

# Modeling boreal forest soil dynamics with the microbially explicit soil model MIMICS+ (v1.0)

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**Abstract.** Understanding carbon exchange processes between land reservoirs and the atmosphere is essential for predicting carbon-climate feedbacks. Still, considerable uncertainty remains in the representation of the terrestrial carbon cycle in Earth System Models. An emerging strategy to constrain these uncertainties is to include the role of different microbial groups explicitly. Following this approach, we extend the framework of the Microbial-MIneral Carbon Stabilization (MIMICS) model

5 with additional mycorrhizal groups and a nitrogen cycle that includes a novel representation of inorganic nitrogen sorption to particles via a Langmuir isotherm. MIMICS+ v1.0 is designed to capture and quantify relationships between soil microorganisms and their environment, with a particular emphasis on boreal ecosystems. We evaluated MIMICS+ against podzolic soil profiles in Norwegian forests as well as the conventional Community Land Model (CLM). MIMICS+ matched observed carbon stocks better than CLM, and gave a broader range of C:N ratios, more in line with observations. This is mainly explained  
10 by a higher direct-plant-derived fraction into the Soil Organic Matter (SOM) pools. The model produces microbial biomass estimates in line with numbers reported in the literature. MIMICS+ also showed better representation of climate gradients than CLM, especially in terms of temperature. To investigate responses to changes in nutrient availability, we performed an N enrichment experiment, and found that nitrogen sorbed to particles through the sorption algorithm served as a long-term storage  
15 of nutrients for the microbes. Furthermore, although the microbial groups responded considerably to the nitrogen enrichment, we only saw minor responses for carbon storage and respiration. Together, our results present MIMICS+ as an attractive tool for further investigations of interactions between microbial functioning and their (changing) environment.

## 1 Introduction

Among the carbon (C) stores in the terrestrial biosphere, soils are the largest, containing ca. 1700 GtC, while vegetation accounts for ca. 450 GtC globally (Friedlingstein et al., 2022). The active exchange of C between terrestrial pools and the  
20 atmosphere is affected by elevated CO<sub>2</sub> concentrations and changes in N deposition, but quantifying the responses has proven to be a central challenge in climate science. Arora et al. (2020) highlight the uncertainty in terrestrial carbon-concentration and carbon-climate feedbacks from the last model intercomparison project, CMIP6. The uncertainty of carbon-cycle feedbacks is up to one order of magnitude larger for land than for ocean, illustrating the need to improve model representation of terrestrial processes. To do this, we need to represent complex C and nutrient cycle processes in a modeling framework, a task that

25 requires careful consideration of how to translate real-world processes into an appropriate model form. Fisher and Koven (2020) suggest an approach based on modular complexity. Dividing a full-complexity land model into smaller modules allows investigation of alternatives for structure and parameter choices, which helps in making good modeling choices and thereby constrain sources of uncertainty.

Large variations in responses between different biomes introduce an extra challenge to C cycle modeling. The impact of 30 environmental changes on boreal systems is of particular interest for several reasons. For example, studies show that the kinetics of soil microbes accustomed to cooler climates are more temperature sensitive than microbes in warmer climates (German et al., 2012). Koven et al. (2017) also showed that soil carbon turnover times in cold areas are more sensitive to 35 climatological temperature than in warm areas. Many boreal areas also experience treeline migration caused by an expansion of the temperature-limited area where tree species can grow (Hansson et al., 2021). Often this leads to a shift in mycorrhizal 40 associations, for example from arbuscular mycorrhiza (AM) to ectomycorrhiza (EcM), which again can lead to changes in soil carbon dynamics and below-ground carbon storage (Taylor et al., 2016; Tonjer et al., 2021). EcM have been found to alter decomposition, either negatively through increased nutrient competition with saprotrophs (Gadgil and Gadgil, 1971, 1975) or positively through priming effects (Brzostek et al., 2015; Phillips et al., 2012) based on environmental context (Fernandez and Kennedy, 2016). Recent findings also suggest that differences in decomposability of necromass from different mycorrhizal 45 groups can impact soil C storage more than previously thought (Huang et al., 2022a, b). In Norwegian forests, vegetation is typically dominated by evergreen, coniferous trees, mainly associated with EcM. The dominating soil type in these forests is podzol (Strand et al., 2016). Podzols are typically nutrient-poor, and competition for nutrients is expected to be important for the carbon dynamics in these systems. Despite the importance of boreal systems, many soil model structure and parameter choices are based on temperate or tropical observations. This bias may skew model results, and make the modeled responses 50 to climate change in boreal environments more uncertain.

