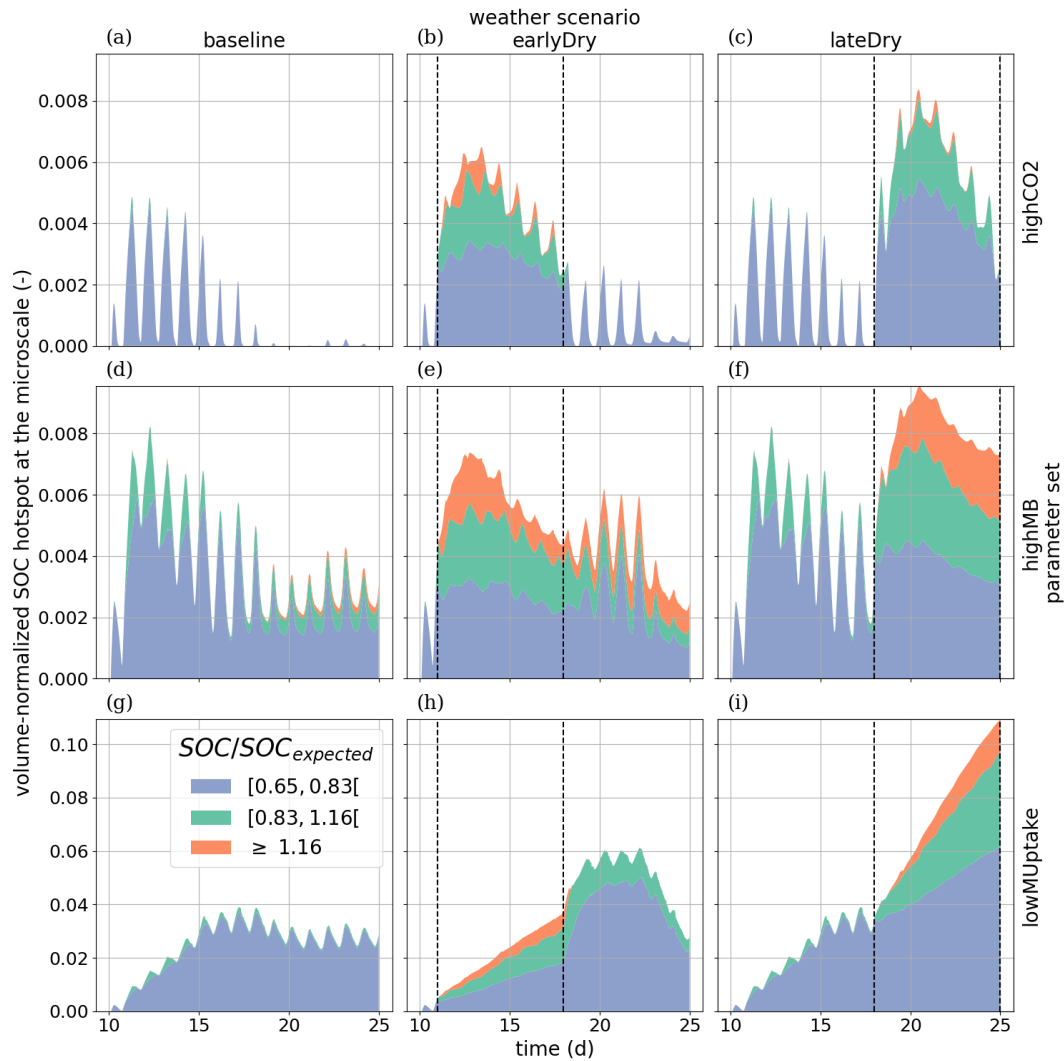


Figure 11. Relative SOC hotspot volume in the perirhizal zone according to time for the three $SOC/SOC_{expected}$ classes. The classes' ranges were defined according to Poeplau and Don (2023). The dotted lines show the beginning and end of the early (day 11 to 18) and late (day 18 to 25) dry spells. The third row is on a scale different from the first and second rows.

3.6 Radial carbon concentration profiles

To analyse how plant-soil system responses to drought affect the carbon stabilisation-fluxes in the perirhizal zone, Fig. 12 shows the connection between the extent of the microscale-1D soil domains and dissolved low-weight organic molecules concentrations C_L^l at the end of the simulation. Just as for the bulk-soil-and-perirhizal-trunc-scales-aggregated-perirhizal volume, the highest C_L^l -low-weight organic molecules-C concentrations were obtained under *lateDry* for *lowMUptake* ($144.0 \mu\text{mol cm}^{-3}$) and under *baseline* for *highCO2* ($33.0 \mu\text{mol cm}^{-3}$) and *highMB* ($24.0 \mu\text{mol cm}^{-3}$) (see Fig. M1). For *lowMUptake*, microscale

1D domains with shorter outer radii (segment index eight, which is nearer to the root surface) had higher radial concentrations. These shorter outer radii were caused by several microscale 1D domains sharing a single macroscale voxel. Therefore, the higher concentrations underline the additive effect on the C_L^l profile due to proximity to the perirhizal zone, which can indicate an overlap of the rhizospheres. This proximity effect was less noticeable for *highCO2* and *highMB*, where we observed concentration profiles less dependent on the extent of the microscale domain. Indeed, with those biokinetic parameter sets, the microbial C uptake was highest and reacted fastest to high influxes of dissolved low molecular weight organic carbon. Therefore, the C_L^l was closer to an asymptote at the outer boundaries of the 1D domain. These steeper gradients can cause a the rhizosphere extent to be smaller, limiting the rhizosphere overlaps. 3D scale voxel. The effect of rapid high microbial C uptake with *highMB* is also visible in Fig. 12(d) and E(e), where some concentration profiles show lower C_L^l low-weight organic molecules concentration values closer to the root surface, where the concentration of C_C copiotrophs-C is highest (see Fig. M). The C_L^l radial profiles radial concentration profiles of the low-weight organic molecules-C are strongly linked to the C_C copiotrophs-C profiles, as presented in appendix M.

The supplementary video "Video S2" shows C_L^l the low-weight dissolved organic molecules-C concentrations and the corresponding C_g^a/C_C active-to-total copiotrophs-C ratios at each time step. We found a high microbial growth during the first pulse of added C followed by a transition to dormancy due to reduced C availability. Microbes remained mostly dormant until reactivated by the next C input pulse while also slowly dying. Reactivation of the microbes during the periods of high C_L^l low-weight dissolved organic molecules-C concentrations led to a rapid consumption of the carbon in dissolved low-weight organic molecules near the root surface. As a result, C_L^l/C_L^l became lower than it initially was once the C input pulse diminished—at night (e.g., day 24.69 of growth) and during the second half of the dry spells. We can also observe how, during the dry spells, C_L^l low-weight dissolved organic molecules-C concentrations were higher than for the other two scenarios due to lower water content, favouring microbial C uptake.

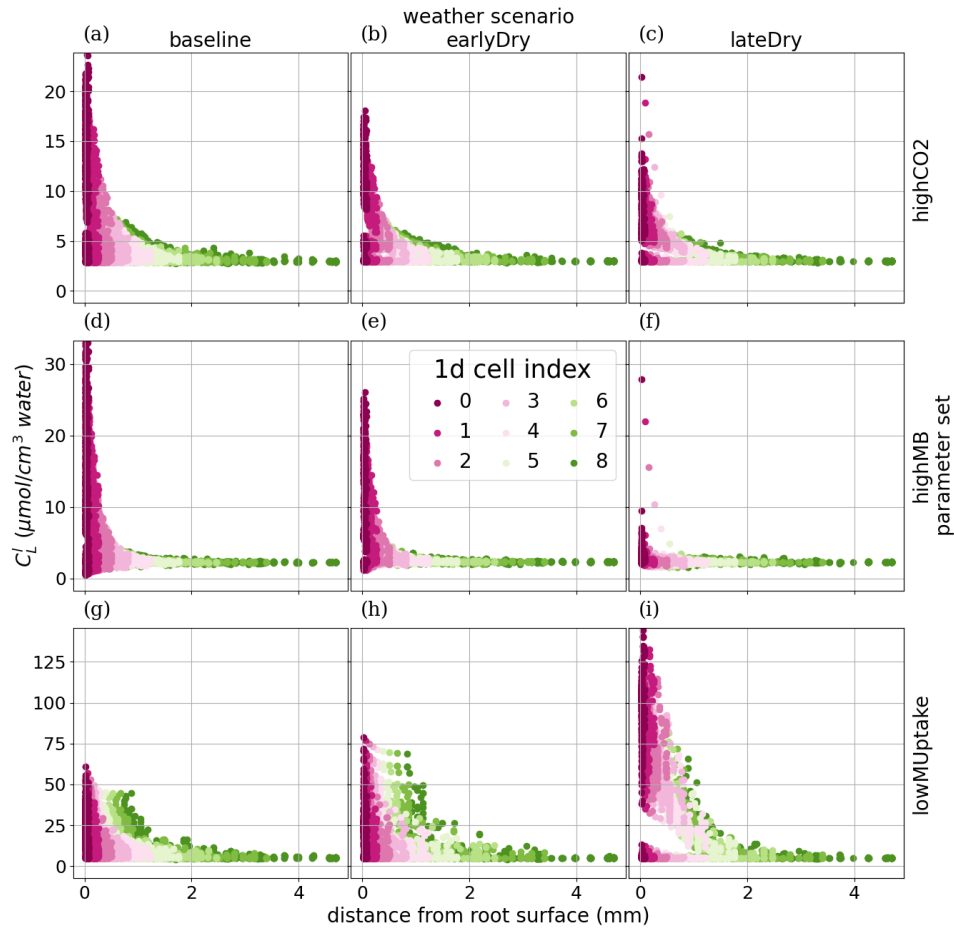


Figure 12. radial concentration profile of carbon from low molecular weight organic compounds at the end of the simulation for each plant perirhizal zone. r (mm) corresponds to the distance to the root surface. The colour gives the segment index of the 1D domain. Each column corresponds to a specific weather scenario and each row corresponds to a specific [biokinetic-kinetic](#) parameterisation.

665 4 Discussion

4.1 Model-setup

Several models have been developed to simulate the root-influenced spatiotemporal distribution of carbon (C) turnover and microbial activity (Pot et al., 2022; Srean et al., 2025), the exudation from a growing 3D root system (?), or the exudation and water uptake by a static plant, based on the C and water balance within the plant and soil (Lacointe and Minchin, 2019; Zhou et al., 2020)

670 ~~Rees et al. (2020) coupled *RhizoDep* with other models on the *OpenAlea* platform to represent the root-soil C and nitrogen exchanges for We developed a growing wheat plant, driven by its inner nitrogen and C balances. Regarding plant-soil coupling for solute flow using FSPMs, several modelling studies have represented the soil as a dynamic environment with water and solute uptake (generally phosphorus) or C releases from a static or growing root system (Ruiz et al., 2021; Schröder et al., 2014; Jorda-Guer~~
~~Braghiere et al. (2020) set up a 3D FSPM accounting for shoot-root-soil interactions for water and C using model coupling.~~
675 ~~However, their framework did not account for either C flow and microbial reactions in the soil or inner plant water and C balance. Some coupled models also represented inner plant water flow (Schröder et al., 2014; Mai et al., 2019; Schwartz et al., 2016)~~
~~, or root-shoot interactions (Heppell et al., 2016; Braghiere et al., 2020), and Mai et al. (2019) used a sequential implementation~~
~~of multiscale soil-rhizosphere-plant framework to evaluate the effects that a dry spell would have on the short-term soil carbon reactions, via shifts in plant processes. The results of our model were used to better understand diverging results observed for~~
680 ~~our selected calibration and scenarios. Our first hypothesis was that a dry spell would cause lower plant exudation. However, we observed that the drought could lead to both a decrease or an increase in cumulative exudation depending on the developmental stage of our simulated plant. Our second hypothesis was that the dry spell would lead to a lower microbial activity. For two~~
~~of our three microbial calibrations, changes in total exudation led to the same trend in the microbial development and CO₂ emission when compared with the multi-scale setup. Our framework stands out by coupling of a dynamic whole plant model~~
685 ~~with a dynamic soil model for both water and C flows (Ruiz et al., 2021; Rees et al., 2018; Schnepf et al., 2022), including biochemical and physical reactions (such as adsorption and the interactions with microbial communities) in the soil domain. The multi-scale framework is a further advantage of this fully open-source setup (Ruiz et al., 2021), increasing the resolution~~
~~in baseline scenario. For our last microbial parameter set *lowMUuptake*, water stress caused by the soil areas near the root-soil interface to balance accuracy and computing speed. Moreover, the use of fixed-point iteration with implicit time stepping~~
690 ~~allowed for larger time steps, increasing therefore not only the model accuracy but also its speed for specific scenarios. However, as the simulation of the water flow was the main bottleneck for model convergence, application of the model to more sandy soil will lead to shorter time steps and, therefore, longer computation time. There remains potential for further optimization the code in order to increase its computing speed. Expanding the framework presented by Schnepf et al. (2018b)~~
~~, this coupled model remains flexible, allowing the user to implement only the modules relevant to their research question.~~
695 ~~The high number of functions represented lead to a high number of parameters. However, several structural plant or root system parameter sets have been established for different species. Moreover analysis of the structural root system module were conducted by Schnepf et al. (2018a) and Morandage et al. (2022). For the soil biokinetics, Sirean et al. (2025) defined a parameter space that fulfills biogeochemical constraints identified from a literature study. They also conducted a sensitivity analysis of the outputs. A user-friendly implementation of the CPlantBox calibration method of Morandage et al. (2021) is~~
700 ~~currently being developed. Finally, this implementation of *TraiRhizo* in *DuMu^x* within CPlantBox sets the building blocks for the simulation of new processes, taking into account solutes such as phosphorus and nitrogen, and the biological, chemical or physical reactions they are subject to~~
~~dry spell was not compensated by higher plant exudation. Our last hypothesis was that drought could cause either an increase or a decrease in short-term carbon storage. For our simulations, we indeed observed~~

both an increase and a decrease in the soil organic carbon.

705

In the first section of this discussion, we look in more detail at the simulated exudation. The main results of our study, regarding the simulated soil carbon dynamic and its changes under drought conditions, are discussed in section 4.2.

4.1 Plant processes

In this implementation of the model, we simulated a simplified plant C ~~dynamic~~ dynamics compared with the one described
710 in the literature, while still being able to ~~re-create qualitative plant dynamic~~. ~~Simulation results indicate that plant expansion growth is strongly influenced by changes in water potential during the diurnal cycle and drought periods, consistent with processes described in previous studies (Verbančič et al., 2018; Barillot et al., 2020; Coussement et al., 2018; De Swaef et al., 2022).~~ However, it is hypothesised that cell wall biosynthesis (biosynthetic growth) predominantly occurs during the day using photoassimilates, while being much more limited at night when it relies on starch reserves (Verbančič et al., 2018). Although
715 our simulations did not explicitly separate biosynthetic growth from expansion growth, the parametrization of the plant C module allowed us to qualitatively reproduce these reproduce qualitative plant dynamics. For this simulation, the virtual plant's starch pool can be interpreted as representing both actual starch reserves and newly synthesised wall material, thereby enabling expansion growth to occur at night. To achieve robust results and better recreate typical C dynamics across a wider range of parameter sets, it could be beneficial to explicitly represent biosynthetic growth in future models. Following the analyses of
720 Verbančič et al. (2018) and Barillot et al. (2020), high biosynthetic growth requires higher C supply but results in the creation of new cells more flexible and capable of expanding under lower turgor pressure. This higher amount of cells with high cell wall flexibility can help maintain plant growth in case of drought (Blum, 2017). Another mechanism supporting growth under low turgor pressure is osmotic adjustment, which has been identified as a critical process during drought conditions (Blum, 2017; Coussement et al., 2018). Not including this process in our model could explain the high growth limitation caused
725 by drought observed in our simulation a more detailed discussion of the plant related processes, see section K3.

Regarding the plant's C release, exudation was found to be positively correlated with plant growth (Přikryl and Vančura, 1980; Canarini et al., 2019). In our study, we assumed that exudation occurs only at the growing root tip, making the two correlated on the diurnal and weekly scales for a specific weather scenario. However, lower growth allowed more C to be available for exudation, making the two anti-correlated on the hourly scale (day vs. night dynamics) and when comparing dryer against wetter scenarios.
730 Moreover, in experimental studies Like for our simulations, a decrease of rhizodeposits was ~~also~~ experimentally observed at night (Kuzyakov and Cheng, 2004). This dynamic was particularly strong in our simulations, as we observed no exudation in the middle of the night outside of the dry spell periods. However, we could still recreate qualitative observations made in other studies. Indeed, plants ~~to~~ regulate internal C concentrations and adjust C allocation for growth and maintenance under fluctuating environmental conditions by modulating starch storage (Tixier et al., 2023; Bazot et al., 2005) and C release rates (Prescott et al., 2020; Canarini et al., 2019). The Also, the upper range of our exudation rates per unit of root surface
735 ($\approx 0.16 \text{ mmol C cm}^{-2} \text{ d}$) was superior to the averaged rates given in the experimental study of Kravchenko et al. (2004) ($0.018 \text{ mmol C cm}^{-2} \text{ d}$) or Trofymow et al. (1987) ($0.01 \text{ mmol C cm}^{-2} \text{ d}$, as presented by Darrah (1991)). However, we

could find similar values in the study of Personeni et al. (2007, Fig3A), due in part to its higher temporal resolution: $\approx 0.68 \text{ mmol C cm}^{-2}$ under high light input and assuming an exuding root tip of 1 cm . The modelling study of Thorpe et al. (2011) for a dwarf bean plant yielded $\approx 0.69 \text{ mmol C cm}^{-2} \text{ d}$, assuming an exuding root tip of 1 cm with radius of 0.02 cm . Moreover, was within the range found in the literature (Kravchenko et al., 2004; Trofymow et al., 1987; Personeni et al., 2007; Thorpe et al., 2011). The microbial C uptake ~~acted~~ affected the exudation rate by acting as a C sink ~~and could affect the exudation rate~~ (Canarini et al., 2019). We also observed a change in the type of organic C released by the plant (in our model, exudation-to-mucilage release ratio) under dry spell (Bazot et al., 2005; Hartmann et al., 2020). In spite of those changes, mucilage remained between 1-10% of exudation, consistent with the observations of Nguyen (2009), and exudation represented between 3-40% of assimilated C (Dilkes et al., 2004; Lynch and Whipps, 1990).

Regarding the plant water balance, the outputs of the model differ from ~~that those~~ presented by Giraud et al. (2023), in part because of the ~~implementation of the multiscale setup with fixed-point iteration~~ change in our solver setup, in part because of the update of the plant parameters, as ~~decrib~~ described in section 2.3. After the update of the parameters, the starch reserves could be used more quickly by the plant when the assimilation rate became lower. This allowed for instance the plant to maintain its exudation rates at night. A further calibration of the plant carbon balance and its effect on the development of the surrounding microbes could be conducted thanks to study using positron emission tomography (Schultes et al., 2025).

4.2 Distribution of soil carbon

4.2 Soil processes

755 4.2.1 Spatial and temporal distribution of microbial functional groups

4.2.1 Resolution of the multiscale setup

Regarding the spatial resolution, we observed a smooth gradient of low-weight dissolved organic molecules-C (C_L^l) along the radial axis for the different biokinetic parametrisation. This suggests that our resolution for the microscale domains is sufficient to represent the C flow in the 1D soil models. For simulations of solutes with low diffusion rates (e.g. phosphorus) and a lower advection (low plant water uptake rate), the root's uptake rate would become strongly dependent on the size of the 1D segment at the root-soil interface. A study could be conducted to evaluate the minimum number of segments necessary to accurately represent the selected soil processes according to the range of expected advection and diffusion rates of solutes under specific scenarios.

The output of the simulations showed diverging results according to the studied scales (~~macro~~ scale, ~~bulk soil~~, ~~perirhizal~~ trunc and microscale). For instance, compared with the bulk soil scale, the ~~perirhizal~~ trunc scale showed a higher influence of the weather scenarios on the soil C balance. Moreover, the microscale showed a high gradient of the C concentration along the radial coordinate of larger scale in the 3D soil, smaller scale with the 1D domains ~~that could not be represented via the perirhizal~~ trunc ~~scale~~ models). These scale-dependent results underline the advantage of using a multi-scale simulation and having a higher precision at the root-soil interface, following the observations of Mai et al. (2019). Moreover, the model

770 showed a strong horizontal and vertical variability in the concentrations, ~~making~~. For our setup, conducting a 3D evaluation remains relevant before, for instance, aggregating the results to a 2D or 1D sink term to be used by higher-scales models. As mentioned in the first implementation of the multiscale framework (Mai et al., 2019), the ~~low-high~~ resolution near the root soil interface (at the *mm* scale) raises the issue of including lower-scale processes, such as the influence of pore size on the water and C balance (Kuppe et al., 2022). As in the study of Landl et al. (2021a), the interactions between the 1D and 3D soil
775 models also allowed us to represent the additive effect of perirhizal zone proximity, which led to higher solute concentration under specific ~~biokinetic-kinetic~~ parameterisation (*lowMUptake*) and favored C hotspot formation from plant releases. Landl et al. (2021a) also evaluated the influence of root traits on the soil's normalised hotspot volume. Our study complements those findings and shows that root and microbial traits have to be evaluated together, as high concentration of soil organic C can be reached for different exudation rates according to the traits of the microbial community.

780

~~Other characteristics of the microbial activity were studied with our model. With the biokinetic-~~

4.2.2 Microbial starving-survival lifestyle

With the kinetic parameter sets *highCO2* and *highMB*, we observed a diurnal dynamic of high microbial growth under higher exudation (daytime), leading to a high consumption of C for microbial growth and maintenance. This was followed by microbial
785 dormancy (high dormant-to-total microbe ratio at nighttime). This dynamic has been observed in the literature and can be described as a "starving-survival lifestyle", where microbes enter dormancy in a low-nutrient environment and quickly react to new inputs of resources (Hobbie and Hobbie, 2013). As highlighted by Hobbie and Hobbie (2013), this strategy represents the more common microbial strategy in natural systems.

790 Although our results depend strongly on modeling choices, this correspondence suggests that the outputs obtained with *highCO2* and *highMB* may be more readily generalisable to real-world conditions.

Despite substantial differences in parameter values, the overall trends remained consistent between these two parameters (but not for the quantitative results). This stability may indicate the presence of conservative system behavior maintained by interactions among processes across a wide parameter space. A future study will include a comprehensive statistical analysis to more rigorously evaluate the influence of parameter variation on model outputs.