Nitrogen (N) is one of the most important nutrients in an ecosystem, and the cycling of nitrogen between above-ground and 55 below-ground reservoirs can greatly affect carbon dynamics. In addition to regulating forest productivity, N availability regulate microbial Carbon Use Efficiency (CUE), as microbes respire excess C to meet their stoichiometrical demand (Mooshammer et al., 2014b). This direct relationship between soil N and the C exchange between the atmosphere and soils emphasizes the importance of including microbial C-N relationships in C cycle models. One factor determining nitrogen availability in an ecosystem is inorganic N deposition from the atmosphere and agricultural fertilization. This inorganic N is subject to physical and chemical processes that affect how readily available the N is to microbes and plants. One such process is cation exchange, which controls storage and release of ammonium ( $\text{NH}_4^+$ ) from negatively charged clay particles and organic molecules (Bonan, 2015), and therefore impacts inorganic N availability for microbes and plants. This is a process that might be extra important 55 in nutrient-poor boreal forest systems. There are studies examining this effect in agricultural soils (Sieczka and Koda, 2016), but few are looking at natural soils.

Traditionally, decomposition processes in models have been represented by first-order kinetics for litter, as well as active, slow, and passive pools of Soil Organic Matter (SOM) (Parton et al., 1988). This approach limits the ability to examine the mechanisms and possible responses of the soil system during climate change (Todd-Brown et al., 2012). Newer work has

60 introduced models that in different ways represent microbial activity explicitly (e.g., Wieder et al., 2015; Sulman et al., 2019; Fatichi et al., 2019; Yu et al., 2020; Huang et al., 2018; Wang et al., 2013). These models increase the possibility to capture carbon climate feedbacks of the future (Tang and Riley, 2014; Hararuk et al., 2015). Wieder et al. (2015) illustrated that by representing the functional traits of microbes in the MIMICS model, one can raise important hypotheses about how microbes can determine responses to for example N enrichment. Kyker-Snowman et al. (2020) further showed that adding an N cycle to 65 the MIMICS model (MIMICS-CN) produced results in line with measurements from North American sites, and comparable models. Wang et al. (2021) presented a vertically resolved C-only version of MIMICS and showed that microbial activity and root carbon inputs were more important than the soil carbon diffusion when simulating soil carbon concentration profiles.

70 Baskaran et al. (2017) introduced a model that emphasized the influence of EcM on decomposition, however without the ability to capture nutrient competition with saprotrophic microbes. We included EcM with parameterizations from Baskaran et al. (2017) in a modeling framework based on the MIMICS model (Wieder et al., 2015) that also includes explicit saprotrophic pools. To capture possible shifts in mycorrhizal associations, we also included an arbuscular mycorrhizal (AM) pool using methods presented by Sulman et al. (2019). In contrast to the always available inorganic N pools in Sulman et al. (2019), we 75 introduced an algorithm for representing sorption of ammonium to soil particles based on the Langmuir isotherm (Sieczka and Koda, 2016), which may be an important but underrepresented process determining the availability of inorganic N to soil microbes in boreal forests. We assume that by including processes and parameters thought to be particularly relevant for climate responses in colder areas we can obtain a better understanding of the C dynamics, and thereby reduce uncertainty connected to soil processes. A future goal is to couple the soil model to a land model/ESM with interactive vegetation, and although our present emphasis is on boreal systems, the incorporated processes are general and representative on a larger scale.