~~We found a specific reaction to resource scarcity for the parameter set-~~

4.2.3 Microbial diversity

For *highMB* with *lateDry*. For this simulation *lateDry*, we found an especially high copiotroph growth and a high ratio of
800 ~~dormant oligotrophs. This dormant-to-total oligotrophs. These~~ diverging results between the two communities are indicative of a possible intra-microbial competition for this simulation. This follows over studies where plant exudation and root growth affected microbial composition (de la Fuente Cantó et al., 2020; Bonkowski et al., 2021). Although only two functional groups

were represented, the higher development of the copiotrophs compared with the oligotrophs at the root-soil surface mimicked the lower microbial alpha-diversity observed experimentally in the rhizosphere (Kuzyakov and Razavi, 2019).

805 ~~Biokinetic-~~

4.2.4 Priming effect

Kinetic parameter sets describing more active microbes (*highCO2* and *highMB*) were also linked to a higher usage of soil solutes in case of higher exudation, when compared with the *baseline* scenarios. This higher usage rate led to a form of "rhizosphere priming effect" (Bonkowski et al., 2021; Kuzyakov and Cheng, 2004) where the root exudation ~~caused~~ can
810 cause an increase in mineralisation of the pre-existing soil organic matter, leading ultimately to a relatively lower amount of C in the solute pool ~~compaired~~ compared with the *baseline*. A more precise evaluation of a possible rhizosphere priming effect could be conducted with this setup, by dividing each C pool between the C originating from the plant additions and the C already present in the system at the beginning of the simulation. ~~Our simulation of the microbial dynamic and competition was however limited by not accounting for other stresses than C and water scarcity, such as nitrogen, phosphorus~~
815 ~~(Drake et al., 2013; Brown et al., 2022), or oxygen (Wiesenbauer et al., 2024) scarcity. Likewise, the diurnal aspect of microbial activity would also be better represented by accounting for soil temperature (Kuzyakov and Cheng, 2004). However, over the time period of this study (two weeks), we assumed the soil temperature varied little. Moreover, in our simulation, the slower decomposing organic carbon pool was not explicitly represented (see section 2.4.4), and we assumed that the pool would vary little over the short time scale represented. Including explicitly this carbon pool would allow for a more complete analysis of~~
820 the carbon allocation and (increased) usage by the microbial community.

4.2.5 Microbial spacial distribution

Regarding the spatial ~~distribution of the microbes, the prescribed exudation near the plant root tips also led to a higher microbial growth in those areas, resulting in a vertical distribution similar to the one observed by McDougall and Rovira (1970).~~
resolution, we observed an effect of the root system structure and growth rate on the C concentration profile. The C concentration
825 peaks observed for *earlyDry* and *lateDry* at deeper depth were due to lower root growth: The root tips remained in that area of soil over a longer period, leading to higher concentrations. Stronger plant growth under *baseline* resulted in perirhizal zones and rhizodeposition at lower depths such that the C accumulation at the root tips was much less pronounced compared to the dry spell scenarios. Moreover, for some of our scenarios, the high concentration of C pools at the outer boundary of our 1D model (*lowMUptake*)
830 underlined the additive effect on the profile of the low-weight organic molecules-C due to proximity to the perirhizal zone, which can indicate an overlap of the rhizospheres. This proximity effect was less noticeable for *highCO2* and *highMB*, where we observed concentration profiles less dependent on the extent of the 1D domain. Indeed, with those kinetic parameter sets, the microbial C uptake was highest and reacted fastest to high influxes of dissolved low molecular weight organic carbon. Therefore, the low-weight dissolved organic molecules-C concentration was closer to an asymptote at the outer boundaries of the 1D domain. These steeper gradients can cause the rhizosphere extent to be smaller, limiting the
835 rhizosphere overlaps.

At the microscale 1D scale, simulations also resulted in very high concentrations of copiotrophs near the root surface. Although high copiotrophic concentration near the root surface following exudation is expected (Bonkowski et al., 2021), representing microbial motility (Kuppe et al., 2022; Schnepf et al., 2022) would help smooth these concentration peaks, especially for older root segment with decreasing exudation and water uptake rates. Representing motility could also potentially lead to a better
840 representation of the priming effect as the microbes (especially from mature roots) might diffuse and consume C in areas further away from the root-soil interface (Dupuy and Silk, 2016).

4.2.6 ~~Effect of dry spells on carbon stabilisation~~

~~The biokinetic-~~

845 4.2.6 Sensitivity to water stress and carbon scarcity

The kinetic parameterisation strongly influenced the effects of the weather scenarios on soil C balance. ~~For more active biokinetic-~~

Our more active kinetic parameter sets (*highCO2* and *highMB*), the microbial growth was limited by C resources. The higher plant C releases and lower soil water content *lateDry* had therefore a positive effect on the microbial community. On
850 the contrary, for our non-C limited microbial communities (*lowMUptake* parameter set), the water stress of *lateDry* ~~affected negatively negatively affected~~ the microbial growth. These opposite reactions influenced how the soil domain was impacted by the short dry spells (see below). Likewise, because of their lower potential maximum C uptake, oligotrophs profited more from wetter conditions.

The microbial ~~dynamic-dynamics~~ will have an even stronger impact on the simulated water and C balance once soil-to-plant
855 feedback mechanisms are implemented, such as nutrient uptake (de la Fuente Cantó et al., 2020) and organic C-dependent soil hydraulic parameters (Landl et al., 2021b; de la Fuente Cantó et al., 2020). Indeed, among other effects, mucilage increases the water holding capacity of the soil and can maintain hydraulic conductivity around the roots (Landl et al., 2021b; Carminati et al., 2016). This change of the hydraulic characteristics of the soil can affect the plant's capacity to withstand drought conditions.

860 4.2.7 Effects of dry spells

We could reproduce qualitatively processes measured by Deng et al. (2021) for the C balance under limited water resources. Indeed, we observed a decrease in root biomass during the dry spells, with, under some scenarios, an increased exudation. The lower root growth then led to lower plant respiration, while the higher concentration of dissolved organic C under the late dry spell (with *highCO2* and *highMB*) led to a higher microbial respiration by the end of the simulation. Contrary
865 to Deng et al. (2021), we did not observe a decrease in total soil organic C. They linked this decrease to a lower soil C addition from plant residue, which is not represented in our model. Under *lateDry*, we could observe (limited) differences in

concentration compared with the *baseline* seven days after the end of the dry spells, underlining the resilience of the simulated environment, especially for the soil zones nearer the roots.

4.2.8 Wetting-drying cycles

870 Our ~~*earlyDry* and *lateDry*~~ *earlyDry* and *lateDry* weather scenarios led to a short decrease in soil water content, creating a wetting-drying cycle. According to the meta-analysis of Borken and Matzner (2009), these ~~wetting-drying-wetting-drying~~ wetting-drying-wetting-drying cycles should lead, under specific environmental conditions and microbial communities, to lower microbial respiration during the spells, followed by a respiration pulse once the soil is rewetted (Birch effect). These pulses may or may not compensate for the earlier CO₂ emission decrease (Borken and Matzner, 2009). In our simulation, the early dry spell always led to a decrease
875 in overall emitted microbial CO₂ at the ~~macroseale 3D scale~~ macroseale 3D scale when compared with the *baseline* scenarios. We then observed in the week following the end of the spell a slight uptick in emitted CO₂ compared with the baseline, which did not compensate the lower earlier emissions. For the later dry spells with *highCO2* and *highMB* ~~parametrisation~~ parameterisation, the emission pulses occurred during the spells. Indeed, while the drying led to a lower portion of microbes being dormant, the soil water content often remained too high to cause high microbial death or deactivation, especially for soil zones further away from
880 the roots (~~macroseale 3D scale~~ macroseale 3D scale). The lower water stress was, therefore, partly compensated by the higher C_T^l concentrations of low-weight dissolved organic molecules-C. We did not simulate an increase of plant and microbial necromass to be used by the soil microbes at the end of the spell (increasing thus the respiration once the soil is rewetted). Another explanation for the divergence is that we do not represent the easier access to "previously protected organic matter" caused by sudden soil rewetting. Moreover, we represented short dry spells (seven days) while longer spells (two weeks) may be necessary to
885 see stronger effects on the microbial community (Borken and Matzner, 2009). The higher C mineralisation during the drying phase of the later dry spell could also be linked to the method used to represent microbial sensitivity to water stress. Indeed, for simplicity, we did not represent the direct effect of soil water content on the potential microbial death (k_{max}). Moreover, we used the same value to calibrate the sensitivity of copiotrophs and oligotrophs to water stress, although oligotrophs could be assumed to be more resistant to low soil water content. Finally, the water scarcity parameters could be recalibrated using
890 experimental data.

~~We could reproduce qualitatively processes measured by Deng et al. (2021) for the C balance under limited water resources. Indeed, we observed a decrease in root biomass during the dry spells, with, under some scenarios, an increased exudation. The lower root growth then led to lower plant respiration, while the higher concentration of dissolved organic C under the late dry spell (with *highCO2* and *highMB*) led to a higher microbial respiration by the end of the simulation. Contrary
895 to Deng et al. (2021), we did not observe a decrease in total soil organic C. They linked this decrease to a lower soil C addition from plant residue, which is not represented in our model. Under *earlyDry*, we could observe (limited) differences in concentration compared with the *baseline* seven days after the end of the dry spells, underlining the resilience of the simulated environment, especially for the soil zones nearer the roots. The *lateDry* scenario led to the lowest plant growth but to a higher SOC hotspot volume, indicating a more resource-efficient root system exudation. We had, therefore,~~

900 4.2.9 Bottlenecks and future development

In theory, two non-microbial organic carbon pools can be used to represent the soil organic carbon (Sircan et al., 2025; Kuppe et al., 2022). However, for this model setup, a trade-off in the C-usage strategy third inert carbon pool was needed to accurately represent the C dynamic of slow decomposing organic matter. The reason for this limitation was twofold. First of all, to represent the movement of mucilage, our high-molecular weight organic C undergoes diffusion. Therefore, we only represent labile organic carbon. Moreover, our soil organic carbon concentration results (even in the perirhizal zone) were below the organic C concentration observed in Selhausen (Wang et al., 2018) and by Poeplau and Don (2023) for healthy bulk soil. This discrepancy was caused by our calibration. In their work Sircan et al. (2025) conducted a review of acceptable ranges for each of their output variables. However the objective of their work was not to conduct an extensive review of the literature for each variable and their maximum organic C concentration values were obtained from two studies conducted in non-German soils. Conducting a more precise calibration for a specific experimental setup and introducing a non-labile carbon pool would allow the model to better represent the soil carbon dynamic. The implicitly represented slow decomposing soil organic carbon was assumed to be inert over the duration of the study, as our simulation time was of two weeks. However, not representing this third pool limits the generalisation that can be made of our results. For instance, the Birch effect discussed below is also caused by the destabilisation of this slow decomposing carbon pool after re-wetting of the soil (Borken and Matzner, 2009). This added pool also shifted the fractions of soil belonging to the different hotspot categories. For this reason, the thresholds of Poeplau and Don (2023) were presented as example categories to compare the scenario and evaluate changes over time rather than absolute value indicative of soil health.

Our simulation of the microbial dynamics and competition could also be improved by accounting for other stresses than C and water scarcity, such as nitrogen, phosphorus (Drake et al., 2013; Brown et al., 2022), or oxygen (Wiesenbauer et al., 2024) scarcity. Likewise, the diurnal aspect of microbial activity would also be better represented by accounting for soil temperature (Kuzyakov and Cheng, 2004).

5 Conclusions

In this paper, we presented the equations and implementation of a coupled model representing ~~the carbon (C)~~ carbon and water flow in the soil-rhizosphere-plant continuum, influenced by ~~the atmospheric conditions via~~ atmospheric conditions through plant transpiration and photosynthesis. This ~~new framework takes into account the effect of the~~ framework accounts for the effects of water content variation on ~~the C~~ carbon flow and microbial reactions. ~~Moreover, the multiscale implementation allowed us to evaluate more precisely the water and C flow activity. The multiscale implementation enables precise evaluation of fluxes and reactions at the soil-plant interface, quantifying the feedback effect of the plant on the soil processes. Thanks to the outputs of the model, we could observe the evolution of variables difficult to measure experimentally at the rhizosphere scale with non-destructive measurements~~ and can capture feedback across domains and scales. Despite the simplified representation of plant and microbial processes and limited calibration, the model reproduces trends reported in the literature, such as ~~exudation and soil respiration. In our simulation, the~~ “starving-survival” lifestyle of some microbial communities. We found

that the effect of ~~dry spells on the soil C balance depended on the time dynamic of~~ water scarcity on soil carbon turnover is an emergent property arising from 1D scale feedbacks between plants, microbes, and local environmental conditions. The varied impacts of water stress highlight the strong influence of local soil and plant characteristics in determining whether carbon or water limitation dominates turnover. By representing multiple sources of plant and microbial stress, the ~~exudation and the characteristics~~ model provides a mechanistic explanation for variability in plant and soil carbon allocation under combined stresses. The model yields a wide range of variables (e.g., short-term carbon storage) across the soil-rhizosphere-plant continuum that can be used to assess the performance of plant phenotypes and management measures under dynamic conditions. By computing the resulting soil and plant conditions and identifying the key processes driving ecosystem responses, this model can both provide parameters for larger-scale models and inform the design of simplified models that focus on the most influential processes. Moreover, this work lays the foundation for a more comprehensive model of the plant-soil interaction cycle, including nutrient exchanges, which are essential to accurately represent the feedback of ~~the microbial community, underlying the site-specific aspect of agricultural practices' effects on the C balance.~~ For instance, compared with the baseline scenarios, ~~dry spells could lead to an increase of C mineralisation with active microbial communities (C-limited) against lower mineralisation for less active microbial communities (water-limited).~~ Adding nutrient (such as nitrogen) flows in the coupled model would be an important extension to represent more accurately the feedback effect of the soil C balance on the plants' development soil on plant processes. Finally, this model is particularly well suited to reproduce the experimental observations of isotopic carbon allocation in the plant and soil.

950 6 Code availability

. For this study, we used the plant model CPlantBox (Schnepf et al., 2025b) and the corresponding soil module (Schnepf et al., 2025a). The full setup is available at: Giraud2025_C_W_Rhizo

7 Video supplement

. A video presenting this graphic for each time step is available in the supplementary documents ("Video S1"). The supplementary video "Video S2" shows the low-weight dissolved organic molecules-C concentrations and the corresponding active-to-total copiotrophs-C ratios at each time step.

Appendix A: List of the parameters and variables

Table A1. Represented carbon pools

Output variables	Definition	Unit
C_L^l/C_L^s	Concentration of carbon from compounds with low molecular weight (in the liquid/solid phase)	$mol\ C\ cm^{-3}\ water /$ $mol\ C\ cm^{-3}\ scv$
C_H^l/C_H^s	Concentration of carbon from compounds with high molecular weight	$mol\ C\ cm^{-3}\ water$
$C_O (C_O^a, C_O^d)$	Concentration of oligotrophs (active, dormant)	$mol\ C\ cm^{-3}\ scv$
$C_C (C_C^a, C_C^d)$	Concentration of copiotrophs (active, dormant)	$mol\ C\ cm^{-3}\ scv$
CO_2	Unit of emitted CO_2 per volume of soil	$mol\ C\ cm^{-3}\ scv$

Appendix B: Soil water flow

In the sections below, we present the implementation of the equations given in section 2.2.2 to the 1D axisymmetric and 3D soil domains.

B1 3D soil model

For the 3D ~~macro~~soil model, the sink term of Eq. (6) $S_\theta = S_{w,root-3DS}$ ($cm^3\ water\ cm^{-3}\ scv\ d^{-1}$), the water uptake or release by the roots:

$$S_{w,root-3DS,init} = \frac{1}{V_{3DS,scv}} \sum_{i=1}^{nr} \frac{\partial \theta_i V_{1DS,scv,i}}{\partial t} \quad (B1)$$

$$S_{w,root-3DS} = \mathbf{Lim}(S_{w,root-3DS,init}) \quad (B2)$$

where nr (–) is the number of root segments in the soil control space of volume $V_{3DS,scv}$ ($cm^3\ scv$), the i subscript is the identification of the i^{th} root, $V_{1DS,scv,i}$ ($cm^3\ scv$) is the volume of the perirhizal zone i and θ_i its water content. Instead of using $q_{w,root-soil}$ (see Eq. (3)) like Mai et al. (2019), in this study $S_{w,root-3DS}$ is obtained from $\frac{\partial \theta_i V_{1DS,scv,i}}{\partial t}$. This helps avoid computation errors when simulating the ~~macro-soil~~ 3D-soil flows and diminishes the divergence between the ~~macro- and microscale models~~ 3D- and 1D-models. Indeed, the plant-prescribed $q_{w,root-soil}$ was already potentially limited during

Table A2. ~~TraiRhizo~~ ~~input~~ ~~Input~~ parameters [for the soil model](#). When the source is not given, the parameter values are taken from the sets defined by Sircan et al. (2025) and selected in section I. HMW: high molecular weight organic carbon compound.

Symbol	Definition (and source)	Value per selected parameter set			Unit
		<i>highCO2</i>	<i>highMB</i>	<i>lowMUptake</i>	
a	Sharpness parameter for the switch function from active to dormancy	0.1			–
$C_{H,init}^l$	Initial concentration of high molecular weight organic carbon concentration	$5.94e^{-4}$	$6.94e^{-4}$	$6.47e^{-4}$	$mol\ C\ cm^{-3}$ <i>water</i>
$C_{L,init}^l$	Initial dissolved low molecular weight organic carbon	$4.71e^{-6}$	$5.15e^{-6}$	$5.1e^{-6}$	$mol\ C\ cm^{-3}$ <i>water</i>
$C_{L,max}^s$	Maximum sorption capacity	$8.9e^{-4}$			$mol\ C\ cm^{-3}$ <i>scv</i>
$C_{thres,C}$	Threshold C_L^l for microbial (de)activation	$1.42e^{-5}$	$9.90e^{-6}$	$2.19e^{-5}$	$mol\ C\ cm^{-3}$ <i>water</i>
$C_{thres,O}$		$8.58e^{-6}$	$7.71e^{-6}$	$9.76e^{-6}$	
D_{HW}	Diffusion coefficient of high molecular weight organic carbon	$3.456e^{-3}$			$cm^2\ d^{-1}$
D_{LW}	Diffusion coefficient of dissolved low molecular weight organic carbon	2.07	0.92	1.45	$cm^2\ d^{-1}$
k_{ads}	Adsorption rate	$2.7e^{-7}$			$cm^3\ scv$ $mol\ C^{-1}\ d^{-1}$
$k_{d,C}$	Deactivation rate coefficient	2.73	1.49	1.37	d^{-1}
$k_{d,O}$		0.24	4.04	0.11	
k_{des}	Desorption rate	$5e^{-5}$			d^{-1}
$k_{C,S}$	Substrate affinity to dissolved low molecular weight organic C	$4.93e^4$	$1.50e^5$	$7.72e^4$	$mol\ C\ cm^{-3}$ <i>scv</i>
$k_{O,S}$		$1.02e^6$	$1.16e^7$	$9.55e^4$	
K_L	Half-saturation coefficients of enzymes targeting HMW	$1.98e^{-6}$	$1.02e^{-7}$	$6.23e^{-4}$	$mol\ C\ cm^{-3}$ <i>scv</i>
$k_{max,C}$	Maximum maintenance rate coefficient	$7.07e^{-3}$	$1.38e^{-3}$	$1.46e^{-3}$	d^{-1}
$k_{max,O}$		$2.30e^{-4}$	$6.57e^{-4}$	$1.14e^{-3}$	
$k_{r,C}$	Reactivation rate coefficient	86.3	95.9	54.3	d^{-1}
$k_{r,O}$		1.17	54.9	0.181	
$v_{max,depoly}$	Maximum reaction rate of enzymes targeting HMW	0.108	$1.31e^{-2}$	$1.69e^{-3}$	d^{-1}

Table A2. Continued

Symbol	Definition (and source)	Value per selected parameter set			Unit
		<i>highCO2</i>	<i>highMB</i>	<i>lowMUptake</i>	
Y	Maintenance yield	0.372	$2.54e^{-2}$	0.397	–
Y_C	Growth yield on dissolved low molecular weight organic carbon	$1.08e^{-2}$	0.108	0.422	–
Y_O		$2.43e^{-2}$	0.142	0.896	
α	Empirical parameter determining the curvature of Eq.D10 (Moyano et al., 2013)	1.47			–
β_C	Reduction factor of maintenance requirements in dormant state	$1.64e^{-3}$	$5.13e^{-3}$	$1.09e^{-4}$	–
β_O		0.592	$9.32e^{-2}$	$3.31e^{-2}$	
$\mu_{max,C}$	Maximum growth rate coefficient	0.83	1.42	0.25	d^{-1}
$\mu_{max,O}$		$1.03e^{-2}$	$3.56e^{-2}$	$1.26e^{-2}$	
ρ_L	Proportion of high molecular weight organic carbon formed from dead microbial biomass due to maintenance	$2.16e^{-2}$	0.579	$1.51e^{-2}$	–
ρ_b	Soil bulk density	1.51			<i>g mineral soil</i> <i>cm⁻³ scv</i>
τ	$\tau = \frac{\psi_{m,soil,D2A}}{\psi_{m,soil,A2D}}$, with $\psi_{m,soil,D2A}$ the threshold soil matric potential for microbial activation (Wang et al., 2021)	0.39			–
$\psi_{m,soil,A2D}$	Threshold soil matric potential for microbial deactivation (Wang et al., 2021)	4600			<i>hPa</i>
$\psi_{m,soil,opt}$	Threshold soil matric potential above which water is not limiting for microbial activity (Moyano et al., 2013)	–30			<i>hPa</i>
$\psi_{m,soil,th}$	Threshold soil matric potential bellow <u>below</u> which microbial activity ceases (Moyano et al., 2013)	–158000			<i>hPa</i>
ω_{DA}	Empirical exponent for Eqs.D29,D30 (Wang et al., 2021)	3.38			–

the computation of $\frac{\partial \theta_i V_{1DS,scv,i}}{\partial t}$ according to $\psi_{x,crit}$ (see Eq. (B12)). We then set a second limitation on the net sink (resp. source) using the maximum potentially available water (resp. space) in the voxel—equal to the water (resp. space) volume at the beginning of the time step and to the maximum amount of water that could be gained (resp. lost) according to the inter-voxel flow at the last time step. The driving equation for the water flow is defined thus at the [macro-scale-3D scale](#) (Mai et al., 2019; 975 Koch et al., 2021):

$$\frac{\partial \theta}{\partial t} = K_{sat} \nabla \cdot (\kappa_m (\nabla \psi_{m,3DS} - \nabla \psi_{g,3DS})) + S_{w,root-3DS} \quad \text{for } \Omega \setminus \partial\Omega, t > t_0 \quad (\text{B3})$$

$$S_{w,root-3DS} = S_{w,root-3DS,0} \quad \text{at } t = t_0 \quad (\text{B4})$$

A boundary condition completes the model and can represent, for instance, the net gain or loss of water resulting from rainfall and soil evaporation.

980 B2 1D axisymmetric soil models

Because of the small radii of the [micro-scale-1D](#) domains, we assume that the gravitational gradient along the radial coordinate is negligible compared with the matric potential gradient: $abs(\frac{\partial \psi_{g,1DS}}{\partial r}) \ll abs(\frac{\partial \psi_{m,1DS}}{\partial r})$. We thus get (Debnath, 2005, Eq. 1.10.4):

$$\frac{\partial \theta}{\partial t} = \frac{1}{r} \frac{\partial}{\partial r} (r K_{sat} \kappa_m \frac{\partial \psi_{m,1DS}}{\partial r}) + \frac{1}{r} S_{w,3DS-1DS} \quad (\text{B5})$$

985 with r (cm) the radial coordinate along the axisymmetric domain, where $r = 0$ corresponds to the center of root segment associated with the [micro-scale-1D](#) soil domain. $r \in [r_{in}, r_{out}]$, with r_{in} (cm) the inner boundary of the domain, corresponding to the root radius, and r_{out} (cm) the outer boundary of the 1D domain.

The water flow at the inner boundary of 1D domain corresponds to $q_{root-soil}$ (see Eq. (3)). $S_{w,3DS-1DS}$ is obtained from the [bulk-soil-water-flow-water flow in the 3D soil](#). For the segment i in the voxel k :

$$990 \quad S_{w,1DS-3DS,i} = - \frac{Q_{w,1DS-3DS,i}}{V_{1DS,scv,i}} \quad (\text{B6})$$

$$Q_{w,1DS-3DS,i} = - Q_{w,3DS-3DS,k} \cdot W_w(i, k) \quad (\text{B7})$$

$$W_w(i, k) = \begin{cases} \frac{(\theta_s - \theta_i) V_{1DS,scv,i} + Q_{w,root-soil,i} dt}{\sum_j^n (\theta_s - \theta_j) V_{1DS,scv,j} + Q_{w,root-soil,j} dt} & \text{if } Q_{w,3DS-3DS,k} > 0 \\ \frac{(\theta_i - \theta_r) V_{1DS,scv,i} - Q_{w,root-soil,i} dt}{\sum_j^{n^*} (\theta_j - \theta_r) V_{1DS,scv,j} - Q_{w,root-soil,j} dt} & \text{else} \end{cases} \quad (\text{B8})$$

$$V_{1DS,scv,i} = \pi (r_{out,i}^2 - r_{in,i}^2) L_{r,i} \quad (\text{B9})$$

$$Q_{w,root-soil,i} = q_{w,root-soil,i} 2\pi r_{in,i} L_i \quad (\text{B10})$$

995 $Q_{w,3DS-3DS,k}$ (cm^3 water d^{-1}) corresponds to the net water change in the sub-control volume caused by the exchange with neighbouring voxels. The second element on the right-hand side of Eq. (B7), $W_w(-)$ is a weighting factor dividing

the net water flow between the ~~microscale domains of the macroscale~~ 1D models of the 3D voxel k . W_w is computed from the maximal potentially available water (resp. space), which is the sum of [a] the net root water release (resp. uptake) over the evaluated period, and [b] the water (resp. air) volume available at the beginning of the evaluated period. In Eq. (B6), the volume of the ~~microscale~~ 1D domain ($V_{1DS,scv}$, cm^3 scv) is used to convert the net water sink from cm^3 $water$ d^{-1} to cm^3 $water$ cm^{-3} scv d^{-1} . $S_{w,1DS-3DS,i}$ is then divided between the 1D segments of the ~~microscale~~ 1D domains using the same weighting method. We obtain the following driving equation for the ~~microscale~~ 1D soil domains (Mai et al., 2019; Koch et al., 2021):

$$r \frac{\partial \theta}{\partial t} = \frac{\partial}{\partial r} \left(r K_{sat} \kappa_m \frac{\partial \psi_{m,1DS}}{\partial r} \right) + r S_{w,1DS-3DS} \quad \text{for } r \in]r_{in}, r_{out}[, t > t_0 \quad (B11)$$

$$K_{sat} \kappa_m \frac{\partial \psi_{m,1DS}}{\partial r} = \max(q_{w,root-soil}, K_{sat} \kappa_m \frac{(\psi_{crit,x} - \psi_{m,1DS})}{\Delta r_{root-1DS}}) \quad \text{at } r = r_{in} \quad (B12)$$

$$K_{sat} \kappa_m \frac{\partial \psi_{m,1DS}}{\partial r} = 0 \quad \text{at } r = r_{out} \quad (B13)$$

$$S_{w,3DS-1DS} = S_{w,3DS-1DS,0} \quad \text{at } t = t_0 \quad (B14)$$

with $\psi_{crit,x}$ (hPa) the critical (minimum) xylem water potential. Therefore, if both $\psi_{m,soil}$ and $q_{w,root-soil}$ are low, the realised net water uptake might be more limited than prescribed via $q_{w,root-soil}$.

Appendix C: Soil carbon transport

In the sections below, we present the implementation of the equations given in section 2.2.4 to the 1D axisymmetric and 3D soil domains.

C1 3D soil model

For the 3D ~~macroscale~~ soil domain, S_X can be computed in two ways:

1. If there are no roots in the voxel i ($len(nr_i) = 0$), S_X corresponds to the sources computed at the voxel scale ($S_{X,3DS}$, mol cm^{-3} scv d^{-1}).
2. If there is at least one root in the voxel, S_X corresponds to the sources computed ~~at the macroscale~~ by the 1D model ($S_{X,1DS}$, mol cm^{-3} scv d^{-1}).

This method is represented by the following equations:

$$S_X = \begin{cases} \frac{1}{V_{3DS}} \sum_{i=1}^{nr_k} V_{1DS} S_{X,1DS} & \text{if } len(nr_k) > 0 \\ S_{X,3DS} & \text{else} \end{cases} \quad (C1)$$

This allows us to achieve similar results [at the micro- and macroscales with the 1D and 3D models](#) and to profit from the higher resolution of the [microscale-1D](#) models, leading to more accurate results in the soil voxels containing roots. $S_{X,1DS}$ is computed similarly to $S_{w,root-3DS}$ (see Eq. (B1)), by computing the mean C variation rate and limiting the sink according to the C potentially available in the voxel. The driving equation for dissolved low molecular weight organic C is (Mai et al., 2019):

$$\frac{\partial \theta C_L^l}{\partial t} - K_{sat} \nabla \cdot (\kappa_m C_L^l (\nabla \psi_m - \nabla \psi_g)) \quad (C2)$$

$$-\nabla \cdot (D_L(\theta, \phi) \nabla C_L^l) - S_L = 0 \text{ for } \Omega \setminus \partial\Omega, t > t_0,$$

$$C_L^l = C_{L,0}^l, \text{ at } t = t_0 \quad (C3)$$

1030 For high molecular weight organic C compounds, we have:

$$\frac{\partial \theta C_H^l}{\partial t} - \nabla \cdot (D_H(\theta, \phi) \nabla C_H^l) - S_H = 0 \quad (C4)$$

$$\text{for } \Omega \setminus \partial\Omega, t > t_0,$$

$$C_H^l = C_{H,0}^l, \text{ at } t = t_0 \quad (C5)$$

For both equation sets, the boundary conditions can be used to represent an input of organic matter in the soil.

1035

C2 1D axisymmetric soil models

For 1D axisymmetric [microscale](#) soil domains, we get (Mai et al., 2019):

$$\frac{\partial \theta C_X^l}{\partial t} = -\frac{1}{r} \frac{\partial}{\partial r} r C_X^l \mathbf{u}_X + \frac{1}{r} \frac{\partial}{\partial r} r (D_X(\theta, \phi) \frac{\partial C_X^l}{\partial r}) + S_X + S_{1DS-3DS,X} \quad (C6)$$

$S_{1DS-3DS,X}$ represents the net source of solutes from inter-voxel exchanges [at the macroscale of the 3D soil model](#). For the segment i in the voxel k :

1040

$$S_{1DS-3DS,X,i} = -\frac{Q_{X,1DS-3DS,i}}{V_{1DS,scv,i}} \quad (C7)$$

$$Q_{X,1DS-3DS,i} = -Q_{X,3DS-3DS,k} W_X(i, k) \quad (C8)$$

$$W_X(i, k) = \begin{cases} \frac{m_{X,pot,i}}{\sum_j^n m_{X,pot,j}} & \text{if } Q_{X,3DS-3DS,k} < 0 \\ \frac{1/\max(m_{X,pot,i}, \epsilon)}{(\sum_j^n 1/\max(m_{X,pot,j}, \epsilon))} & \text{else} \end{cases} \quad (C9)$$

$$m_{X,pot,i} = \theta_i V_{1DS,scv,i} C_{X,i}^l + Q_{X,root-soil,i} dt \quad (C10)$$

$$1045 \quad Q_{X,root-soil,i} = 2\pi r_{in,i} L_i \cdot \begin{cases} q_{exud,i} & \text{if } X \text{ is } C_L^l \\ q_{mucil,i} & \text{if } X \text{ is } C_H^l \end{cases} \quad (C11)$$

$m_{X,pot}$ ($mol\ C$) is the potentially available solute content. $Q_{X,3DS-3DS,k}$ ($mol\ C/d$) corresponds to the net solute transport into the sub-control volume from other soil voxels. The second element of Eq. (C8) is a weighting factor dependent on the solute content of the **microscale-1D** domains in the voxel. $V_{1DS,scv,i}$ is used to convert the net solute transport from $mol\ d^{-1}$ to $mol\ cm^{-3}\ d^{-1}$. For solutes we could have $m_{X,pot} = 0$, we, therefore, set a minimum value of $\epsilon \approx 0$ when using $C_{X,pot}^q$ to avoid a division by 0. The same method is used to divide the net sinks between the **sub-control volumes of the 1D segments of the microscale** domain.

Eq. (C6) for dissolved low molecular weight organic C gives:

$$r \frac{\partial \theta C_L^l}{\partial t} - \frac{\partial}{\partial r} (r C_L^l K_{sat} \kappa_m(S_w) \frac{\partial \psi_m}{\partial r}) - \frac{\partial}{\partial r} r (D_L(\theta, \phi) \frac{\partial C_L^l}{\partial r}) - r (S_L + S_{1DS-3DS,L}) = 0 \text{ for } \Omega \setminus \partial\Omega, t > t_0, \quad (C12)$$

$$K_{sat} \frac{\partial}{\partial r} \left(\kappa_m C_L^l \frac{\partial \psi_m}{\partial r} \right) + \frac{\partial}{\partial r} (D_L(\theta, \phi) \frac{\partial C_L^l}{\partial r}) = q_{exud,1DS-root} \text{ for } r = r_{in}, \quad (C13)$$

$$K_{sat} \frac{\partial}{\partial r} \left(\kappa_m C_L^l \frac{\partial \psi_m}{\partial r} \right) + \frac{\partial}{\partial r} (D_L(\theta, \phi) \frac{\partial C_L^l}{\partial r}) = 0 \text{ for } r = r_{out}, \quad (C14)$$

$$C_L^l = C_{L,0}^l, \text{ at } t = t_0 \quad (C15)$$

1060

Eq. (C6) for high molecular weight organic C compounds gives:

$$r \frac{\partial \theta C_H^l}{\partial t} - \frac{\partial}{\partial r} r (D_H(\theta, \phi) \frac{\partial C_H^l}{\partial r}) - r (S_H + S_{1DS-3DS,H}) = 0 \text{ for } \Omega \setminus \partial\Omega, t > t_0, \quad (C16)$$

$$\frac{\partial}{\partial r} (D_H(\theta, \phi) \frac{\partial C_H^l}{\partial r}) = q_{mucil,1DS-root} \text{ for } r = r_{in}, \quad (C17)$$

$$\frac{\partial}{\partial r} (D_H(\theta, \phi) \frac{\partial C_H^l}{\partial r}) = 0 \text{ for } r = r_{out}, \quad (C18)$$

$$C_H^l = C_{H,0}^l, \text{ at } t = t_0 \quad (C19)$$

The solute transport at the inner boundary of the domain corresponds to the exchange with the root (q_{exud} or q_{mucil} , see Eq. (16), (20)).

Appendix D: Soil reactions and microbial pools

1070 Microorganisms can be modeled as a system of ordinary differential equations in time, (Pot et al., 2022), representing the to SOC **mineralisaiton-mineralisation** and CO_2 release caused by microbes (Pagel et al., 2020). We can describe the microbial behaviour by looking at community-level microbial traits (Bardgett and Caruso, 2020): recent studies indicate that soil bacterial communities are dominated by relatively few taxa with strong environmental preferences. Therefore a focus on the functional

traits of dominant taxa can improve our understanding.

1075 The equations in this section were adapted from the ones setup by Sircan et al. (2025). They present the microbe-driven soil reactions. The microbial community is divided in two main groups: oligotrophs (slower development and higher substrate affinity) and copiotrophs (quicker development and lower substrate affinity). The main differences with the equations presented by Sircan et al. (2025) are the explicit representation of the soil water content and its influence on microbial processes, and the non-instantaneous C sorption. We present below the list of reactions which sum up to $S_{microbe,X}$ for according to the
 1080 concentration of the organic C ~~concentration~~ of dissolved low molecular weight ~~concentration~~ (C_L^l) and high molecular weight compounds (C_H^l), as well as the connected microbial pools:

$$S_{microbe,L} = S_{depoly} + (1 - p_L)S_{decay} - S_{growth,L} - S_{Muptake,L} \quad (D1)$$

$$S_{microbe,H} = -S_{depoly} + p_L S_{decay} \quad (D2)$$

with S_{depoly} the depolymerisation rate of high molecular weight organic C compounds, S_{decay} the solute gain from microbial
 1085 decay, $S_{growth,S}$ the dissolved low molecular weight organic C uptake for microbial growth, and $S_{Muptake,L}$ the dissolved low molecular weight organic C uptake for maintenance (all in $mol\ C\ C^{-3}\ scv\ d^{-1}$). p_L (mol/mol) is the proportion of high molecular weight organic C compounds to total C compounds formed from dead microbial biomass due to maintenance.

The reaction rates are dependent on the concentration of microbial pools. Like Pagel et al. (2020), we define four microbial pools in the solid, described by their C concentration in the soil phase C_Z^Y ($mol\ C\ microbes/cm^3\ scv$). Y stands either for a
 1090 (active microbes) or d (dormant microbes). Z stands for O (oligotrophesoligotrophs) or C (copiotrophescopiotrophs).

We define their respective variation thus:

$$\frac{\partial C_O^a}{\partial t} = S_{growth,O} - S_{deact,O} + S_{react,O} - \frac{1}{Y_M} S_{Mdecay,O}^a \quad (D3)$$

$$\frac{\partial C_O^d}{\partial t} = S_{deact,O} - S_{react,O} - \frac{1}{Y_M} S_{Mdecay,O}^d \quad (D4)$$

$$\frac{\partial C_C^a}{\partial t} = S_{growth,C} - S_{deact,C} + S_{react,C} - \frac{1}{Y_M} S_{Mdecay,C}^a \quad (D5)$$

$$1095 \quad \frac{\partial C_C^d}{\partial t} = S_{deact,C} - S_{react,C} - \frac{1}{Y_M} S_{Mdecay,C}^d \quad (D6)$$

$S_{growth,Z}$ corresponds to the growth of C_Z^a , $S_{decay,Z}^Y$ corresponds to the decay of $C_Z^{Y,s}$, $S_{deact,Z}$ represents the switch of C_Z^a to C_Z^d and $S_{react,Z}$ represents the switch of C_Z^d to C_Z^a . $S_{Mdecay,Z}^Y$ represents the loss of $C_Z^{Y,s}$ C because of maintenance. All

rates are in $\text{mol C cm}^{-3} \text{ scv d}^{-1}$. $Y_M(-)$ is the maintenance yield. The growth reaction rates are defined thus:

$$S_{\text{growth},L} = \frac{1}{Y_O} S_{\text{growth},O} + \frac{1}{Y_C} S_{\text{growth},C} \quad (\text{D7})$$

1100

$$S_{\text{growth},O} = f_A(\psi_{m,\text{soil}}) \frac{\mu_{\text{max},O} C_L^l k_{O,L}}{\mu_{\text{max},O} + C_L^l k_{O,L}} C_O^a \quad (\text{D8})$$

$$S_{\text{growth},C} = f_A(\psi_{m,\text{soil}}) \frac{\mu_{\text{max},C} C_L^l k_{C,L}}{\mu_{\text{max},C} + C_L^l k_{C,L}} C_C^a \quad (\text{D9})$$

$$f_A(\psi_{m,\text{soil}}) = \begin{cases} 1, & \psi_{m,\text{soil}} > \psi_{m,\text{soil,opt}} \\ 1 - \left(\frac{\log_{10}(\psi_{m,\text{soil}}/\psi_{m,\text{soil,opt}})}{\log_{10}(\psi_{m,\text{soil,th}}/\psi_{m,\text{soil,opt}})} \right)^\alpha, & \psi_{m,\text{soil}} \in [\psi_{m,\text{soil,th}}, \psi_{m,\text{soil,opt}}] \\ 0, & \psi_{m,\text{soil}} < \psi_{m,\text{soil,th}} \end{cases} \quad (\text{D10})$$

With $Y_Z(-)$ the growth yield of C_Z . $\mu_{\text{max},Z}$ ($1/d$) is the maximum growth rate of C_Z^a . $k_{O,L}$ ($\text{cm}^3 \text{ scv d mol}^{-1} \text{ C}$) is the C_Z^a affinity to C_L^l . $f_A(\psi_{m,\text{soil}})(-)$ defines the water limitation on the access of the microorganisms and enzymes to the solutes (Moyano et al., 2013). The maintenance decay rates are defined thus:

1105

$$S_{M\text{decay}} = S_{M\text{decay},O}^a + S_{M\text{decay},O}^d + S_{M\text{decay},C}^a + S_{M\text{decay},C}^d \quad (\text{D11})$$

$$S_{M\text{decay},O}^a = k_{\text{max},O} \overline{C_O^a} - S_{M\text{uptake},L,O}^a \quad (\text{D12})$$

$$S_{M\text{decay},O}^d = k_{\text{max},O} \beta_O C_O^d - S_{M\text{uptake},L,O}^d \quad (\text{D13})$$

$$S_{M\text{decay},C}^a = k_{\text{max},C} \overline{C_C^a} - S_{M\text{uptake},L,C}^a \quad (\text{D14})$$

1110

$$S_{M\text{decay},C}^d = k_{\text{max},C} \beta_C C_C^d - S_{M\text{uptake},L,C}^d \quad (\text{D15})$$

$$\overline{C_O^a} = \max(0., C_O^a - C_{O,\text{lim}}^a) \quad (\text{D16})$$

$$\overline{C_C^a} = \max(0., C_C^a - C_{C,\text{lim}}^a) \quad (\text{D17})$$

$k_{\text{max},Z}$ ($1/d$) is the maximum maintenance rate coefficient for C_Z . $\beta_Z(-)$ is the reduction factor of maintenance requirements in dormant state for C_Z . $C_{Z,\text{lim}}^a$ ($\text{mol C/cm}^3 \text{ scv}$) corresponds to the minimal value of C_Z^a below which the microbial pool can only be subjected to growth. This limitation allows us to have a [re-development](#) [redevelopment](#) of the pool even if we have no solute in the soil for a period of time. We use consequently the limited $\overline{C_Z^a}$ ($\text{mol C/cm}^3 \text{ scv}$) value for all reaction rates except for $S_{\text{growth},Z}$. The uptake rates are defined thus:

1115

$$S_{\text{uptake},L} = S_{\text{uptake},L,O}^a + S_{\text{uptake},L,O}^d + S_{\text{uptake},L,C}^a + S_{\text{uptake},L,C}^d \quad (\text{D18})$$

$$S_{\text{uptake},L,O}^a = f_A(\psi_{m,\text{soil}}) \frac{k_{\text{max},O} C_L^l k_{O,L}}{k_{\text{max},O} + C_L^l k_{O,L}} \overline{C_O^a} \quad (\text{D19})$$

1120

$$S_{\text{uptake},L,O}^d = f_A(\psi_{m,\text{soil}}) \frac{k_{\text{max},O} C_L^l k_{O,L}}{k_{\text{max},O} + C_L^l k_{O,L}} \beta_O C_O^d \quad (\text{D20})$$

$$S_{\text{uptake},L,C}^a = f_A(\psi_{m,\text{soil}}) \frac{k_{\text{max},C} C_L^l k_{C,L}}{k_{\text{max},C} + C_L^l k_{C,L}} \overline{C_C^a} \quad (\text{D21})$$

$$S_{\text{uptake},L,C}^d = f_A(\psi_{m,\text{soil}}) \frac{k_{\text{max},C} C_L^l k_{C,L}}{k_{\text{max},C} + C_L^l k_{C,L}} \beta_C C_C^d \quad (\text{D22})$$

The deactivation and reactivation rates are defined thus:

$$S_{deact,O} = \max((1 - \phi_O), f_{A2D}(\psi_{m,soil}))k_{d,O}\overline{C}_O^a \quad (D23)$$

$$1125 \quad S_{deact,C} = \max((1 - \phi_C), f_{A2D}(\psi_{m,soil}))k_{d,C}\overline{C}_C^a \quad (D24)$$

$$S_{react,O} = \min(\phi_O, f_{D2A}(\psi_{m,soil}))k_{r,O}C_O^d \quad (D25)$$

$$S_{react,C} = \min(\phi_C, f_{D2A}(\psi_{m,soil}))k_{r,C}C_C^d \quad (D26)$$

$$\phi_O = \frac{1}{e^{\frac{C_{thres,O}^l - C_L^l}{aC_{thres,O}^l}} + 1} \quad (D27)$$

$$\phi_C = \frac{1}{e^{\frac{C_{thres,C}^l - C_L^l}{aC_{thres,C}^l}} + 1} \quad (D28)$$

$$1130 \quad f_{A2D}(\psi_{m,soil}) = \frac{1}{1 + [\psi_{m,soil,A2D}/\psi_{m,soil}]^{\omega_{DA}}} \quad (D29)$$

$$f_{D2A}(\psi_{m,soil}) = \frac{1}{1 + [\psi_{m,soil}/(\tau \psi_{m,soil,A2D})]^{\omega_{DA}}} \quad (D30)$$

with $k_{d,Z}$ and $k_{r,Z}$ (1/d) respectively the deactivation and reactivation rate coefficients for C_Z . $C_{thres,Z}^l$ (mol C/cm³ water) is the C_L^l threshold for ϕ_Z . ϕ_Z (–) is the switch function defining the effect of C_L^l on microbial dormancy ($S_{deact,Z}$ and $S_{react,Z}$), while f_{A2D} and f_{D2A} (–) define the effect of soil water content (see Wang et al. (2021)). The increase of the deactivation (resp. decrease of the activation) is defined by the most limiting resource between dissolved low molecular weight organic C and water. For instance, when water limiting for microbial activation, we will have $\min(\phi_Z, f_{D2A}(\psi_{m,soil})) = f_{D2A}(\psi_{m,soil})$ or $\max((1 - \phi_Z), f_{A2D}(\psi_{m,soil})) = f_{A2D}(\psi_{m,soil})$. Finally, we get the depolymerisation rate (Pagel et al., 2020):

$$S_{depoly} = f_A(\psi_{m,soil})v_{max,depoly} \frac{C_H^l}{K_L + C_H^l} \overline{C}_O^a \quad (D31)$$

With $v_{max,depoly}$ (1/d) is the maximum reaction rate of enzymes targeting high molecular weight organic C compounds, K_L (mol/cm³ water) half-saturation coefficients of enzymes targeting high molecular weight organic C compounds.