80 We introduce a vertically resolved, microbially explicit soil decomposition model, MIMICS+, that represents C and N flows between litter, microbial, and SOM pools. In this study the model is offline, and forced with data produced by the Community Land Model v5.1 (CLM; Lawrence et al. (2019)). C and N stock estimates from the CLM simulations represent a microbially implicit approach based on the traditional CENTURY model (Parton et al., 1988). Therefore, we compare the CLM and MIMICS+ results to investigate the implications of including the processes and mechanisms mentioned above. To evaluate the model, we use a collection of soil profile data from forested, podzolic sites in Norway, covering a range of conditions 85 representative of boreal systems (Strand et al., 2016). Our experimental setup is as follows; for a selection of 50 sites in Norway, we ran simulations with the CLM model to produce a) input data needed to run MIMICS+, and b) estimates of C and N stocks. We then ran MIMICS+ with the produced forcing data. The aims of the study are 1) to formulate a standalone, microbially explicit model capable of representing soil processes in boreal systems, 2) to evaluate model performance and model structure by comparing simulated vertical soil C content along a climatic gradient with observations and simulated soil 90 carbon from the microbially implicit model CLM 3) Apply the model to perform an N enrichment experiment to investigate below-ground responses to nutrient changes.

## 2 Model and Methods

### 2.1 Model Description

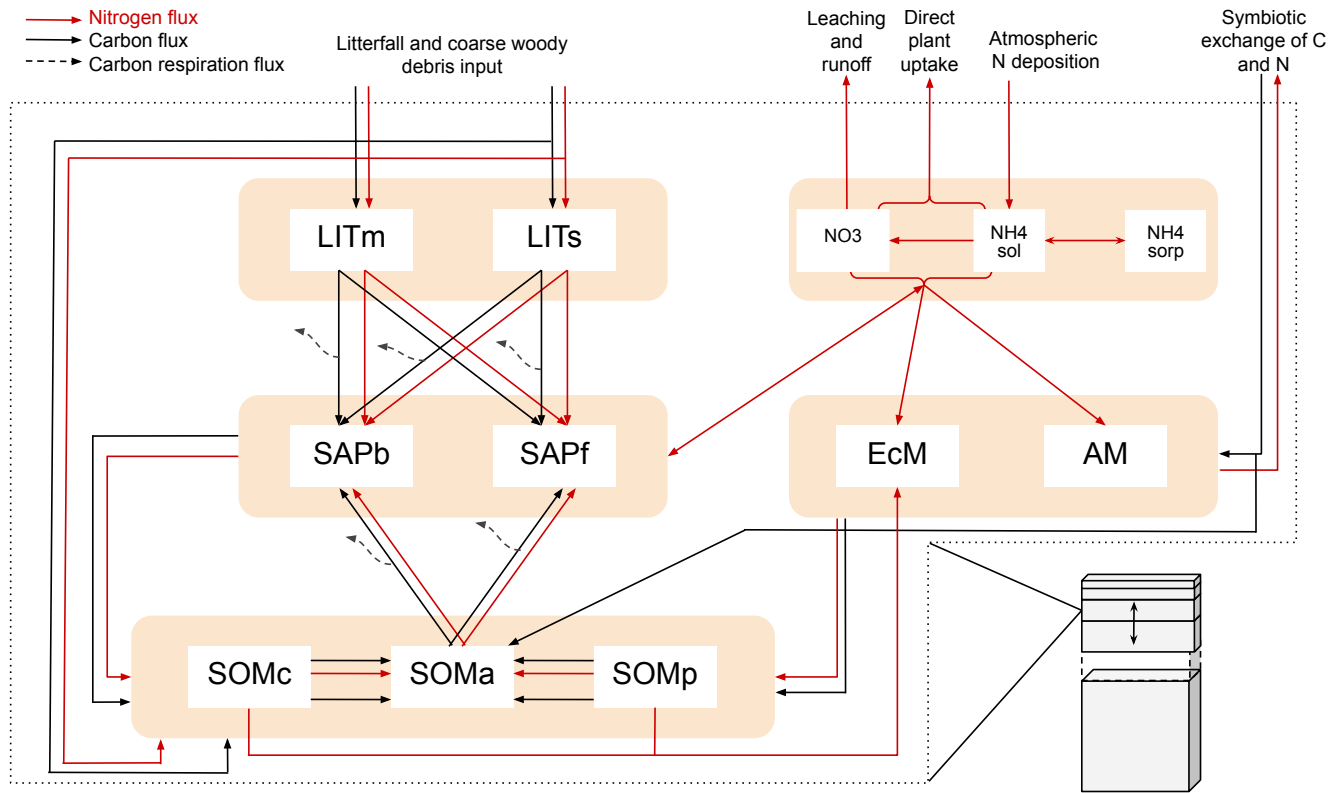
MIMICS+ is based on the MIMICS framework where microbial groups, litter and soil organic matter are represented as separate pools (Wieder et al., 2015). In its current state MIMICS+ is not coupled to a comprehensive land model, and therefore needs prescribed C and N input, and soil temperature and moisture, which it is set up to read from CLM history files. Mass balance equations,  $dP/dt = \text{Sources} - \text{Sinks}$ , determine the change at each timestep for each pool, P. The model structure with pools and fluxes is illustrated in Fig. 1, and a detailed overview of mass balance and rate equations are provided in the Appendix; Tables A1 and A2 contain mass balance equations for C and N pools, respectively, while Tables A3 and A4 list C and N rate equations. Throughout the model description, fluxes referred to as "CX" or "NX", where X is a number, can be found in those tables, and are illustrated as arrows in Fig. 1 and Fig. A1. A list of parameters is given in Table A5. By representing the same hydrologically and biogeochemically active layers as in CLM, MIMICS+ can represent the depth discretization of temperature and moisture dependent processes. For each layer the fluxes between the pools within the layer are calculated first, before the vertical transport is calculated. Unless otherwise stated, the equations below describe transport within one layer. The vertical transport is described in Sect. 2.1.4.