From the reactions defined above, we also get C_{co2} (mol C/cm³ scv), the amount of C released:

$$\frac{\partial C_{co2}}{\partial t} = \frac{1 - Y_O}{Y_O} S_{growth,O} + \frac{1 - Y_C}{Y_C} S_{growth,C} + \frac{1 - Y_M}{Y_M} S_{Mdecay} + S_{Muptake,L} \quad (D32)$$

Although we do compute the amount of C released, we do not simulate its transport in the domain.

Appendix E: Growth of existing 1D axisymmetric soil models

1145 Once the 1D domain volumes at time step N have been computed (see Eq. (26)), the redistribution of the water and C between the [microscale-1D](#) domains of a voxel are done in 2 steps:

1. Compute the new water volume and solute content in the pre-existing [microscale-1D](#) domains that have shrunk while trying to maintain the radial concentration gradients.

2. Compute the water volume and C content of the new **microscale-1D** domains and of the pre-existing **microscale-domaines**
 1150 **1D domains** that have grown. We define for each voxel a mean water content and C concentration from the leftover water
 volume, C content, and voxel space. There is no concentration gradient in the newly added **microscale-space** **space of the**
1D models.

We set m_X (mol) as the content of the C element X with the concentration C_X , and V_w (cm^3 water) the water volume.

1155 E1 Shrinking 1D axisymmetric soil models

For perirhizal zone with a volume decrease between time step $N - 1$ and N , we adapt m_X in each soil voxel so that, for the
 segment k :

$$ChangeRatioS_k = \frac{m_{x,k}^N}{m_{X,k}^{N-1}}$$

$$ChangeRatioW_k = \frac{V_{w,k}^N}{V_{w,k}^{N-1}}$$

We set $ChangeRatioS$ and $ChangeRatioW$ so that the **microscale-domain-1D models** will lose or gain water and solute according
 to the volume change. $ChangeRatioW$ is also adapted to keep $\theta \in [\theta_{wilt\,point}, \theta_s]$. $\theta_{wilt\,point}$ is the soil water content when
 $\psi = \psi_{crit}$ the critical plant water potential. Consequently:

$$ChangeRatioS_k = \min\left(\frac{V_{1DS,scv,k}^N}{V_{1DS,scv,k}^{N-1}}, 1\right) \quad (E1)$$

$$1160 \quad ChangeRatioW_k = \max\left(\min\left(ChangeRatioS_k, \frac{\theta_s V_{1DS,scv,k}^N}{V_{w,k}^{N-1}}\right), \frac{\theta_{wilt\,point} V_{1DS,scv,k}^N}{V_{w,k}^{N-1}}\right) \quad (E2)$$

We also want to keep the radial concentration gradients:

$$\frac{\partial C_k^N}{\partial r} = \frac{\partial C_k^{N-1}}{\partial r} \quad (E3)$$

$$\frac{\partial \theta_k^N}{\partial r} = \frac{\partial \theta_k^{N-1}}{\partial r} \quad (E4)$$

For the discretised **microscale-1D** domain made of n segments, this gives us a set of equations that can be solved analytically.

1165 In matricial form:

$$B = Q^{-1}G \quad (E5)$$

With Q a matrix of size $[n, n]$:

$$Q = \begin{bmatrix} -1 & 1 & 0 & \dots & 0 \\ 0 & -1 & 1 & \dots & 0 \\ 0 & \dots & \dots & \dots & 0 \\ 0 & \dots & \dots & -1 & 1 \\ V_{0,k}^N & V_{1,k}^N & \dots & V_{n-1,k}^N & V_{n-1,k}^N \end{bmatrix} \quad (E6)$$

And G an array of size n :

$$1170 \quad G = \begin{bmatrix} b_{r_{1,k}^N}^{N-1} - b_{r_{0,k}^N}^{N-1} \\ \dots \\ \dots \\ b_{r_{n,k}^N}^{N-1} - b_{r_{n-1,k}^N}^{N-1} \\ B_k^{N-1} \text{ChangeRatio}_k^N \end{bmatrix} \quad (E7)$$

B_k^N is the total amount of the solute or of water (in mol or cm^3) and b is the solute concentration or water content. The last lines of G and Q assure that we obtain the correct total solute and water amount, the other lines represent the wished for gradient between the new location of the segment centers. The value of $b_{r_{1,k}^N}^{N-1} - b_{r_{0,k}^N}^{N-1}$ give the changes of b (at $N - 1$) between the segment centers (distance at N). It is obtained by interpolation of the concentration gradients between the segments centers

1175 at the end of time step $N - 1$.

$\text{ChangeRatio}W_k$ and $\text{ChangeRatio}S_k$ ensure that the overall θ and m_k in the whole microscale-1D domain remains within the needed range, but not for each segment. Consequently, when implementing the new water value for each microscale segment, we adapt the distribution of the water between the segments to have for each segment 1D sub-control volume $\theta \in [\theta_{wilt\text{ingpoint}}, \theta_s]$.

Likewise, we ensure that $\min(m_k) \geq 0$.

1180 This yields $C_{1DS}^{N,0,-1}$ and $\theta_{1DS}^{N,0,-1}$ to be used at the beginning of the next fixed point loop iteration, see section F.

E2 New and expanding 1D axisymmetric soil models

From the concentration of the existing microscale-1D domains per voxel, we compute the mean θ_{new} and C_{new} , the water and concentration in the newly freed soil space. θ_{new} and C_{new} will define the initial conditions of the added volume in the expanding or new microscale-1D domains. For this, we compute the total volume of the newly freed volume in the voxel

1185 ($V_{3DS,scv,lefter}^N, cm^3$):

$$V_{3DS,scv,i,lefter}^N = \sum_j^{n_{shrunk,i}^N} L_{j,N} \pi (r_{out,j}^N{}^2 - r_{in,j}^N{}^2) \quad (E8)$$

with $nr_{shrunk,i}^N$ the sets of [microscale-1D models](#) domains in voxel i that have shrunk at time step N . We then get the content of water and C that are not in the old [microscale domains-1D models](#) of voxel i anymore:

$$V_{3DS,w,i,leftover}^N = \sum_j^{nr_{shrunk,i}^N} V_{1DS,w,j}^{N-1} - V_{1DS,w,j}^N \quad (\text{E9})$$

$$1190 \quad m_{3DS,X,i,leftover}^N = \sum_j^{nr_{shrunk,i}^N} m_{1DS,X,j}^{N-1} - m_{1DS,X,j}^N \quad (\text{E10})$$

We can then get θ_{new} and C_{new} :

$$\theta_{k,new}^N = \begin{cases} \frac{V_{3DS,w,i,leftover}^N}{V_{3DS,scv,i,leftover}^N} & \text{if } \text{len}(nr_i^{N-1}) > 0 \\ \frac{V_{3DS,w,i}^{N-1}}{V_{3DS,scv,i}^{N-1}} & \text{else} \end{cases} \quad (\text{E11})$$

$$C_{k,new}^{y,N} = \begin{cases} \frac{X_{i,leftover}^N}{V_{3DS,i,leftover}^N} & \text{if } \text{len}(nr_i^{N-1}) > 0 \\ \frac{X_i^{N-1}}{V_{3DS,i}^{N-1}} & \text{else} \end{cases} \quad (\text{E12})$$

$$V_{3DS,i,leftover} = \begin{cases} V_{3DS,w,i,leftover} & \text{if } C^y = C^l \\ V_{3DS,scv,i,leftover} & \text{if } C^y = C^s \end{cases} \quad (\text{E13})$$

$$1195 \quad V_{3DS,i} = \begin{cases} V_{3DS,w,i} & \text{if } C^y = C^l \\ V_{3DS,scv,i} & \text{if } C^y = C^s \end{cases} \quad (\text{E14})$$

We can then add the C and water to the growing [microscale-1D soil](#) domains (the C concentration is null for newly created domains):

$$\theta_{1DS}^{N,0,-1} = \theta_{1DS}^{N-1} + \theta_{k,new}^N \Delta V_{1DS,scv,k} \quad (\text{E15})$$

$$C_{1DS}^{N,0,-1} = C_{1DS}^{N-1} + C_{k,new}^N \Delta V_{1DS,scv,k} \quad (\text{E16})$$

1200 We then use the method presented in section E1 to obtain the θ and C_X value for each [microscale segment-1D sub-control volume](#) respecting the new content and trying to maintain the old concentration gradients. This yields $C_{1DS}^{N,0,-1}$ and $\theta_{1DS}^{N,0,-1}$ to be used at the beginning of the next fixed point loop iteration, see section F.

Appendix F: Pseudo-code of the iterative computation loop

The three models (plant, 1D [microscale](#)-soil, 3D [macroscale](#)-soil) are coupled using an iterative approach based on the method
 1205 Jorda Guerra et al. (2021). In the following section, we present the implementation of the equations defined above, in matricial form and for one time step N . Although the water and C balances are computed together, for clarity, the descriptions of their respective computation loops are presented separately.

F1 Definition of matrices

For the section below, M represents a matrix of size $[ns, nr]$, with ns the number of soil voxels and nr the number of root segments belowground (equal to the number of [microscale-1D soil](#) domains). Other variables include vectors of size nr with root, root-soil interface or [microscale-1D soil](#) values (with [respectively-respectively](#) the subscripts $root, rsi, 1DS$). The root-soil interface corresponds to the 1D segment at the inner boundary of the [microscale-1D](#) models. Vectors with the subscript $3DS$ give the [macroscale-3D](#) soil values and are of size ns . $V_{V,w}$ (cm^3 water), θ , ψ , C_X and \mathbf{m}_x ($mol C$), K_{sri} ($cm^2 hPa^{-1} d^{-1}$) are vectors giving respectively the water volume, water content, water potential, component concentration, component content, and hydraulic conductivity at the soil-root interface. $V_{surf,out}$ (cm^2) is a vector of size nr and $V_{surf,out}[i] = 2\pi r_{out,i} L_{r,i}$, giving the surface at the outer boundary of the perirhizal zone i . Similarly, $V_{surf,in}[i] = 2\pi r_{in,i} L_{r,i}$ gives the surface at the inner boundary. $V_{V,scv,3DS}$ (resp. $V_{V,scv,1DS}$) is a vector of size ns (resp. nr) giving the volume of each voxel (resp. of each [microscale-1D soil](#) domain). The source vector is obtained from $\mathbf{S} = SV_V$ and is thus in cm^3 water d^{-1} or in $mol d^{-1}$. M_c is a binary matrix. M_c is filled with 0 except for $M_c[i, j] = 1$, when the j^{th} root segment (and corresponding 1D soil domain) is in the i^{th} soil voxel. $V_{hasRoot}$ is a vector of size ns with 1 for the [macroscale-3D](#) soil voxels with at least one root and 0 elsewhere. We set:

$$M_{surf,in} = M_c \circ V_{surf,in}$$

with \circ the hadamard product and $M_{surf,in}$ a matrix of size $[ns, nr]$ giving the inner surface of the [microscale-1D](#) domain
 1210 for each soil voxel. $Q_{w,3DS-3DS}$ (cm^3 water d^{-1}), and $Q_{X,3DS-3DS}$ ($mol C d^{-1}$) are vectors of size ns giving, for all the voxels containing at least one root, the changes in water or C (for a specific component) caused by exchanges with other voxels (i.e., not caused by input from a root or from biochemical reactions). $M_{weight,X}$ is the weight matrices used to divide voxel flows ($Q_{3DS-3DS}$) between the [microscale-1D soil](#) domains in each voxel (see Eq. (B8),(C9))

F2 Fixed point iteration for the water flow

1215 The pseudo-code below presents the implementation of the fixed-point iteration loop for the water flow.

$$\begin{aligned}
\text{(A)} \quad & L_{root}^N, M_c^N = \mathbf{Growth}(\mathbf{Env}^N, \mathbf{C}_L^{st,N-1}, \psi_x^{N-1}) \\
& M_{surf,in}^N, V_{surf,in}^N, V_{surf,out}^N, \theta_{1DS}^{N,0,-1}, \psi_{rsi}^{N,0,-1} = \\
& \quad \mathbf{Distribute}(L_{root}^N, M_c^N, \theta_{1DS}^{N-1}) \\
& Q_{w,3DS}^{N,0,-1} = Q_{w,3DS}^{N-1}, \theta_{3DS}^{N,0,-1} = \theta_{3DS}^{N-1}
\end{aligned} \tag{F1}$$

1220 for n in $\text{range}(\frac{\Delta t}{\delta t})$:

while $\text{any}(\text{err}\dots > \epsilon\dots)$, and $k < k_{max}$:

$$\text{(B)} \quad q_{w,root-soil}^{N,n,k} \psi_x^{N,n,k} = \mathbf{FlowPhoto}(\mathbf{Env}^{N,n}, \psi_{rsi}^{N,n,k-1}) \tag{F2}$$

$$\text{(C)} \quad S_{w,1DS-3DS}^{N,n,k} = -M_{weight,w}^{N,n,k,T} Q_{w,3DS}^{N,n,k-1} \tag{F3}$$

$$\begin{aligned}
& K_{sri}^{N,n,k}, \theta_{1DS}^{N,n,k}, q_{w,root-soil}^{N,n,k,limited} = \mathbf{Flow}(V_{surf,in}^N q_{w,root-soil}^{N,n,k} \delta t, \\
& \quad \mathbf{S}_{w,1DS-3DS}^{N,n,k} \delta t, \theta_{1DS}^{N,n,k-1}, V_{V,scv,1DS}^N)
\end{aligned} \tag{F4}$$

$$\mathbf{S}_{w,3DS}^{N,n,k} = M_{surf,in}^N q_{w,root-soil}^{N,n,k,limited} \tag{F5}$$

$$\text{(D)} \quad \theta_{3DS}^{N,n,k}, Q_{w,3DS}^{N,n,k} = \mathbf{Flow}(\mathbf{S}_{w,3DS}^{N,n,k} \delta t, \theta_{3DS}^{N,n,k-1}, V_{V,scv,3DS}^N) \tag{F6}$$

$$\begin{aligned}
\text{(E)} \quad & \text{errW}_{1DS,3DS,cumul}^{N,n,k} = \mathbf{Err}(M_c[\theta_{1DS}^{N,n,k} V_{V,scv,1DS}^N] - \\
& \quad [\theta_{3DS}^{N,n,k} V_{V,scv,3DS}])
\end{aligned} \tag{F7}$$

$$1230 \quad \text{errW}_{1DS,3DS}^{N,n,k} = \text{errW}_{1DS,3DS,cumul}^{N,n,k} - \text{errW}_{1DS,3DS,cumul}^{N,n-1} \tag{F8}$$

$$\text{errW}_{k,k-1} = \mathbf{Err}(X^{N,n,k} - X^{N,n,k-1}) \tag{F9}$$

$$\text{errW}_{qroot} = \mathbf{Err}(q_{w,root-soil}^{N,n,k,limited} - q_{w,root-soil}^{N,n,k}) \tag{F10}$$

$$\begin{aligned}
\text{errW}_{mass1DS} = \mathbf{Err}(\theta_{1DS}^{N,n,k} V_{V,scv,1DS}^N - [\theta_{1DS}^{N,n,IC} V_{V,scv,1DS}^N + \\
(V_{surf,in}^N q_{w,root-soil}^{N,n,k} + \mathbf{S}_{w,1DS-3DS}^{N,n,k} S_{w,1DS-1DS}^{N,n,k}) \Delta t])
\end{aligned} \tag{F11}$$

$$\begin{aligned}
1235 \quad \text{errW}_{mass3DS} = \mathbf{Err}(\sum_{segments} (\theta_{3DS}^{N,n,k} V_{V,scv,3DS} - [\theta_{3DS}^0 V_{V,scv,3DS} \\
+ \sum_j^N (M_{surf,in}^j q_{w,root-soil}^j \delta t^j)])
\end{aligned} \tag{F12}$$

$$\text{assert } \text{err}\dots \leq \epsilon\dots \tag{F13}$$

$$\begin{aligned}
\psi_{rsi}^{N,n+1,-1} &= \psi_{rsi}^{N,n,k}, \theta_{1DS}^{N,n+1,-1} = \theta_{1DS}^{N,n,k} \\
Q_{w,3DS}^{N,n+1,-1} &= Q_{w,3DS}^{N,n,k}, \theta_{3DS}^{N,0,-1} = \theta_{3DS}^{N-1}
\end{aligned} \tag{F14}$$

1240 N (resp. Δt) is the step index (resp. time step) of the whole computation loop and n (resp. δt) is the step index (resp. time step) of the fixed point iteration. k is the iteration number of the fixed point iteration and k_{max} the maximum amount of iteration allowed. \mathbf{Env} represents other environmental conditions, L_{root} (cm) is a vector which contains the length of each root segment (and thus microscale-1D domain).

The T superscript gives the transpose of a matrix. **Flow** represents a function computing the water flow and **FlowPhoto** the coupled plant water flow and photosynthesis. **Distribute** corresponds to the division of the voxel volume and water between the **microscale-1D** domains according to the methods presented in section 2.2.6. $err_{1DS,3DS,cumul}$ gives the difference between the water content values computed **at the micro- and macroscale by the 1D- and 3D-models** and $err_{1DS,3DS}$ the difference added at the last time step. err_{root} is the difference between the root water uptake rates computed by the plant and the **microscale-1D** soil. err_{mass} gives the mass balance error, $err_{k,k-1}$ the output differences between the last two iterations (non-convergence). **Err** is a function to sum and normalise the error. $err_{...}$ corresponds to each of the errors computed within the iteration loop, and $\epsilon_{...}$ is their respective maximum value allowed.

F3 Fixed point iteration for the solutes transport

The pseudo-code below presents the implementation of the fixed-point iteration loop for the C transport.

1255 (A)
$$L_{root}^N, M_c^N, q_{X,root-soil}^N, C_{X,root}^{st,N} = \mathbf{Growth} \circ \mathbf{TransportReac}(\text{Env}^N, \sum_j^n (A_n^{N-1,j} \delta t^{N,j}), C_{X,rsi}^{N-1}) \quad (\text{F15})$$

$$M_{surf,in}^N, V_{surf,in}^N, V_{surf,out}^N, C_{X,1DS}^{N,0,-1} = \mathbf{Distribute}(L_{root}^N, M_c^N, C_{X,1DS}^{N-1}) \quad (\text{F16})$$

$$Q_{X,3DS}^{N,0,-1} = Q_{X,3DS}^{N-1}, C_{X,3DS}^{N,0,-1} = C_{X,1DS}^{N-1} \quad (\text{F17})$$

for n in range($\frac{\Delta t}{\delta t}$):

1260 while any($\text{err}... > \epsilon...$), and $k < k_{max}$:

(B)
$$q_{X,root-soil}^{N,n,k}, A_n^{N,n,k} = \mathbf{TransportPhoto}(\text{Env}^{N,n}, \psi_{rsi}^{N,n,k-1}) \quad (\text{F18})$$

(C)
$$\mathbf{S}_{X,1DS-3DS}^{N,n,k} = -M_{weight,X}^{N,n,k,T} Q_{X,3DS}^{N,n,k-1} \quad (\text{F19})$$

$$C_{X,1DS}^{N,n,k}, \mathbf{m}_{X,1DS}^{N,n,k} = \mathbf{TransportReac}(V_{surf,in}^N q_{X,root-soil}^N \delta t, \mathbf{S}_{X,1DS-3DS}^{N,n,k} \delta t, C_{X,1DS}^{N,n,k}, \theta_{1DS}^{N,n,k}) \quad (\text{F20})$$

1265
$$\mathbf{S}_{X,3DS}^{N,n,k} = M_c^N ((\mathbf{m}_{X,1DS}^{N,n,k} - \mathbf{m}_{X,1DS}^{N,n-1}) / \delta t + \mathbf{S}_{X,1DS-3DS}^{N,n,k}) \quad (\text{F21})$$

(D)
$$C_{X,3DS}^{N,n,k}, Q_{X,3DS}^{N,n,k} = V_{hasRoot} \mathbf{Transport}(\mathbf{S}_{X,3DS}^{N,n,k} \delta t, \mathbf{C}_{X,3DS}^{N,n-1}) + (1 - V_{hasRoot}) \mathbf{TransportReac}(\mathbf{C}_{X,3DS}^{N,n-1}) \quad (\text{F22})$$

(E)
$$\text{err}C_{1DS,3DS,cumul} = \mathbf{Err}(M_c^N \mathbf{m}_{x1DS}^{N,n,k} - \mathbf{m}_{x3DS}^{N,n,k}) \quad (\text{F23})$$

$$\text{err}C_{1DS,3DS}^{N,n,k} = \text{err}C_{1DS,3DS,cumul}^{N,n,k} - \text{err}C_{1DS,3DS,cumul}^{N,n-1} \quad (\text{F24})$$

1270
$$\text{err}C_{k,k-1} = \mathbf{Err}(X^{N,n,k} - X^{N,n,k-1}) \quad (\text{F25})$$

$$\text{err}C_{mass1DS} = \mathbf{Err}(\sum_{comp}^{NComp} (\mathbf{m}_{X,1DS}^{N,k} - [\mathbf{m}_{X,1DS}^{N-1} + (V_{surf,in}^N q_{X,root-soil}^{N,k} + S_{x,1DS-1DS}^{N,k} + S_{x,1DS-3DS}^{N,k}) \Delta t])) \quad (\text{F26})$$

$$\text{err}C_{mass3DS} = \mathbf{Err}(\sum_{segments} (\sum_{comp}^{NComp} (\mathbf{m}_{X,3DS}^N - [\mathbf{m}_{X,3DS}^0 + \sum_n^N M_{surf,in}^n q_{X,root-soil}^n \Delta t]))) \quad (\text{F27})$$

1275 assert $\text{err}... \leq \epsilon...$ (F28)

$$\mathbf{m}_{X,1DS}^{N,n+1,-1} = \mathbf{m}_{X,1DS}^{N,n,k}, C_{X,1DS}^{N,n+1,-1} = C_{X,1DS}^{N,n,k} \quad (\text{F29})$$

$$Q_{X,3DS}^{N,n+1,-1} = Q_{X,3DS}^{N,n,k}, C_{X,3DS}^{N,n+1,-1} = C_{X,3DS}^{N,n,k} \quad (\text{F30})$$

TransportReac represents the implementation of a function computing biochemical reactions and solute transport according to input variables. We need to sum all the components for the solute mass balance (err_{mass}). A_n is the net assimilation of

1280 sucrose, thus $A_n \neq 0$ only for dissolved low molecular weight organic C (C_L^l and \mathbf{m}_L). $Q_{X,3DS} \neq 0$ only for the C of
 1285 dissolved low molecular weight organic compounds and high molecular weight organic compounds.