#### 2.1.1 Litter and SOM pools

Organic C and N enters the litter and SOM pools as dead plant material. As in MIMICS (Wieder et al., 2015) and ORCHIDEE-SOM (Camino-Serrano et al., 2018), we separate between metabolic (labile) litter mainly originating from leaves and fine roots, and structural litter, in which we also include Coarse Woody Debris (CWD). For SOM we again follow the MIMICS approach with two protected SOM pools, and one pool that is available for saprotrophic decomposition. Depolymerization and desorption ~~moves~~move organic matter from chemically and physically protected pools, respectively, to the available pool (C11, C12, N11, N12). The depolymerization process ~~represent~~represents the enzymatic breakdown of recalcitrant SOM, and is thus modeled with an rMMK mechanism, while the desorption is a function of clay content, as this rate is representing the physical desorption from mineral surfaces (Wieder et al., 2015). 50 % of the incoming metabolic and structural litter goes to physically and chemically protected SOM, respectively, as direct plant-derived SOM (C3, C4, N3, N4). The direct litter fluxes, together with microbial necromass (C13-C24, N13-N24) and a flux representing EcM enzyme production (C27) are the sources of input to the SOM pools. The microbial pools determine the rates of decomposition, and thereby the transfer rates between the main storage pools; SOM and litter.

#### 2.1.2 Microbial processes

MIMICS+ represents two different types of microbes: saprotrophs and mycorrhizal fungi. Within these two main groups we separate between two functional traits, giving four different microbial pools in total. We divide between saprotrophic fungi (SAPf; analog to MIMICS k-strategists) and bacteria (SAPb; analog to MIMICS r-strategists). Temperature-sensitive reverse



**Figure 1.** Schematic showing C and N flows within each layer of the model. Black arrows indicate carbon fluxes ( $\text{gCm}^{-3}\text{h}^{-1}$ ) while red arrows indicate nitrogen fluxes ( $\text{gNm}^{-3}\text{h}^{-1}$ ). The dashed black arrows symbolize C leaving the system as heterotrophic respiration. LITm, LITs: metabolic and structural litter. SAPb, SAPf: saprotrophic bacteria and fungi. EcM, AM: ecto- and arbuscular mycorrhizal fungi. SOMc, SOMa, SOMp: chemically protected, available and physically protected soil organic matter. NO<sub>3</sub>, NH<sub>4</sub><sub>sol</sub>, NH<sub>4</sub><sub>sorb</sub>: Inorganic N in the form of NO<sub>3</sub>, NH<sub>4</sub> in solution and NH<sub>4</sub> sorbed to particles, respectively.

Michaelis-Menten kinetics, together with a moisture modifier (Wieder et al., 2017), determines the rates at which saprotrophs decompose substrate from the two litter pools, and the available SOM (C5-C10, N5-N10). The N fluxes are determined by the stoichiometry of the substrate pools. During decomposition, a fraction of the incoming C is lost from the soil as heterotrophic respiration (HR), while the rest is contributing to saprotrophic biomass. The respiration fraction is determined by the carbon use efficiencies CUE<sub>b</sub> and CUE<sub>f</sub> which have maximum values of 0.4 and 0.7 for bacteria and fungi respectively, but is reduced under low nutrient conditions. This is based on the theory that microbes adjust their efficiencies to maintain a relatively constant, low C:N ratio despite the higher C:N ratio of substrates (Mooshammer et al., 2014b). The C:N ratio of the model saprotrophs is assumed to be constant ( $\text{CN}_b = 5$  and  $\text{CN}_f = 8$ , Table A5). To ensure that this ratio is fulfilled in each layer and time step (in addition to potentially reducing CUE) N is exchanged between the saprotrophs and the inorganic pools (N36 and N37). The

exchange rates can be positive or negative, leading to either immobilization or mineralization of inorganic N. We first calculate the uptake and demand of N to determine if there is enough to meet the requirement for optimal saprotrophic functioning.

$$N_{demand,x} = \frac{CUE_x \cdot (FC_{LITm,SAPx} + FC_{LITs,SAPx} + FC_{SOMa,SAPx})}{CN_x} \quad (1)$$

135

$$N_{uptake,x} = NUE \cdot (FN_{LITm,SAPx} + FN_{LITs,SAPx} + FN_{SOMa,SAPx}) \quad (2)$$

Here, x represents either b (bacteria) or f (fungi) and NUE is Nitrogen Use Efficiency, further described below. This results in one of four possibilities:

1. The N demand is greater than the uptake for both bacteria and fungi, meaning both groups will immobilize inorganic N. In this case we check if there is enough available inorganic N to fulfill the demand from both groups. If not, CUE is reduced (according to Eq. (3) and (4)) so that the saprotrophs utilize all N that is available to them, before the demand is recalculated. Here,  $N_{for\_sap}$  is referring to the sum of the available N pools,  $N_{NH4,sol}$  and  $N_{NO3}$ :

$$CUE_b = \frac{(f_b \cdot N_{for\_sap} + N_{uptake,b} \cdot dt) \cdot CN_b}{(FC_{LITm,SAPb} + FC_{LITs,SAPb} + FC_{SOMa,SAPb}) \cdot \Delta t} \quad (3)$$

$$145 \quad CUE_f = \frac{((1 - f_b) \cdot N_{for\_sap} + N_{uptake,f} \cdot dt) \cdot CN_f}{(FC_{LITm,SAPf} + FC_{LITs,SAPf} + FC_{SOMa,SAPf}) \cdot \Delta t} \quad (4)$$

where  $f_b$  determines the division of the available inorganic N between bacteria and fungi, and is calculated as:

$$f_b = \frac{(N_{demand,b} - N_{uptake,b})}{((N_{demand,b} - N_{uptake,b}) + (N_{demand,f} - N_{uptake,f}))} \quad (5)$$

Equation (3) and (4) reduces CUE (and increase the respired fraction) enough to maintain the C:N ratios under the prevailing conditions, and the resulting exchange rates is:

$$150 \quad FN_{IN,SAPb} = f_b \cdot N_{for\_sap} \quad (6)$$

$$FN_{IN,SAPf} = (1 - f_b) \cdot N_{for\_sap} \quad (7)$$

2. N uptake is larger than demand for both saprotrophic groups, meaning both will mineralize inorganic N. The mineralized N will enter the  $N_{NH4,sol}$  pool.