Appendix G: Soil parameters and process constraints

The tables below present the parameter and process constraints defined for [TraiRhizo-the soil model](#) by Sircan et al. (2025).

1285 Baseline soil corresponds to the soil outside the rhizosphere.

Table G1. Parameter constraints defined by Sircan et al. (2025). [These parameter constraints represent the relative differences between the characteristics of the two functional microbial groups represented \(oligotrophs and copiotrophs\).](#)

	Constraint	Description/ Explanation
1	$\mu_{max,C} > \mu_{max,O}$	Maximum growth rate of copiotrophs is higher than maximum growth rate of oligotrophs.
2	$k_{O,S} > k_{C,S}$	Specific substrate affinity to small molecules for oligotrophs is higher than for copiotrophs.
3	$k_{max,C} > k_{max,O}$	Maximum maintenance rate coef. must be higher for copiotrophs than for oligotrophs.
4	$C_{thresh,C} > C_{thresh,O}$	Oligotrophs (K-strategists) are adapted to low carbon and nutrient availability.
5	$k_{max,i} < \mu_{max,i}$ $i \in \{O, C\}$	It is a logical constraint to ensure organisms are "fit to survive".
6	$k_{r,i} \geq k_{d,i}$ $i \in \{O, C\}$	Transition to dormant (or potentially active) state is typically slower than reactivation.
7	$k_{j,i} > \mu_{max,i}$ $i \in \{O, C\}, j \in \{r, d\}$	Changes of metabolic state are faster than growth, death, and changes in composition
8	$Y_O > Y_C$	Oligotrophs are slow growing at high yield, copiotrophs are fast-growing at low yield

Table G2. Process constraints adapted from the ones defined by Sircan et al. (2025). ^a: The lower bound of the ~~constraints-constraint~~ was decreased by 10% compared with the value defined by Sircan et al. (2025). ^b: The ~~lower-upper~~ bound of the ~~constraints-constraint~~ was ~~decreased-increased~~ by 10% compared with the value defined by Sircan et al. (2025).

	Constraint	Description/ Explanation
1	$3.33e^{-6} < (C_O + C_C)_R < 1e^{-4} \text{ mol cm}^{-3}$	The range for the concentration of microbial biomass in the rhizosphere from literature for different soil types, textures, experimental conditions.
2 ^a	$0.9 < \left(\frac{C_O + C_C}{C_O + C_C}\right)_B < 2.5$	Microbial biomass is higher in the rhizosphere than baseline soil and lower than a threshold value which is set up from literature.
3 ^b	$3.33e^{-4} < (C_L^l + C_H^l)_R < 1.33e^{-3} \times 1.1 \text{ mol cm}^{-3}$	The range for soil organic carbon in the rhizosphere for different soil types, textures, experimental conditions.
4 ^b	$0.66 < \left(\frac{C_L^l + C_H^l}{C_L^l + C_H^l}\right)_B < 2 \times 1.1$	The range for ratio of soil organic carbon in the rhizosphere to baseline soil for different soil types, textures, experimental conditions.
5	$6e^{-4} < \left(\frac{C_O^a + C_C^a}{C_O + C_C}\right)_R < 0.6$	The range for ratio of active microbial biomass to total microbial biomass for different soil types, textures, experimental conditions.
6	$C_{L,R}^l < 4.58e^{-5} \text{ mol cm}^{-3}$	Based on water extractable organic carbon measurements. Water extractable organic carbon should include dissolved organic carbon (DOC) together with some sorbed carbon mass therefore it must be higher than the DOC concentration.
7 ^a	$0.9C_{L,B}^l < C_{L,R}^l$	Based on water extractable organic carbon measurements.
8 ^a	$0.9C_{C,R} > C_{O,R}$	
9	$\frac{C_{C,B}}{C_{O,B}} < \frac{C_{C,R}}{C_{O,R}}$	The rhizosphere is dominated by copiotrophs.

Appendix H: Lateral root water conductivity in the mature root zone

The mature root area as well as the root water conductivity ($k_{lat,x}$, $cm \text{ hPa}^{-1} d^{-1}$) values for the immature zone are taken from Giraud et al. (2023). However, in that study, $k_{lat,x} = 0$ in the mature root area (more than 0.8 cm away from the root

tip). To obtain more realistic results, we used the work of Bramley et al. (2007) to evaluate the $\frac{k_{lat,x,mature}}{k_{lat,x,immature}}$ ratio. Bramley et al. (2007, Fig. 4.15) defines the surface-normalised root resistance $k_{tot,x}$ in $m s^{-1} MPa 10^{-8}$ over the length l (m) from the tip as: $k_{tot,x}(l) = (-83 \times l + 20.9) \times 1e - 8$ and the mean root conductance (in $m^3 s^{-1} MPa 10^{-11}$) as $K_{tot,x}(l) = (0.7 \times l + 2.4) \times 1e - 11$. We can therefore obtain the side surface of the root segment (A in m) up to l cm from $A(l) = \frac{K_{tot,x}(l)}{k_{tot,x}(l)}$. Moreover, in their study, the root axial resistance was found to be negligible, and we get therefore $k_{lat,x}(l) = k_{tot,x}(l)$ and $K_{lat,x}(l) = K_{tot,x}(l)$. We assumed that the mean $k_{lat,x}$ over the assumed immature root zone could be defined as $k_{lat,x,immature} \approx k_{lat,x}(0.008)$. We then evaluated $K_{lat,x}$ between $l_1 = 0.07$ m from the tip and $l_2 = 0.18$ m from the tip, which were the range of distances used in their study. We then computed $\int_{l_1}^{l_2} A(l)dl$ and $\int_{l_1}^{l_2} K_{lat,x}(l)dl$. From this we could get $\int_{l_1}^{l_2} k_{lat,x}(l)dl = \frac{\int_{l_1}^{l_2} K_{lat,x}(l)dl}{\int_{l_1}^{l_2} A(l)dl}$. This yielded $\frac{k_{lat,x,mature}}{k_{lat,x,immature}} = \frac{\int_{l_1}^{l_2} k_{lat,x}(l)dl}{k_{lat,x}(0.008)} \times 100 = 1.45\% \approx 1.5\%$. We consequently set $k_{lat,x,mature} = 0.015k_{lat,x,immature}$ for our modelling study.

Appendix I: Calibration of the soil carbon dynamic

This section describes the method used to calibrate the soil C dynamic. The objective was to implement a simple method allowing us to select sets representing different dynamics. A summary of the characteristics of each of the three selected microbial parameter sets is given in table 2.

I1 Selection of the **FraiRhizo** microbial parameter sets

In their study, Sircan et al. (2025) defined 1650 parameter sets for **FraiRhizo** soil model according to a series of parameter and process constraints, which are given in appendix G. The 1650 parameter sets were divided in three groups according to the resulting gradients of C_L^l directly at the root surface—small, medium, or strong gradient. [a] We randomly selected 33 sets in each of the three gradient categories. Then, we ran the coupled model on the (adapted) parameterised plant of Giraud et al. (2023) between days 10 and 14 of growth (with dynamic soil water and C flows). [b] Sets that did not respect the process constraints defined by Sircan et al. (2025) were removed—see Table G2. [c] From the 12 remaining sets, we selected three sets representing different soil C dynamics. For step [a], we ran the simulation without water scarcity effect—meaning, without implementing the effect of water on microbial (de)activation (Eq. (D29), (D30)) and access to soil solutes (Eq. (D10)). For step [b], following the method of (Sircan et al., 2025) and the analysis of Kuzyakov and Razavi (2019), we defined the rhizosphere as the soil volume up to 3 mm from the root surface. The baseline soil (soil outside of the rhizosphere) values were taken from the **maeroseale-3D** voxel data at the bottom of the soil column. We retained the sets that respected all the constraints for all the rhizosphere units at the end of the simulation. In some cases, some perirhizal zone units were slightly outside the constraints' bounds. To retain those sets, some constraint bounds were enlarged by 10%. These bounds-relaxations are indicated in table G2.

Figure I1 gives the concentration of different C pools along the soil depth. The first set led to a high CO_2 production in spite of a low microbial development, and was named *highCO2*. The second set led to a high development of the microbial biomass and was named *highMB*. The last set led to low solute mineralisation and microbial development. It was named *lowMUuptake*.

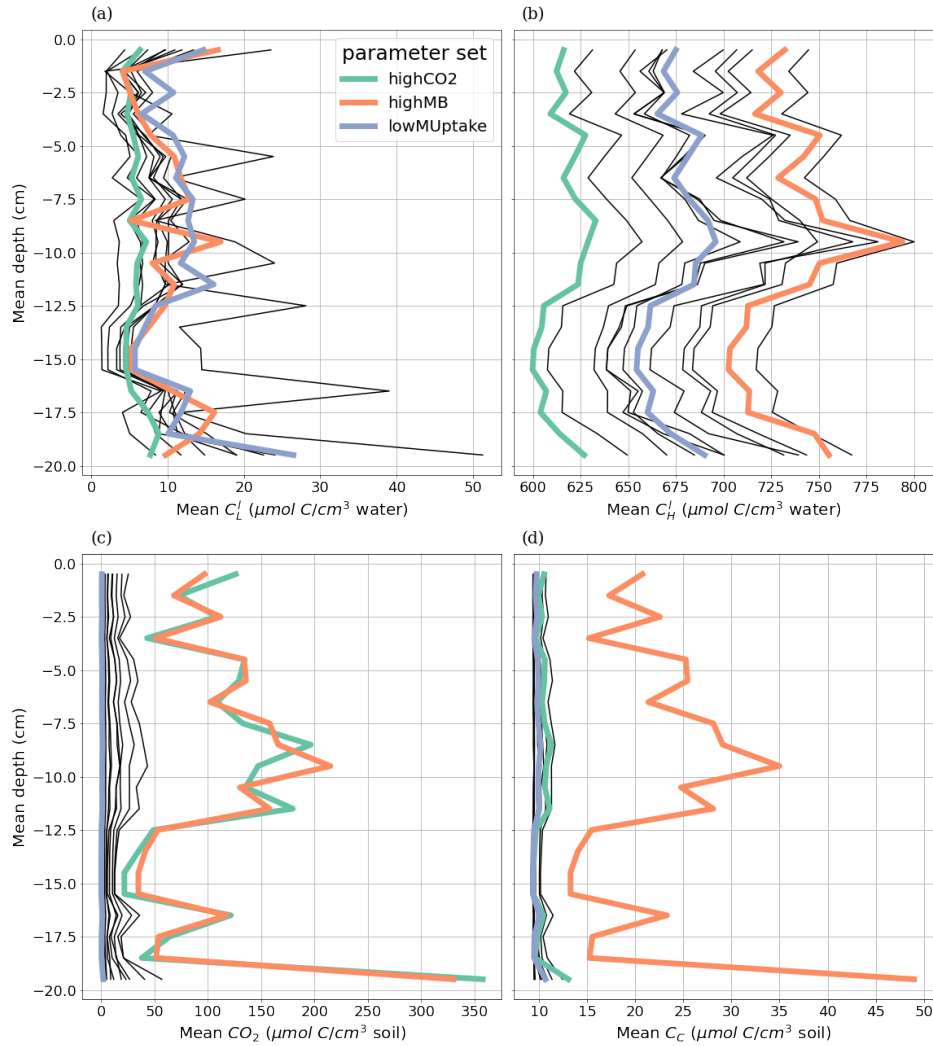


Figure I1. Mean carbon concentration per soil after four days of simulation (day 10 to 14) for concentrations of carbon from high (C_H^l) and dissolved low (C_L^l) molecular weight organic compounds, emitted CO_2 , and copiotroph carbon (C_C), in the 3D soil voxels which contain at least one root segment (perirhizal zone). The line colour gives the index of the parameter set. Only parameter sets that respect the process constraints defined by Sırcan et al. (2025) after four simulation days are represented. The selected sets are *highCO2* (green line), *highMB* (orange line) and *lowMUptake* (purple line).

I2 Description of the selected **TraiRhizo** microbial parameter sets

I2.1 Distribution of the parameter values

Figs. I2 and I3 present frequency histograms for each of the parameters in the 1650 sets. The parameters pointed by the numbered arrows correspond to the parameters of the three selected sets (*highCO2*, *highMB*, *lowMUptake*). As we can see,

1325 several of the parameter distributions are strongly skewed to the right, making some of the higher values outliers. Therefore, the sets containing the higher parameter values are more likely to give different outputs when compared with the other sets.

From the parameter values, we can give a description of the selected sets:

All three parameter sets offer low β_C (activity of dormant copiotrophs), $k_{C,S}$ (copiotroph affinity to dissolved low molecular weight organic C compounds), $k_{d,C}$, $k_{d,O}$ (deactivation rates), $k_{max,C}$, $k_{max,O}$ (maintenance rates), $\mu_{max,C}$, $\mu_{max,O}$ (growth rates), $v_{max,depoly}$ (depolymerisation rates of high molecular weight organic C compounds) values. This follows the average values of the 1650 sets and lead to a lower microbial activity and slower deactivation. Moreover, the sets have higher than average D_{LW} values, which should smooth the distribution of C_L^l . As can be seen more clearly in Fig. I3, we have an equilibrium between $k_{d,O}$ and $k_{r,O}$: the sets with higher $k_{d,O}$ also have higher $k_{r,O}$. All sets have a higher $k_{r,C}$ leading to a quick reactivation rate of the copiotrophs. Finally, the three sets have similar C_{init}^l values, and, in particular, a higher than average $C_{L,init}^l$.

1335 *highCO2* stands out by its higher β_O value, leading to a higher activity for dormant oligotrophs. Moreover, this sets has the higher D_{LW} values. Finally, the low growth yield (Y_C , Y_O) may lead to higher C usage and CO₂ emission rates.

highMB has higher activation rate parameters ($k_{r,O}$, $k_{r,C}$) linked with low C_{thres} parameters, which should lead to a high activated-to-total microbial biomass, although the deactivation parameter for oligotrophs ($k_{d,O}$) is also high. The relatively higher affinity parameters ($k_{C,S}$, $k_{O,S}$) and growth parameters ($\mu_{max,C}$, $\mu_{max,O}$) should lead to a quicker increase of the microbial biomass. The low Y (yield parameters) should also lead to higher C usage and CO₂ emissions.

1340 *lowMUptake* stands out for its low activation rate parameters ($k_{r,O}$, $k_{r,C}$) linked with high C_{thres} parameters, which should lead to a high activated-to-total microbial biomass, although the deactivation parameters ($k_{d,C}$, $k_{d,O}$) are also low. Low maintenance need for dormant microbes (β_C , β_{O}) and high yield parameters (Y , Y_C , Y_O) should also lead to a lower C usage and CO₂ emissions.

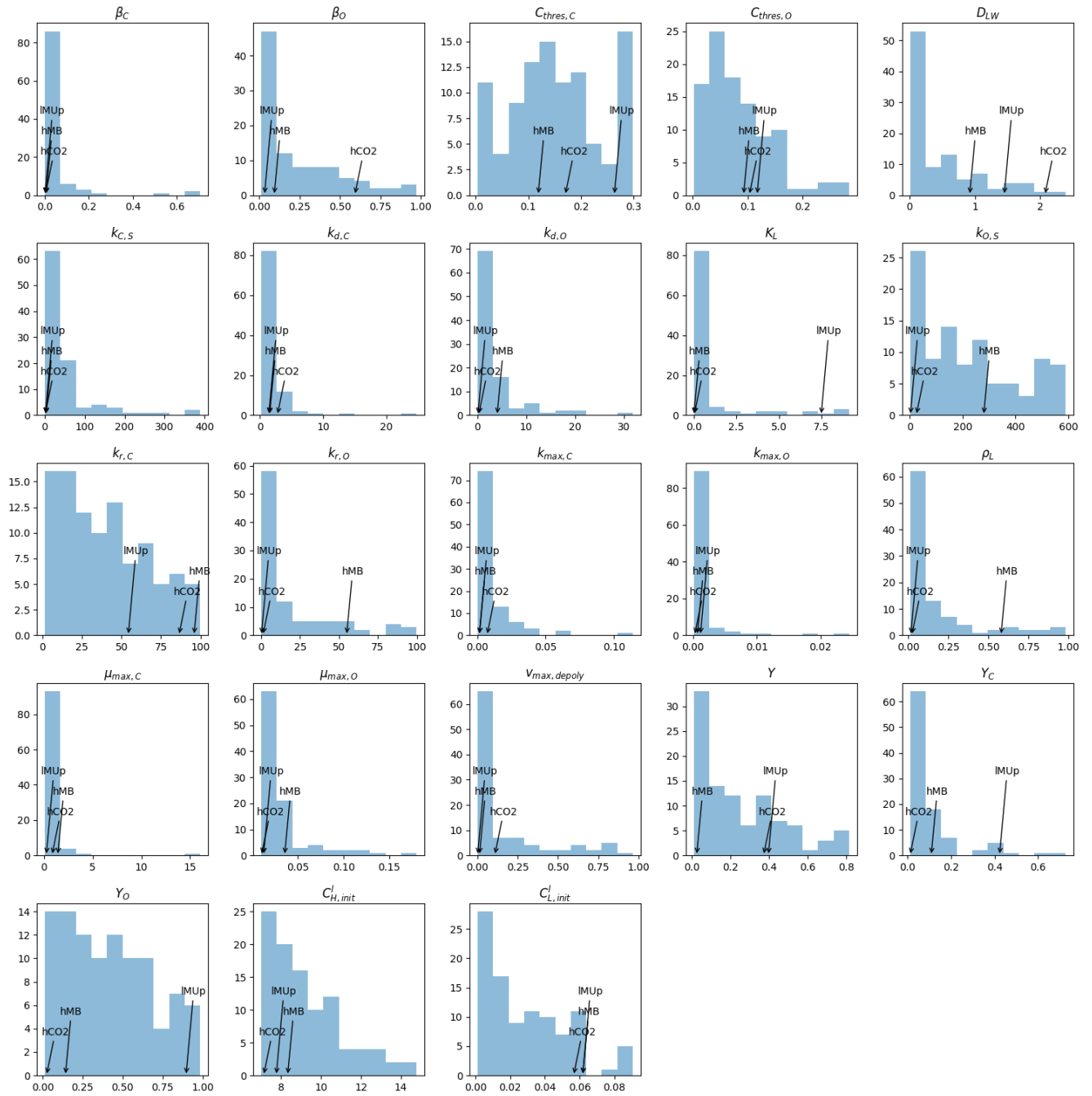


Figure 12. Histogram of the TraiRhizo input parameters Histogram of the soil input parameters. The parameters pointed by the numbered arrows correspond to the parameters of the three selected sets, *highCO2*, *highMB*, *lowMUptake*, respectively called *hCO2*, *hMB*, *IMUp*.