155 3. Fungi will mineralize N (uptake > demand), while bacteria immobilizes N (uptake < demand). In this case bacteria can access the N mineralized by fungi in addition to the inorganic N if needed.

4. Bacteria will mineralize N (uptake > demand), while fungi immobilizes N (uptake < demand). In this case fungi can access the N mineralized by bacteria in addition to the inorganic N if needed.

160 Saprotrrophic necromass is transferred to the SOM pools, and is partitioned between the three pools based on clay content of the soil and the metabolic fraction of incoming litter (C13-C18 and N13-N18). Only a fraction of the N released during decomposition is directly available to saprotrophs, determined by the NUE (constant NUE = 0.8, Mooshammer et al. (2014a)). The remaining fraction is transferred to N<sub>NH4,sol</sub>.

165 The model represent two different types of mycorrhizal fungi: EcM and AM. The mycorrhizal pools receive a C supply from plants, and in return provides N to its associated plants. How the incoming carbon ( $I_{veg,MyC}$ , cf. C28 and C29) is partitioned between EcM and AM is determined dynamically through a Return Of Investment (ROI) function based on the method from Sulman et al. (2019). The partition between EcM and AM is determined as a fraction,

$$f_{alloc,i} = \frac{ROI_i}{\sum_i ROI_i} \quad (8)$$

where  $ROI_i$  is the nitrogen return of the carbon investment from mycorrhizal association  $i$  (EcM or AM);

$$ROI_i = \frac{N_{acquired,i} \cdot \tau_{myc,som} \cdot CUE_i}{C_i} \quad (9)$$

170 EcM acquires N from the protected SOM and inorganic N pools ( $N_{acquired,EcM} = N25 + N26 + N27$ ) while AM only acquires inorganic N ( $N_{acquired,AM} = N28$ ).  $\tau_{myc,som}$  is the mycorrhizal turnover time, while  $CUE_i$  is the growth efficiency for mycorrhizal association  $i$ . N25 and N26 represent ectomycorrhizal mining for N (Lindahl and Tunlid, 2015). By releasing enzymes (C27) EcM access N from protected SOM, and at the same time release C to the available SOM pool (C25 and C26). The enzyme production is modeled as a fraction of the incoming carbon (C28) that is directed into the SOMa pool instead of the EcM pool (C27). The mining algorithm is based on Baskaran et al. (2017), with mycorrhizal "decomposition" modeled as a multiplicative function of mycorrhizal biomass, SOM, and a decay rate ( $K_{mo}$ , Table A5). We use this expression together with the C:N ratio of the substrate pool to determine the amount of nitrogen acquired through ectomycorrhizal mining (N25 and N26).

175 As the mycorrhizal pools are assumed to have constant C:N ratios, a part of the acquired N is used to fulfill the stoichiometric constraint. Any additional acquired N is leaving the soil system as N supply to the plant. The prescribed C supply from CLM is zero during the winter months, so to ensure that the mycorrhizal fungi do not provide "free" N to the plant during this time, we introduce a scaling factor;

$$r_{myc} = \frac{I_{veg,myc}(t)}{\max(I_{veg,myc})} \quad (10)$$

180 Here,  $I_{veg,myc}(t)$  (gCm<sup>-2</sup>h<sup>-1</sup>), is the time varying C supply from vegetation (prescribed from CLM), and  $\max(I_{veg,myc})$  is the maximum value of  $I_{veg,myc}$  in the current year. This scaling factor means that the mycorrhizal fungi are most effective when they receive most energy in the form of C. Since  $I_{veg,myc}(t)$  is prescribed in the current model version, the input flux will not respond to changes in soil N availability. Constant mortality rates determine the transfer from mycorrhizal fungi to the SOM pools (C19-C24 and N19-N24).