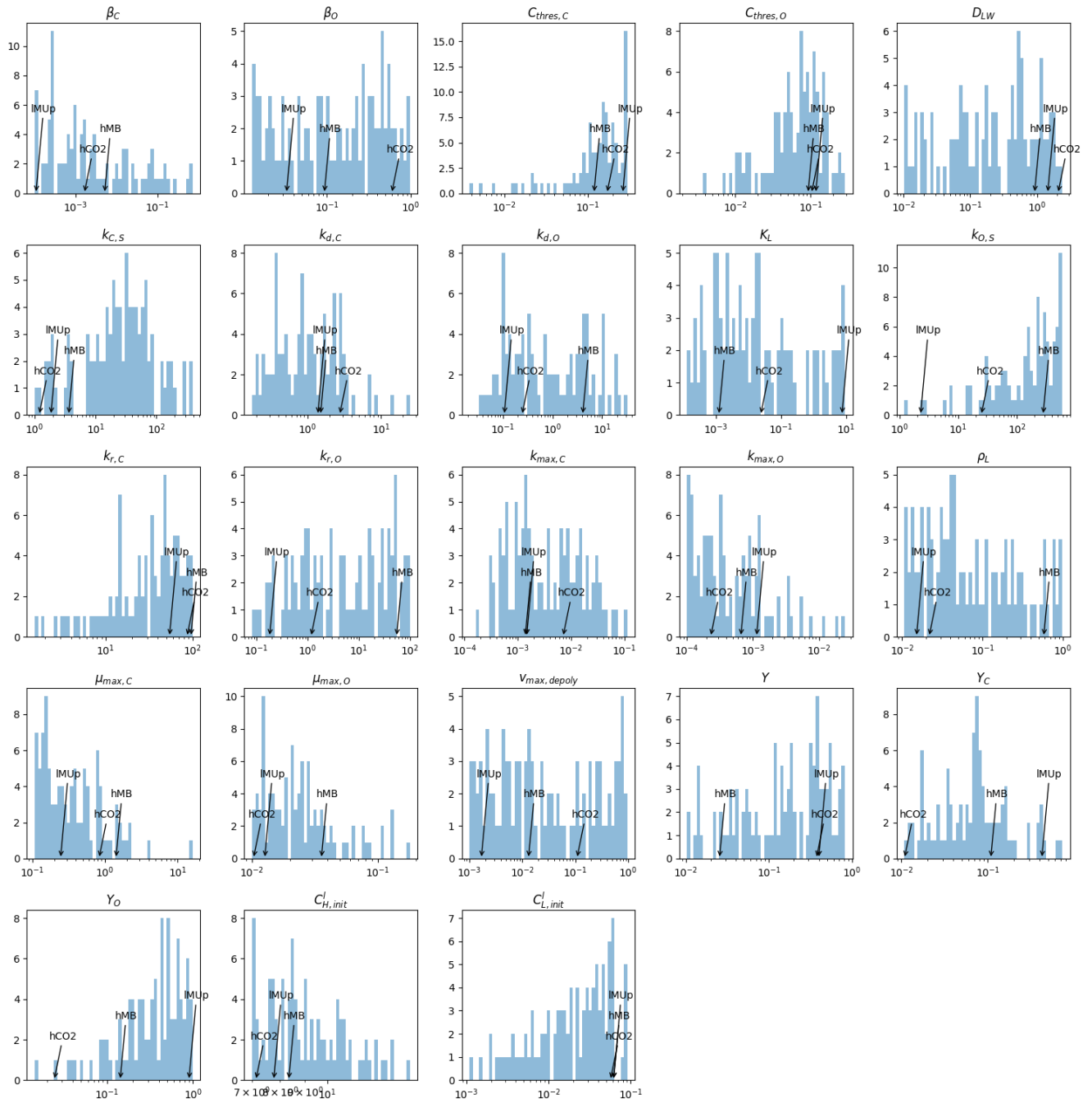


Figure 13. ~~Logarithmic-histogram of the TraiRhizo input parameters~~ Logarithmic histogram of the soil input parameters. he parameters pointed by the numbered arrows correspond to the parameters of the three selected sets, *highCO2*, *highMB*, *lowMUptake*, respectively called *hCO2*, *hMB*, *lMUptake*.

12.2 Distribution of output variables under different values of dissolved low molecular weight organic carbon

We computed for a range of C_L^l values intermediary outputs describing the microbial dynamic:

- ϕ_O (ϕ_C), -: ratio of oligotrophs (copiotrophs) which will be activated
- S_{decay} , $mol\ C\ cm^{-3}\ d^{-1}$: decay rate
- 1350 – S_{growth} , $mol\ C\ cm^{-3}\ d^{-1}$: growth rate
- S_{deact} , $mol\ C\ cm^{-3}\ d^{-1}$: deactivation rate of active microbes
- S_{react} , $mol\ C\ cm^{-3}\ d^{-1}$: reactivation rate of dormant microbes
- dCO_2dt , $mol\ C\ cm^{-3}\ d^{-1}$: production rate of CO₂
- $dMBdt$, $mol\ C\ cm^{-3}\ d^{-1}$: total absolute amount of change in the mass of microbial biomass C per unit of time

1355 Fig.I4 give the ratio between the related intermediary variables for all the parameter sets with the selected sets being colored. Fig. I5 shows the same data but only for the selected sets. Subgraphics d to i use a logarithmic scale on the y-axis. As we can observe, *highCO2* led to a higher relative advantage of the oligotroph compared with the copiotroph regarding their fitness (decay-to-growth rate ratio). This was driven by the higher copiotroph decay-to-growth rate ratio. This was partly compensated by the lower ratio $\frac{S_{growthC}+S_{decayC}}{S_{deactC}+S_{reactC}}$ indicating that the copiotroph dynamic was more driven by (de)activation than by growth and decay. Moreover, we can note a higher CO₂ emission rate per unit of transformed microbial C.

1360 *highMB* had lower ϕ values for both copiotrophs and oligotrophs, indicating that lower C_L^l values was necessary for an activation of the microbial community. This led in part to the lower $\frac{S_{deact}}{S_{react}}$ for both microbial group, causing a quicker reactivation compared with the deactivation. *highMB* also led to relatively low decay-to-growth ratio when compared with the other two sets. Finally, for lower C_L^l values, *highMB* had the highest relative importance of (de)activation on the copiotroph dynamic, when compared with the other two sets. *lowMUptake* offered the highest copiotroph fitness and the lowest oligotroph-to-copiotroph fitness ratio, giving a relative advantage to the copiotrophs. However, this set had the highest ϕ and $\frac{S_{deact}}{S_{react}}$ values, leading to a stronger deactivation of the microbes. For this set, we have a higher importance of the growth and decay over the (de)activation when comparing with the other two sets.

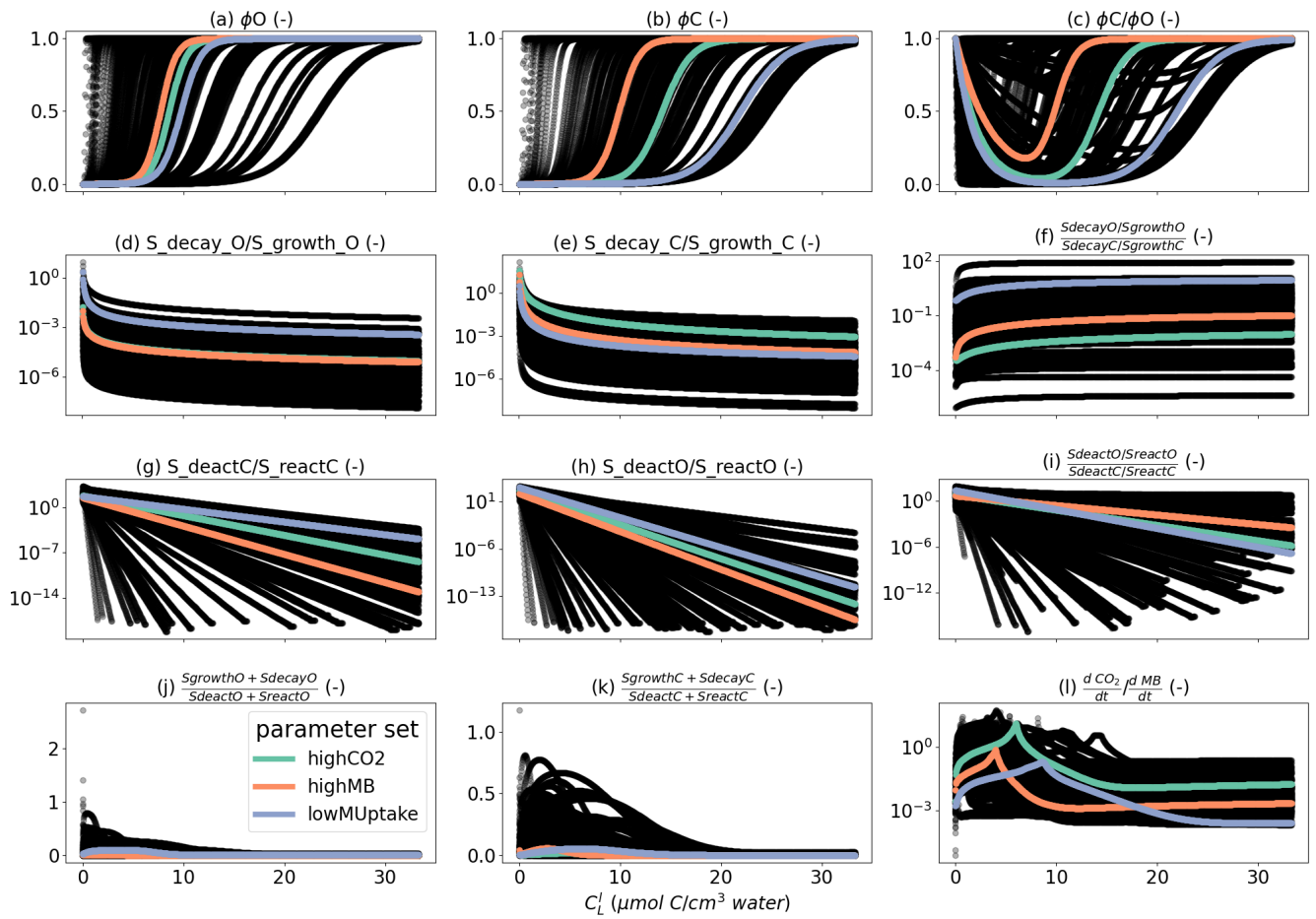


Figure I4. Distribution of specific microbial-related reaction rate for a range of C_L^l for all 99 parameter sets. The coloured values correspond to the three selected sets: *highCO2* (green line), *highMB* (orange line) and *lowMUptake* (purple line). Graphics d to i use a logarithmic scale for the y-axis.

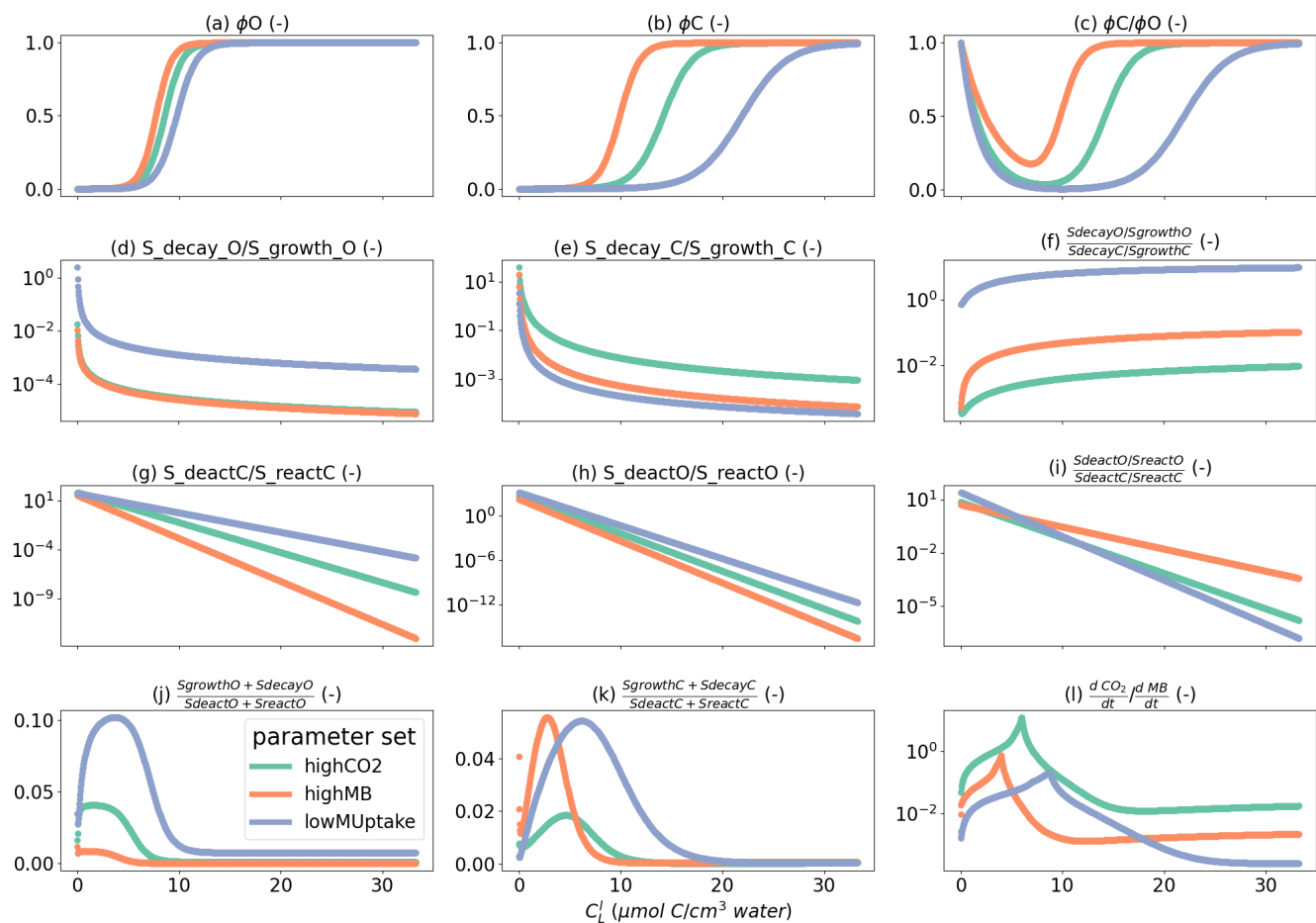


Figure 15. Distribution of specific microbial-related reaction rate for a range of C_L^l for the selected parameter sets. The coloured values correspond to the three selected sets: *highCO2* (green line), *highMB* (orange line) and *lowMUuptake* (purple line). Graphics d to i use a logarithmic scale for the y-axis.

Appendix J: Selected scales for the simulation outputs

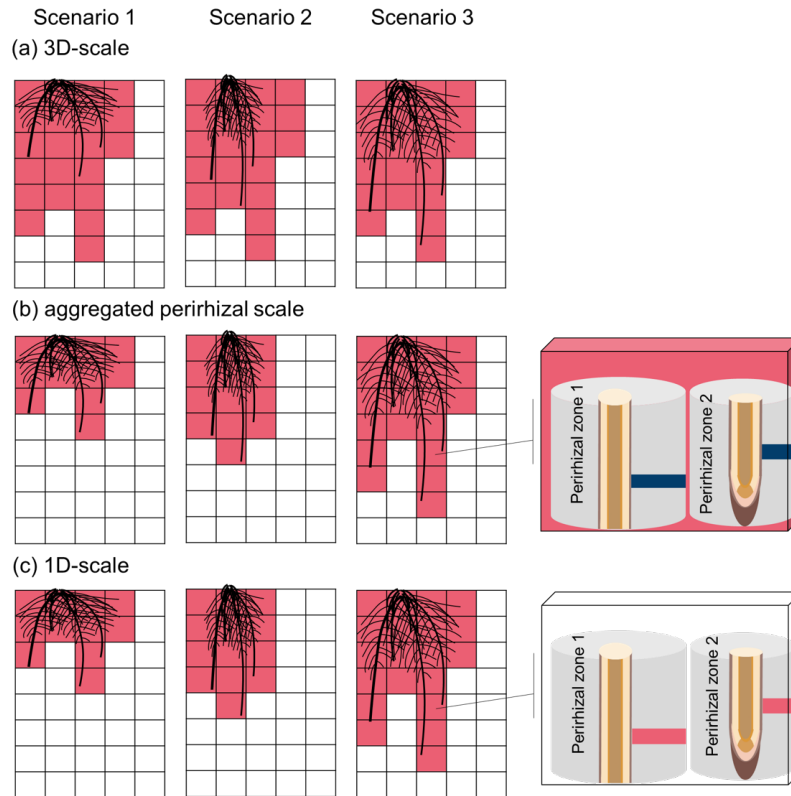


Figure J1. Graphical representation of the three scales used to analyse the simulation outputs in the main body of the text, using three example scenarios. (a) The 3D scale includes all the 3D voxels that contain at least one root at one time step of the simulations in any of the scenarios. This volume is constant over time. (b) The volume of the aggregated perirhizal scale can change at each time step and between scenarios. It includes the 3D voxels that contain at least one root for the studied time point of the studied scenario. (c) The 1D scale includes all the sub-control volume of the 1D soil model for the studied time point of the studied scenario. The 1D scale covers the same area as the aggregated perirhizal volume but represent a higher resolution: one data point for each sub-control volume of the 1D-models, against one data point for each sub-control volume of the selected 3D soil voxel. The definition of the $perirhizal_{trunc}$ presented in the appendix is not included in this graphic.

1370 Appendix K: Complementary results

K1 Model runtime

The model was implemented on the Institute of Bio- and Geosciences (IBG-3) computer cluster, Forschungszentrum Jülich. Each simulation was run on one node (shared memory) and 32 cores (Mainboard Supermicro H12DSU-iN), using all 32 cores for the simulation of the 1D and 3D soil domains (~~respectively the micro- and macroscale~~) and one core for the plant domain.

1375 The run-time of the simulation lasted between 9 h (*baseline* with *highSolutes*) and 13 h (*lateDry* with *highCO2*). The time step of the fixed point iteration remained at 20 mn for the baseline scenarios, went several times down to 7 mn for the *earlyDry* and *lateDry* scenarios. The computation of the water flows and content was the main bottleneck for the convergence of the iteration loop. For *baseline*, respectively 17%, 10%, and 27% of the runtime was spent solving the 3D soil domain, 1D soil domain, and plant models. For *earlyDry*, on average, 36%, 9%, and 14% of the runtime was spent solving the 3D soil domain, 1D soil domains, and plant models. For *lateDry*, on average, 38%, 11%, and 15% of the runtime is spent solving the 3D soil domain, 1D soil domain, and plant model. In total, between 54% and 64% of the runtime was dedicated to solving the models.

K2 Normalised root length density

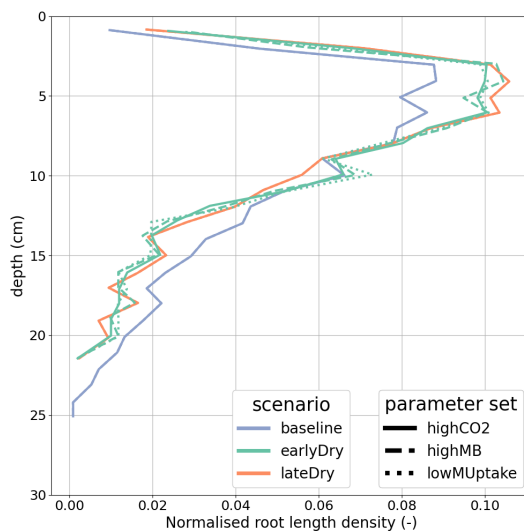


Figure K1. Normalised root length density at the end of the simulation. Line colours indicate the weather scenarios: baseline (blue) against an early (green) and late (red) dry spell. Line types represent different **biokinetic-kinetic** parameter sets.

K3 Plant processes

1385 Fig. 7.A presents the plant xylem mean water potential (ψ_x , cm) for each kinetic parameterisation and weather scenario. The simulation results reflect the diurnal dynamics of the xylem water potential in response to the truncated sinusoidal curve used to describe the atmospheric conditions (light input, air relative humidity and temperature). These boundary conditions affected the plant stomatal regulation and, thus, the C assimilation and transpiration rate. We observed an increase of the xylem water potential over time for *baseline* as the leaf:root ratio decreased. With drought, the xylem water potential rapidly decreased at the beginning of the dry spells and recovered after its end in response to changes in soil water content and mean air relative humidity during the dry spells (see Table 1). Towards the end of the simulation with *earlyDry*, the xylem water potential

1390

1395 converged to the values of *baseline*, indicating a full recovery of the plant water uptake capacity. As a consequence of the lower leaf:root ratio, the xylem water potential was lower at the beginning of the dry spells for *earlyDry* (11th day) than for *lateDry* (18th day). However, in comparison to *earlyDry*, the higher plant transpiration under *lateDry* with higher total leaf and root surface area led to a quicker decrease in soil water content. This mechanism induced lower xylem water potential in the *lateDry* compared to the *earlyDry* scenario from the third day after the start of the dry spells onward (13th and 20th day). Fig. 7.B shows the cumulative amount of C used for growth (and growth respiration) during the simulation. The diurnal cycle of the xylem water potential affected the plant growth. During the day, the transpiration-induced lower xylem water potential limited plant growth compared to the night, where increased xylem water potential permitted faster, albeit still water-limited, C assimilation and biomass production. The required C was taken from the starch reserves built up during the day. We can also observe trends over the simulation. Unlike younger plants, older plants could maintain growth during the diurnal period outside of the dry spells. During the dry spells, growth occurred only at night and at a severely reduced rate. On the last three nights of the late dry spell, we observed no night-growth due to lower plant night xylem water potential. Plant maintenance respiration increased during the spells due to the higher air temperatures. In the *earlyDry* scenario, plant growth was severely limited compared to *baseline*, resulting in a lower plant maintenance respiration rate after the dry spell ended (18th day). For *lateDry*, we saw a lower maintenance respiration rate from the fourth day of the spell. The total cumulative respiration (growth and maintenance related) at the end of the simulation was highest for the *baseline* scenario. It was about 29% lower for *earlyDry* and 18% lower for *lateDry*. The kinetic parameterisation had no significant effect on the plant water processes or growth and maintenance.

1410

Simulation results indicate that plant expansion growth is strongly influenced by changes in water potential during the diurnal cycle and drought periods, consistent with processes described in previous studies (Verbančič et al., 2018; Barillot et al., 2020; Coussement et al., 2018). However, it is hypothesised that cell wall biosynthesis (biosynthetic growth) predominantly occurs during the day using photoassimilates, while being much more limited at night when it relies on starch reserves (Verbančič et al., 2018). The expansion growth corresponds to the dilatation of those walls, driven by the cell's turgor pressure. Therefore, expansion growth depends on the supply of C but does not need to be concurrent to C supply. For this simulation, the virtual plant's starch pool, which varied according to the C supply, can be interpreted as representing both actual starch reserves and newly synthesised wall material. The carbon belonging to the newly synthesised wall material was moved to the carbon used for growth once these walls undergo expansion growth, thereby enabling the expansion growth to occur at night. To achieve robust results and better recreate typical C dynamics across a wider range of parameter sets, it could be beneficial to explicitly represent biosynthetic growth in future models. Following the analyses of Verbančič et al. (2018) and Barillot et al. (2020), high biosynthetic growth requires higher C supply but results in the creation of new cells more flexible and capable of expanding under lower turgor pressure. This higher amount of cells with high cell wall flexibility can help maintain plant growth in case of drought (Blum, 2017). Another mechanism supporting growth under low turgor pressure is osmotic adjustment, which has been identified as a critical process during drought conditions (Blum, 2017; Coussement et al., 2018). Not including this process in

1425

our model could explain the high growth limitation caused by drought observed in our simulation.

1430 In spite of these limitations, the outputs of these study follow more closely experimental results than those of Giraud et al. (2023)
: expansion growth occurring at night, starch supply and exudation being the regulatory variables for plant carbon balance.

Appendix L: Complementary cumulative volume distributions

1435 For the perirhizal_{trunc} analysis, we aggregated the mean concentrations for each 1D domain truncated at a specified distance from the root surface, denoted as $\Delta r_{analysis}$. The value of $\Delta r_{analysis}$ was pragmatically set to 0.06 cm. It corresponds to the minimum distance between the inner and outer boundaries of the 1D domains obtained during the simulation. By setting a consistent distance between the inner and outer boundaries of the truncated domains, we obtained a partial uniformity between the domains that reduced variations that could arise from differences in soil allocation to each domain (see Eq. (26)). Moreover, using these truncated 1D models allowed us to evaluate a soil area nearer to the root surface, for comparison with the aggregated perirhizal scale.

1440 We created complementary cumulative volume distribution plots of the concentration at the perirhizal_{trunc} scale. In these plots, the x-axis represents different concentration values. The y-axis shows the absolute or relative volume of the selected perirhizal zone (up to 0.06 cm from the root surface) with concentrations equal to or greater than the corresponding x-axis value. This representation provides insight into the spatial variability and distribution of concentrations within the specified zone. Additionally, these graphs allow us to evaluate how much of the perirhizal zone volume exceeds certain concentration levels, such as hotspot thresholds.

1445 Fig. L1 shows the perirhizal_{trunc} volume with a concentration inferior or equal to the value on the x-axis. The description and motivation of the evaluation method for this figure are given in section 2.4.3. A video presenting this graphic for each time step is available in the supplementary documents ("Video S1"). Root exudation and mucilage release triggered concentration peaks of solutes and microbial C in the perirhizal_{trunc} volume (Fig. L1). Particularly copiotrophs were strongly stimulated reaching high abundances up to $580 \mu\text{mol}/\text{cm}^3 \text{ soil}$ for *lateDry* with *highMB*, for a volume of $2.9e^{-2} \text{ cm}$. Both dry spells led to strongly reduced total perirhizal_{trunc} volumes at the end of the simulation of 42% for *earlyDry* and 34% for *lateDry* in relation to *baseline*. As a consequence, maximum concentrations of high molecular weight organic compounds-C and copiotrophs-C concentrations were lowest with *baseline* because the released plant C triggering copiotrophic growth was distributed in a larger volume.

1455 While *lateDry* with *highCO2* and *highMB* caused the highest CO₂, copiotrophs-C and dormant copiotrophs-C peaks at the aggregated perirhizal scale (see respectively Figs. 8, 9), for the perirhizal_{trunc} zone the highest values for these parameter sets were obtained with *earlyDry* (Fig. L1(c), (d), (f)). The simulated copiotroph growth during the dry spell occurred earlier under *earlyDry*, creating a larger pool of dormant copiotrophs-C ready to be re-activated and grow during periods of high exudation. As the growth rate is a function of the current active copiotrophs-C, high dormant copiotrophs-C early in the simulation favoured the overall growth in those hotspots.

1460 Parameterizing the model with *highMB* under *lateDry* caused an especially high copiotrophs-C over most of the perirhizal *trunc*
 volume with an increase of active and a strong build-up of dormant copiotrophs (Fig. L1(d), (e), (f)). Conversely, although
highMB under *lateDry* also led to a slight increase in the total and dormant oligotrophic concentrations (Fig. L1(g), (i)),
 it resulted in a decrease in the amount of active oligotrophs (Fig. L1(h)). This simulated pattern indicates a pronounced
 competition between oligotrophs and copiotrophs for this scenario-parameter set combination. Indeed, the active oligotrophs-C
 1465 values started decreasing strongly during the second half of the late dry spell, when the exudation rate became also lower.

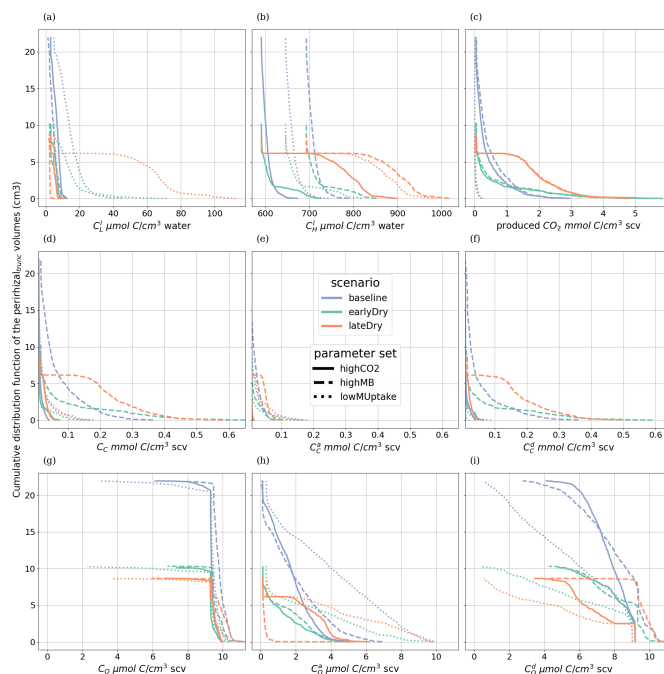


Figure L1. Perirhizal zone volume according to the minimal concentration of organic carbon in the truncated perirhizal zones, for (a) carbon of low-weight dissolved organic molecules (C_L), and (b) carbon of high-weight organic molecules (C_H), (c) emitted CO_2 , (d) total (C_C) (e) active (C_C^a) (f) dormant (C_C^d) copiotrophs in the soil phase, (g) total (C_O) (h) active (C_O^a) (i) dormant (C_O^d) oligotrophs in the soil phase. The line colors give the weather scenario: baseline (blue) against an early (green) and late (red) dry spell. The line types represent the different kinetic parameterisation. Subplots d, e, f and g, h, i have the same scale on the x-axis.

Appendix M: Radial carbon concentration profiles

To evaluate how C_L —the concentration of low-weight dissolved organic molecules-C is linked with the development and
 1470 activation of the most important microbial pool (copiotrophs), Fig. M1 presents the radial concentration profile of the C_C
 and $C_C^a:C_C$ copiotrophs-C and active-to-total copiotrophs-C ratio at the end of the simulations. We observed a very steep

1475 gradient of C_c -copiotrophs-C according to the distance from the root surface for *highMB* and *highCO2*. Indeed, the microbes developed around the C source. As they were not motile, they remained on this segment even once the root segment had become older and its C releases and water uptake decreased. For *lowMUptake*, the C_c -copiotrophs-C gradient (like the C_c -low-weight dissolved organic molecules-C concentration gradient, see Fig. 12) was smoother than the other two parameter sets for all weather scenarios. Therefore, even though C_c -copiotrophs-C directly at the root-soil interface is higher with *highCO2* than with *lowMUptake*, we observed similar C_c -at-the-bulk-soil-and-perirhizal-trunc-scales (Figs.??(e)and-L1-copiotrophs-C values at the aggregated perirhizal scales (Fig. 9).

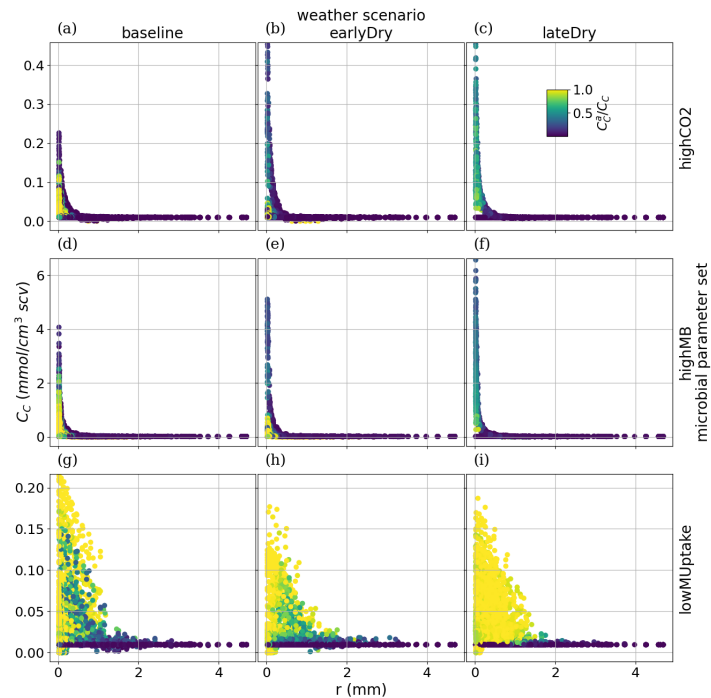


Figure M1. radial concentration profile of copiotroph carbon at the end of the simulation for each plant perirhizal zone. r (mm) corresponds to the distance to the root surface. The colour gives the ratio of active-to-total oligotroph biomass. Each column corresponds to a specific weather scenario and each row corresponds to a specific kinetic parameterisation. We have a different y-axis for each row.

M1 Effects of drought in the bulk soil

1480 To better understand the effect of drought on the perirhizal zone, Fig. M2 presents the relative change of carbon allocation caused by the droughts in the bulk soil. To represent the bulk soil, we selected the soil voxels at the bottom of the simulated soil column, which were separated by the nearest root by over 13cm. We can observe that the relative changes in carbon allocation was much more limited than within the perirhizal zone, and less affected by the day-night cycle. The lower soil water content

allowed for a higher concentration of soil solute, which led to a higher activation of the soil microbes (Fig. M2.(a)). However, the lower soil water potential also slowed the microbial activity, leading to a lower uptake of resources (Fig. M2.(d)), CO₂ emissions (Fig. M2.(c)), and microbial growth (Fig. M2.(b)). At this distance from the roots, the microbes are not affected by the changes in plant C releases, which lowers the differences between the *early dry* and *late dry* scenarios.

radial concentration profile of copiotroph carbon at the end of the simulation for each plant perirhizal zone. r (mm) corresponds to the distance to the root surface. The colour gives the ratio of active-to-total oligotroph biomass. Each column corresponds to a specific weather scenario and each row corresponds to a specific biokinetic parameterisation. We have a

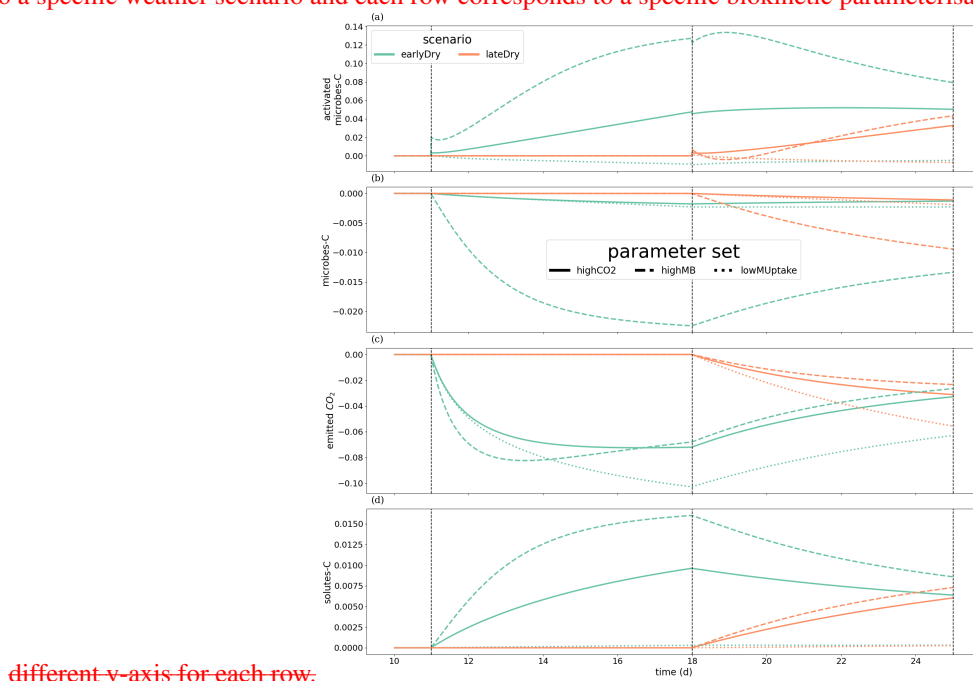


Figure M2. Relative change of carbon allocation in response to drought scenarios according to the kinetic parameters in the bulk soil.

The panels present the difference of C between each drought treatment and the baseline in relation to the baseline C usage for (a) microbes-C, (b) activated microbes-C, (c) CO₂ emitted by microbes, (d) soil solutes-C. The line types represent the different kinetic parameterisation. The dotted lines show the beginning and end of the early (day 11 to 18) and late (day 18 to 25) dry spells. Please note that the scales of the y-axis differ in each subplot.

. Author contribution

MG: Development and implementation of the new modules, Output visualization, Writing – original draft, Writing – review and editing.
 1490 AKS: Definition of the parameter space. MG, AKS: Setup of the equation set, Output analysis, Writing – review and editing. AS, GL, HP,

TS: Output analysis, Setup equation set, Writing – review and editing, Setup of the project and Funding acquisition, Supervision. GL, AS, HP: Support with the model development. MG and AKS contributed equally to the study. AS and HG supervised equally this study.

. Competing interests

The authors declare that they have no conflict of interest.

1495 **. Acknowledgement**

The model was implemented on the Institute of Bio- and Geosciences (IBG-3) computer cluster, Forschungszentrum Jülich. The authors acknowledge the DuMu^x developers for answering the DuMu^x-related questions, as well as Dr. Magdalena Landl for her help during this study.

Financial support

1500 The authors acknowledge funding by the German Federal Ministry of Education and Research (BMBF) in the framework of the funding measure "Plant roots and soil ecosystems, significance of the rhizosphere for the bio-economy" (Rhizo4Bio), subproject CROP phase I (FKZ 031B0909). Further funding is appreciated from the German Research Foundation (DFG) within the priority program 2322 "Soil Systems" (STR 481/12-1) and under Germany's Excellence Strategy (EXC 2070-390732324).

1505 **References**

- Ahrens, B., Guggenberger, G., Rethemeyer, J., John, S., Marschner, B., Heinze, S., Angst, G., Mueller, C. W., Kögel-Knabner, I., Leuschner, C., Hertel, D., Bachmann, J., Reichstein, M., and Schrumpf, M.: Combination of energy limitation and sorption capacity explains 14C depth gradients, *Soil Biology and Biochemistry*, 148, <https://doi.org/10.1016/j.soilbio.2020.107912>, 2020.
- Ahusborde, E., Kern, M., and Vostrikov, V.: Numerical simulation of two-phase multicomponent flow with reactive transport in porous media: application to geological sequestration of CO₂, *ESAIM: Proceedings and surveys*, 49, 21–39, <https://doi.org/10.1051/proc/201550002>, 2015.
- 1510
- Badri, D. V. and Vivanco, J. M.: Regulation and function of root exudates, *Plant, Cell & Environment*, 32, 666–681, <https://doi.org/10.1111/j.1365-3040.2009.01926.x>, 2009.
- Bardgett, R. D. and Caruso, T.: Soil microbial community responses to climate extremes: resistance, resilience and transitions to alternative states, *Philosophical Transactions of the Royal Society B: Biological Sciences*, 375, <https://doi.org/10.1098/rstb.2019.0112>, 2020.
- 1515
- Bardgett, R. D., Manning, P., Morriën, E., and De Vries, F. T.: Hierarchical responses of plant–soil interactions to climate change: consequences for the global carbon cycle, *Journal of Ecology*, 101, 334–343, <https://doi.org/10.1111/1365-2745.12043>, 2013.
- Barillot, R., De Swaef, T., Combes, D., Durand, J.-L., Escobar-Gutiérrez, A. J., Martre, P., Perrot, C., Roy, E., and Frak, E.: Leaf elongation response to blue light is mediated by stomatal-induced variations in transpiration in *Festuca arundinacea*, *Journal of Experimental Botany*, 72, 2642–2656, <https://doi.org/10.1093/jxb/eraa585>, 2020.
- 1520
- Bazot, S., Mikola, J., Nguyen, C., and Robin, C.: Defoliation-induced changes in carbon allocation and root soluble carbon concentration in field-grown *Lolium perenne* plants: do they affect carbon availability, microbes and animal trophic groups in soil?, *Functional Ecology*, 19, 886–896, <https://doi.org/10.1111/j.1365-2435.2005.01037.x>, 2005.
- Blum, A.: Osmotic adjustment is a prime drought stress adaptive engine in support of plant production, *Plant, Cell & Environment*, 40, 4–10, <https://doi.org/10.1111/pce.12800>, 2017.
- 1525
- Bonkowski, M., Tarkka, M., Razavi, B., Schmidt, H., Blagodatskaya, E., Koller, R., Yu, P., Knief, C., Hochholdinger, F., and Vetterlein, D.: Spatiotemporal Dynamics of Maize (*Zea mays* L.) Root Growth and Its Potential Consequences for the Assembly of the Rhizosphere Microbiota, *Frontiers in Microbiology*, 12, <https://doi.org/10.3389/fmicb.2021.619499>, 2021.
- Borken, W. and Matzner, E.: Reappraisal of drying and wetting effects on C and N mineralization and fluxes in soils, *Global Change Biology*, 15, 808–824, <https://doi.org/10.1111/j.1365-2486.2008.01681.x>, 2009.
- 1530
- Braghiere, R. K., Gérard, F., Evers, J. B., Pradal, C., and Pagès, L.: Simulating the effects of water limitation on plant biomass using a 3D functional-structural plant model of shoot and root driven by soil hydraulics, *Ann Bot*, 126, 713–728, <https://doi.org/10.1093/aob/mcaa059>, 2020.
- Bramley, H., Turner, D., Tyerman, S., and Turner, N.: Water Flow in the Roots of Crop Species: The Influence of Root Structure, Aquaporin Activity, and Waterlogging, in: *Advances in Agronomy*, vol. 96 of *Advances in Agronomy*, pp. 133–196, Academic Press, [https://doi.org/10.1016/S0065-2113\(07\)96002-2](https://doi.org/10.1016/S0065-2113(07)96002-2), 2007.
- 1535
- Brown, R. W., Chadwick, D. R., Bending, G. D., Collins, C. D., Whelton, H. L., Daulton, E., Covington, J. A., Bull, I. D., and Jones, D. L.: Nutrient (C, N and P) enrichment induces significant changes in the soil metabolite profile and microbial carbon partitioning, *Soil Biology and Biochemistry*, 172, <https://doi.org/10.1016/j.soilbio.2022.108779>, 2022.
- 1540
- Canarini, A., Kaiser, C., Merchant, A., Richter, A., and Wanek, W.: Root Exudation of Primary Metabolites: Mechanisms and Their Roles in Plant Responses to Environmental Stimuli, *Front Plant Sci*, 10, 157, <https://doi.org/10.3389/fpls.2019.00157>, 2019.

- Carminati, A., Kroener, E., Ahmed, M. A., Zarebanadkouki, M., Holz, M., and Ghezzehei, T.: Water for Carbon, Carbon for Water, *Vadose Zone Journal*, 15, vjz2015.04.0060, <https://doi.org/https://doi.org/10.2136/vjz2015.04.0060>, 2016.
- 1545 Coussement, J., Swaef, T., Lootens, P., Roldán-Ruiz, I., and Steppe, K.: Introducing turgor-driven growth dynamics into functional-structural plant models, *Annals of botany*, 121, <https://doi.org/10.1093/aob/mcx144>, 2018.
- Damour, G., Simonneau, T., Cochard, H., and Urban, L.: An overview of models of stomatal conductance at the leaf level, *Plant, Cell and Environment*, 33, 1419–1438, <https://doi.org/10.1111/j.1365-3040.2010.02181.x>, 2010.
- Darrah, P. R.: Models of the rhizosphere, *Plant and Soil*, 133, 187–199, <https://doi.org/10.1007/BF00009191>, 1991.
- 1550 de la Fuente Cantó, C., Simonin, M., King, E., Moulin, L., Bennett, M., Castrillo, G., and Laplaze, L.: An extended root phenotype: the rhizosphere, its formation and impacts on plant fitness, *The Plant Journal*, 103, <https://doi.org/10.1111/tpj.14781>, 2020.
- De Swaef, T., Pieters, O., Appeltans, S., Borra-Serrano, I., Coudron, W., Couvreur, V., Garré, S., Lootens, P., Nicolaï, B., Pols, L., Saint Cast, C., Šalagovič, J., Van Haeverbeke, M., Stock, M., and wyffels, F.: On the pivotal role of water potential to model plant physiological processes, in *silico Plants*, 4, <https://doi.org/10.1093/insilicoplants/diab038>, 2022.
- 1555 Debnath, L.: *Linear Partial Differential Equations*, pp. 1–148, Birkhäuser Boston, Boston, MA, ISBN 978-0-8176-4418-5, https://doi.org/10.1007/0-8176-4418-0_1, 2005.
- Deng, L., Peng, C., Kim, D.-G., Li, J., Liu, Y., Hai, X., Liu, Q., Huang, C., Shangguan, Z., and Kuzyakov, Y.: Drought effects on soil carbon and nitrogen dynamics in global natural ecosystems, *Earth-Science Reviews*, 214, <https://doi.org/10.1016/j.earscirev.2020.103501>, 2021.
- Dilkes, N. B., Jones, D. L., and Farrar, J.: Temporal Dynamics of Carbon Partitioning and Rhizodeposition in Wheat, *Plant Physiology*, 134, 706–715, <https://doi.org/10.1104/pp.103.032045>, 2004.
- 1560 Drake, J. E., Darby, B. A., Giasson, M.-A., Kramer, M. A., Phillips, R. P., and Finzi, A. C.: Stoichiometry constrains microbial response to root exudation- insights from a model and a field experiment in a temperate forest, *Biogeosciences*, 10, 821–838, <https://doi.org/10.5194/bg-10-821-2013>, 2013.
- Dupuy, L. X. and Silk, W. K.: Mechanisms of Early Microbial Establishment on Growing Root Surfaces, *Vadose Zone Journal*, 15, <https://doi.org/10.2136/vjz2015.06.0094>, 2016.
- 1565 Farquhar, G. D., von Caemmerer, S., and Berry, J. A.: A biochemical model of photosynthetic CO₂ assimilation in leaves of C₃ species, *Planta*, 149, 78–90, <https://doi.org/10.1007/BF00386231>, 1980.
- Galindo-Castañeda, T., Lynch, J., Six, J., and Hartmann, M.: Improving Soil Resource Uptake by Plants Through Capitalizing on Synergies Between Root Architecture and Anatomy and Root-Associated Microorganisms, *Frontiers in Plant Science*, 13, <https://doi.org/10.3389/fpls.2022.827369>, 2022.
- 1570 Galindo-Castañeda, T., Hartmann, M., and Lynch, J. P.: Location: root architecture structures rhizosphere microbial associations, *Journal of Experimental Botany*, 75, 594–604, <https://doi.org/10.1093/jxb/erad421>, 2023.
- Gaudio, N., Louarn, G., Barillot, R., Meunier, C., Vezy, R., and Launay, M.: Exploring complementarities between modelling approaches that enable upscaling from plant community functioning to ecosystem services as a way to support agroecological transition, in *silico Plants*, 4, <https://doi.org/10.1093/insilicoplants/diab037>, 2021.
- 1575 George, T. S., Bulgarelli, D., Carminati, A., Chen, Y., Jones, D., Kuzyakov, Y., Schnepf, A., Wissuwa, M., and Roose, T.: Bottom-up perspective – The role of roots and rhizosphere in climate change adaptation and mitigation in agroecosystems, *Plant and Soil*, <https://doi.org/10.1007/s11104-024-06626-6>, 2024.

- Giraud, M., Le Gall, S., Harings, M., Javaux, M., Leitner, D., Meunier, F., Rothfuss, Y., van Dusschoten, D., Vanderborght, J., Vereecken, H., Lobet, G., and Schnepf, A.: CPlantBox: a fully coupled modeling platform for the water and carbon fluxes in the Soil-Plant-Atmosphere-Continuum, in *silico Plants*, <https://doi.org/10.1093/insilicoplants/diad009>, 2023.
- 1580 Gérard, F., Blitz-Frayret, C., Hinsinger, P., and Pagès, L.: Modelling the interactions between root system architecture, root functions and reactive transport processes in soil, *Plant and Soil*, 413, 161–180, <https://doi.org/10.1007/s11104-016-3092-x>, 2017.
- Hartmann, H., Bahn, M., Carbone, M., and Richardson, A. D.: Plant carbon allocation in a changing world – challenges and progress: introduction to a Virtual Issue on carbon allocation, *New Phytologist*, 227, 981–988, <https://doi.org/10.1111/nph.16757>, 2020.
- 1585 Helmig, R.: *Multiphase Flow and Transport Processes in the Subsurface: A Contribution to the Modeling of Hydrosystems*, Springer, <https://doi.org/http://dx.doi.org/10.1007/978-3-642-60763-9>, 1997.
- Heppell, J., Payvandi, S., Talboys, P., Zygalkis, K. C., Langton, D., Sylvester-Bradley, R., Edwards, A. C., Walker, R., Withers, P., Jones, D. L., and Roose, T.: Use of a coupled soil-root-leaf model to optimise phosphate fertiliser use efficiency in barley, *Plant and Soil*, 406, 341–357, <https://doi.org/10.1007/s11104-016-2883-4>, 2016.
- 1590 Hirschberg, K., Miller, C. M., Ellenberg, J., Presley, J. F., Siggia, E. D., Phair, R. D., and Lippincott-Schwartz, J.: Kinetic analysis of secretory protein traffic and characterization of golgi to plasma membrane transport intermediates in living cells, *J Cell Biol*, 143, <https://doi.org/10.1083/jcb.143.6.1485>, 1998.
- Hobbie, J. E. and Hobbie, E. A.: Microbes in nature are limited by carbon and energy: the starving-survival lifestyle in soil and consequences for estimating microbial rates, *Frontiers in Microbiology*, 4, <https://doi.org/10.3389/fmicb.2013.00324>, 2013.
- 1595 Jiang, C., Séquaris, J.-M., Wacha, A., Bóta, A., Vereecken, H., and Klumpp, E.: Effect of metal oxide on surface area and pore size of water-dispersible colloids from three German silt loam topsoils, *Geoderma*, 235-236, 260–270, <https://doi.org/10.1016/j.geoderma.2014.07.017>, 2014.
- Jorda Guerra, H., Huber, K., Kunkel, A., Vanderborght, J., Javaux, M., Oberdörster, C., Hammel, K., and Schnepf, A.: Mechanistic modeling of pesticide uptake with a 3D plant architecture model, *Environmental Science and Pollution Research*, 28, <https://doi.org/10.1007/s11356-021-14878-3>, 2021.
- 1600 Khare, D., Selzner, T., Leitner, D., Vanderborght, J., Vereecken, H., and Schnepf, A.: Root System Scale Models Significantly Overestimate Root Water Uptake at Drying Soil Conditions, *Frontiers in Plant Science*, 13, <https://doi.org/10.3389/fpls.2022.798741>, 2022.
- Koch, T., Glaser, D., Weishaupt, K., Ackermann, S., Beck, M., Becker, B., Burbulla, S., Class, H., Coltman, E., Emmert, S., Fetzer, T., Grüninger, C., Heck, K., Hommel, J., Kurz, T., Lipp, M., Mohammadi, F., Scherrer, S., Schneider, M., Seitz, G., Stadler, L., Utz, M., Weinhardt, F., and Flemisch, B.: DuMux 3 – an open-source simulator for solving flow and transport problems in porous media with a focus on model coupling, *Computers & Mathematics with Applications*, 81, 423–443, <https://doi.org/10.1016/j.camwa.2020.02.012>, 2021.
- 1605 Kravchenko, L. V., Strigul, N. S., and Shvytov, I. A.: Mathematical Simulation of the Dynamics of Interacting Populations of Rhizosphere Microorganisms, *Microbiology*, 73, 189–195, <https://doi.org/10.1023/B:MICI.0000023988.11064.43>, 2004.
- 1610 Kuppe, C. W., Schnepf, A., von Lieres, E., Watt, M., and Postma, J. A.: Rhizosphere models: their concepts and application to plant-soil ecosystems, *Plant and Soil*, 474, 17–55, <https://doi.org/10.1007/s11104-021-05201-7>, 2022.
- Kutschera-Mitter, L., Barmicheva, K. M., and Sobotik, M.: *The Importance of Root-Cap Mucilage for Plant And Soil*, pp. 673–683, Springer Netherlands, Dordrecht, ISBN 978-94-011-5270-9, https://doi.org/10.1007/978-94-011-5270-9_60, 1998.
- Kuzyakov, Y. and Cheng, W.: Photosynthesis controls of CO₂ efflux from maize rhizosphere, *Plant and Soil*, 263, 85–99, 1615 <https://doi.org/10.1023/B:PLSO.0000047728.61591.fd>, 2004.

- Kuzyakov, Y. and Razavi, B. S.: Rhizosphere size and shape: Temporal dynamics and spatial stationarity, *Soil Biology and Biochemistry*, 135, 343–360, <https://doi.org/10.1016/j.soilbio.2019.05.011>, 2019.
- Lacointe, A. and Minchin, P. E. H.: A Mechanistic Model to Predict Distribution of Carbon Among Multiple Sinks, pp. 371–386, Springer New York, New York, NY, https://doi.org/10.1007/978-1-4939-9562-2_28, 2019.
- 1620 Landl, M., Haupenthal, A., Leitner, D., Kroener, E., Vetterlein, D., Bol, R., Vereecken, H., Vanderborght, J., and Schnepf, A.: Simulating rhizodeposition patterns around growing and exuding root systems, *in silico Plants*, 3, <https://doi.org/10.1093/insilicoplants/diab028>, 2021a.
- Landl, M., Phalempin, M., Schlüter, S., Vetterlein, D., Vanderborght, J., Kroener, E., and Schnepf, A.: Modeling the Impact of Rhizosphere Bulk Density and Mucilage Gradients on Root Water Uptake, *Frontiers in Plant Science*, 3, <https://doi.org/10.3389/fagro.2021.622367>, 2021b.
- 1625 Leuning, R.: A critical appraisal of a combined stomatal-photosynthesis model for C₃ plants, *Plant, Cell and Environment*, 18, 339–355, <https://doi.org/10.1111/j.1365-3040.1995.tb00370.x>, 1995.
- Lynch, J.: Harnessing root architecture to address global challenges, *The Plant Journal*, 109, <https://doi.org/10.1111/tpj.15560>, 2021.
- Lynch, J. M. and Whipps, J. M.: Substrate flow in the rhizosphere, *Plant and Soil*, 129, 1–10, <https://doi.org/10.1007/BF00011685>, 1990.
- 1630 Lynch, J. P., Strock, C. F., Schneider, H. M., Sidhu, J. S., Ajmera, I., Galindo-Castañeda, T., Klein, S. P., and Hanlon, M. T.: Root anatomy and soil resource capture, *Plant and Soil*, 466, 21–63, <https://doi.org/10.1007/s11104-021-05010-y>, 2021.
- Ma, W., Tang, S., Dengzeng, Z., Zhang, D., Zhang, T., and Ma, X.: Root exudates contribute to belowground ecosystem hotspots: A review, *Frontiers in Microbiology*, 13, <https://doi.org/10.3389/fmicb.2022.937940>, 2022.
- Mai, T. H., Schnepf, A., Vereecken, H., and Vanderborght, J.: Continuum multiscale model of root water and nutrient uptake from soil with explicit consideration of the 3D root architecture and the rhizosphere gradients, *Plant and Soil*, 439, 273–292, <https://doi.org/10.1007/s11104-018-3890-4>, 2019.
- 1635 McDougall, B. M. and Rovira, A. D.: Sites of exudation of ¹⁴C-labelled compounds from wheat roots, *New Phytologist*, 69, 999–1003, <https://doi.org/10.1111/j.1469-8137.1970.tb02479.x>, 1970.
- Meunier, F., Draye, X., Vanderborght, J., Javaux, M., and Couvreur, V.: A hybrid analytical-numerical method for solving water flow equations in root hydraulic architectures, *Applied Mathematical Modelling*, 52, 648–663, <https://doi.org/10.1016/j.apm.2017.08.011>, 2017.
- 1640 Millington, R. J. and Quirk, J. P.: Permeability of porous solids, *Transactions of the Faraday Society*, 57, 1200–1207, <https://doi.org/10.1039/TF9615701200>, 1961.
- Morandage, S., Laloy, E., Schnepf, A., Vereecken, H., and Vanderborght, J.: Bayesian inference of root architectural model parameters from synthetic field data, *Plant and Soil*, 467, 1–23, <https://doi.org/10.1007/s11104-021-05026-4>, 2021.
- 1645 Morandage, S., Schnepf, A., Leitner, D., Javaux, M., Vereecken, H., and Vanderborght, J.: Correction to: Parameter sensitivity analysis of a root system architecture model based on virtual field sampling, *Plant and Soil*, 477, 849–850, <https://doi.org/10.1007/s11104-021-05089-3>, 2022.
- Moyano, F. E., Manzoni, S., and Chenu, C.: Responses of soil heterotrophic respiration to moisture availability: An exploration of processes and models, *Soil Biology and Biochemistry*, 59, 72–85, <https://doi.org/10.1016/j.soilbio.2013.01.002>, 2013.
- 1650 Mualem, Y.: A new model for predicting the hydraulic conductivity of unsaturated porous media, *Water Resources Research*, 12, 513–522, <https://doi.org/10.1029/WR012i003p00513>, 1976.
- Neumann, G. and Römheld, V.: The release of root exudates as affected by the plant's physiological status, pp. 23–54, CRC Press, 2009.

- Nguyen, C.: Rhizodeposition of Organic C by Plant: Mechanisms and Controls, vol. 23, pp. 97–123, Springer Netherlands, ISBN 978-90-1655 481-2665-1, https://doi.org/10.1007/978-90-481-2666-8_9, 2009.
- Pagel, H., Kriesche, B., Uksa, M., Poll, C., Kandeler, E., Schmidt, V., and Streck, T.: Spatial Control of Carbon Dynamics in Soil by Microbial Decomposer Communities, *Frontiers in Environmental Science*, 8, <https://doi.org/10.3389/fenvs.2020.00002>, 2020.
- Personeni, E., Nguyen, C., Marchal, P., and Pagès, L.: Experimental evaluation of an efflux–influx model of C exudation by individual apical root segments, *Journal of Experimental Botany*, 58, 2091–2099, <https://doi.org/10.1093/jxb/erm065>, 2007.
- 1660 Poelplau, C. and Don, A.: A simple soil organic carbon level metric beyond the organic carbon to clay ratio, *Soil Use and Management*, 39, <https://doi.org/10.1111/sum.12921>, 2023.
- Pot, V., Portell, X., Otten, W., Garnier, P., Monga, O., and Baveye, P. C.: Accounting for soil architecture and microbial dynamics in microscale models: Current practices in soil science and the path ahead, *European Journal of Soil Science*, 73, <https://doi.org/10.1111/ejss.13142>, 2022.
- 1665 Prescott, C. E., Grayston, S. J., Helmisaari, H.-S., Kaštovská, E., Körner, C., Lambers, H., Meier, I. C., Millard, P., and Ostonen, I.: Surplus Carbon Drives Allocation and Plant–Soil Interactions, *Trends in Ecology & Evolution*, 35, 1110–1118, <https://doi.org/10.1016/j.tree.2020.08.007>, 2020.
- Putten, W., Bardgett, R., Bever, J., Bezemer, T., Casper, B., Fukami, T., Kardol, P., Klironomos, J., Kulmatiski, A., Schweitzer, J., Suding, K., Voorde, T., and Wardle, D.: Plant-Soil Feedbacks: the past, the present and future challenges., *Journal of Ecology*, 101, 265–276, <https://doi.org/10.1111/1365-2745.12054>, 2013.
- 1670 Přikryl, Z. and Vančura, V.: Root exudates of plants, *Plant and Soil*, 57, 69–83, <https://doi.org/10.1007/BF02139643>, 1980.
- Rakshit, A., Singh, S., Abhilash, P. C., and Biswas, A., eds.: *Soil Science: Fundamentals to Recent Advances*, Springer, Singapore, <https://doi.org/10.1007/978-981-16-0917-6>, 2021.
- Rees, F., Richard-Molard, C., Chenu, C., and Andrieu, B.: Modelling Rhizodeposition with Functional-Structural Plant Models, in: 10th 1675 International Society on Root Research Conference (ISRR-10), 2018.
- Rees, F., Barillot, R., Gauthier, M., Pagès, L., Pradal, C., and Andrieu, B.: Simulating rhizodeposition as a function of shoot and root interactions within a new 3D Functional-Structural Plant Model, in: *FSPM 2020 - 9th International Conference on Functional-Structural Plant Models*, pp. 22–23, Hanover / Virtual, Germany, <https://hal.inrae.fr/hal-02964060>, 2020.
- Rees, F., Gérard, T., Gauthier, M., Barillot, R., Richard-Molard, C., Jullien, A., Chenu, C., Pradal, C., and Andrieu, B.: Deciphering 1680 spatiotemporal patterns of rhizodeposition with a functional-structural root model: RhizoDep, *Plant and Soil*, 516, 777–795, <https://doi.org/10.1007/s11104-025-07766-z>, 2025.
- Richards, L. A.: Capillary conduction of liquids through porous mediums, *Physics*, 1, 318–333, <https://doi.org/10.1063/1.1745010>, 1931.
- Roose, T., Keyes, S., Daly, K., Carminati, A., Otten, W., Vetterlein, D., and Peth, S.: Challenges in imaging and predictive modeling of rhizosphere processes, *Plant and Soil*, 407, <https://doi.org/10.1007/s11104-016-2872-7>, 2016.
- 1685 Rougier, M.: Secretory Activity of the Root Cap, vol. 13/B of *Encyclopedia of Plant Physiology*, p. 772, Springer, Berlin, Heidelberg, ISBN 978-3-642-68234-6, <https://doi.org/10.1007/978-3-642-68234-6>, 1981.
- Ruiz, S., McKay Fletcher, D., Williams, K., and Roose, T.: Plant–Soil Modelling, vol. 4, pp. 127–198, John Wiley & Sons, Ltd, ISBN 9781119312994, <https://doi.org/10.1002/9781119312994.apr0755>, 2021.
- Schey, H. M.: *Div, grad, curl, and all that*, ISBN 0-393-96997-5, 1973.
- 1690 Schnepf, A., Huber, K., Landl, M., Meunier, F., Petrich, L., and Schmidt, V.: Statistical Characterization of the Root System Architecture Model CRootBox, *Vadose Zone Journal*, 17, <https://doi.org/10.2136/vzj2017.12.0212>, 2018a.

- Schnepf, A., Leitner, D., Landl, M., Lobet, G., Mai, T. H., Morandage, S., Sheng, C., Zörner, M., Vanderborght, J., and Vereecken, H.: CRootBox: a structural–functional modelling framework for root systems, *Annals of Botany*, 121, 1033–1053, <https://doi.org/10.1093/aob/mcx221>, 2018b.
- 1695 Schnepf, A., Carminati, A., Ahmed, M., Ani, M., Benard, P., Bentz, J., Bonkowski, M., Knott, M., Diehl, D., Duddek, P., Kroener, E., Javaux, M., Landl, M., Lehndorff, E., Lippold, E., Lieu, A., Mueller, C., Oburger, E., Otten, W., and Vetterlein, D.: Linking rhizosphere processes across scales: Opinion, *Plant and Soil*, 478, <https://doi.org/10.1007/s11104-022-05306-7>, 2022.
- Schnepf, A., Leitner, D., Landl, M., Khare, D., Heck, A., Giraud, M., Selzner, T., Helmrich, D., Lobet, G., Zhou, X., Bouvri, A., Ullah, S., Feron, T., Heymans, A., and Koch, T.: CPlantBox, branch Giraud2025_CarbonStabilisation, <https://doi.org/10.5281/zenodo.14809628>,
 1700 zenodo [code], 2025a.
- Schnepf, A., Leitner, D., Landl, M., Khare, D., Heck, A., Giraud, M., Selzner, T., Helmrich, D., Lobet, G., Zhou, X., Bouvri, A., Ullah, S., Feron, T., Heymans, A., and Koch, T.: dumux-rosi, branch Giraud2025_CarbonStabilisation, <https://doi.org/10.5281/zenodo.14809414>,
 zenodo [code], 2025b.
- Schröder, N., Lazarovitch, N., Vanderborght, J., Vereecken, H., and Javaux, M.: Linking transpiration reduction to rhizosphere salinity using a 3D coupled soil-plant model, *Plant and Soil*, 377, 277–293, <https://doi.org/10.1007/s11104-013-1990-8>, 2014.
- Schultes, S., Rüger, L., Niedeggen, D., Freudenthal, J., Frindte, K., Becker, M., Metzner, R., Pflugfelder, D., Chlubek, A., Hinz, C., Dusschoten, D., Bauke, S., Bonkowski, M., Watt, M., Koller, R., and Knief, C.: Photosynthate distribution determines spatial patterns in the rhizosphere microbiota of the maize root system, *Nature Communications*, 16, <https://doi.org/10.1038/s41467-025-62550-y>, 2025.
- Schwartz, N., Carminati, A., and Javaux, M.: The impact of mucilage on root water uptake—A numerical study, *Water Resources Research*,
 1710 52, 264–277, <https://doi.org/10.1002/2015WR018150>, 2016.
- Silva, L. C. R. and Lambers, H.: Soil-plant-atmosphere interactions: structure, function, and predictive scaling for climate change mitigation, *Plant and Soil*, 461, 5–27, <https://doi.org/10.1007/s11104-020-04427-1>, 2021.
- Sircan, A. K., Streck, T., Schnepf, A., Giraud, M., Lattacher, A., Kandeler, E., Poll, C., and Pagel, H.: Trait-based Modeling of Microbial Interactions and Carbon Turnover in the Rhizosphere, *Soil Biology and Biochemistry*, <https://doi.org/10.1016/j.soilbio.2024.109698>, 2025.
- 1715 Thorpe, M. R., Lacoite, A., and Minchin, P. E. H.: Modelling phloem transport within a pruned dwarf bean: a 2-source-3-sink system, *Funct Plant Biol*, 38, 127–138, <https://doi.org/10.1071/fp10156>, 2011.
- Tixier, A., Forest, M., Prudent, M., Durey, V., Zwieniecki, M., and Barnard, R. L.: Root exudation of carbon and nitrogen compounds varies over the day–night cycle in pea: The role of diurnal changes in internal pools, *Plant, Cell & Environment*, 46, 962–974, <https://doi.org/10.1111/pce.14523>, 2023.
- 1720 Trofymow, J. A., Coleman, D. C., and Cambardella, C.: Rates of rhizodeposition and ammonium depletion in the rhizosphere of axenic oat roots, *Plant and Soil*, 97, 333–344, <https://doi.org/10.1007/BF02383223>, 1987.
- van Genuchten, M. T.: A Closed-form Equation for Predicting the Hydraulic Conductivity of Unsaturated Soils, *Soil Science Society of America Journal*, 44, 892–898, <https://doi.org/10.2136/sssaj1980.03615995004400050002x>, 1980.
- Van Rossum, G. and Drake Jr, F. L.: Python tutorial, Centrum voor Wiskunde en Informatica Amsterdam, The Netherlands, 1995.
- 1725 Verbančič, J., Lunn, J. E., Stitt, M., and Persson, S.: Carbon Supply and the Regulation of Cell Wall Synthesis, *Molecular Plant*, 11, 75–94, <https://doi.org/10.1016/j.molp.2017.10.004>, 2018.
- Wang, G., Li, W., Wang, K., and Huang, W.: Uncertainty quantification of the soil moisture response functions for microbial dormancy and resuscitation, *Soil Biology and Biochemistry*, 160, <https://doi.org/10.1016/j.soilbio.2021.108337>, 2021.

- 1730 Wang, J., Bogena, H. R., Vereecken, H., and Brüggemann, N.: Characterizing Redox Potential Effects on Greenhouse Gas Emissions Induced by Water-Level Changes, *Vadose Zone Journal*, 17, <https://doi.org/10.2136/vzj2017.08.0152>, 2018.
- Wiesenbauer, J., König, A., Gorka, S., Marchand, L., Nunan, N., Kitzler, B., Inselsbacher, E., and Kaiser, C.: A pulse of simulated root exudation alters the composition and temporal dynamics of microbial metabolites in its immediate vicinity, *Soil Biology and Biochemistry*, 189, <https://doi.org/10.1016/j.soilbio.2023.109259>, 2024.
- 1735 Zhou, X.-R., Schnepf, A., Vanderborght, J., Leitner, D., Lacoïnte, A., Vereecken, H., and Lobet, G.: CPlantBox, a whole-plant modelling framework for the simulation of water- and carbon-related processes, in *silico Plants*, 2, <https://doi.org/10.1093/insilicoplants/diaa001>, 2020.