### 2.1.3 Inorganic N processes

190 Inorganic N is divided between nitrate and ammonium dissolved in soil water ( $N_{NO_3}$  and  $N_{NH_4, sol}$ ), and ammonium sorbed to soil particles ( $N_{NH_4, sorb}$ ). Reactive nitrogen from atmospheric deposition enters  $N_{NH_4, sol}$  (N32) where it can undergo nitrification to  $N_{NO_3}$  (N34) or become sorbed to particles (N35).  $N_{NO_3}$  is exposed to leaching and runoff based on CLM algorithms (N31). Both dissolved pools,  $N_{NH_4, sol}$  and  $N_{NO_3}$ , can be taken up by mycorrhizal fungi (N27, N28) or directly by plants (N33). Since the model is not coupled to above-ground vegetation, direct plant uptake is a constant loss rate of the 195 available inorganic N ( $k_{plant}$ ). We assume that processes in boreal forests are relatively slow, and that the residence times of the pools are much longer than the one-hour time step. We therefore apply a sequential approach to model the mass balance of the inorganic N pools. Within a time step (1 hour) the different processes affecting inorganic N is calculated in a sequence: 1) Deposition, leaching and runoff 2) nitrification 3) N from decomposition 4) direct uptake by vegetation 5) uptake by mycorrhiza 6) exchange with saprotrophs 7) Langmuir sorption algorithm. The Langmuir sorption algorithm is based on Sieczka and Koda 200 (2016) and described below. The basis for this process is cation exchange, where positively charged ammonium is adsorbed to negatively charged clay particles. Before pt. 7) the total concentration of ammonium is

$$N_{NH_4, tot} = N_{NH_4, sorp} + N_{NH_4, sol} \quad (11)$$

Using Eq. (11) together with the Langmuir isotherm equation, we find the equilibrium partition between  $N_{NH_4, sol}$  and  $N_{NH_4, sorp}$  given the total concentration  $N_{NH_4, tot}$ . The Langmuir isotherm equation is given by

$$205 N_{NH_4, sorp, eq} = \frac{NH_4_{sorp, max} \cdot K'_L \cdot N_{NH_4, sol, eq}}{1 + K'_L \cdot N_{NH_4, sol, eq}} \quad (12)$$

where  $K'_L$  is a Langmuir constant related to adsorption energy, and a function of soil water content.  $NH_4_{sorp, max}$  is the maximum adsorption capacity. We assume that the system moves towards the equilibrium value during the timestep, via the following mechanism, derived from the pseudo-second order kinetic model in Sieczka and Koda (2016):

$$N_{NH_4, sorp} = \begin{cases} N_{NH_4, sorp, eq} - \frac{1}{N_{NH_4, sorp, eq} - N_{NH_4, sorp, prev.} + k \cdot \Delta t} & N_{NH_4, sorp, eq} > N_{NH_4, sorp, prev}, \\ N_{NH_4, sorp, eq} + \frac{1}{N_{NH_4, sorp, prev} - N_{NH_4, sorp, eq} + k \cdot \Delta t} & N_{NH_4, sorp, eq} < N_{NH_4, sorp, prev}, \\ N_{NH_4, sorp, prev} & N_{NH_4, sorp, eq} = N_{NH_4, sorp, prev} \end{cases} \quad (13)$$

Here  $k$  is a rate constant and  $\Delta t$  is the timestep. The top option corresponds to absorption, the middle option to desorption and the third if equilibrium is already reached. All parameter values are from Sieczka and Koda (2016), converted to appropriate 210 model units (see Table A5).

## 2.1.4 Vertical structure

The discrete vertical layers of the model follow the same structure as CLM with increasing layer thickness with depth (Lawrence et al., 2019). This allows incoming litter and N deposition to be distributed following the same vertical profile as in CLM. We use vertically resolved soil temperature and soil moisture from CLM as input to MIMICS+. We also use 215 drainage and runoff rates from CLM to determine N leaching. Within each time step the fluxes between the pools are calculated and applied first, then vertical transport is calculated and applied. This transport is calculated as a simple diffusion equation between adjacent layers (Soetaert and Herman, 2009), using a constant diffusion coefficient from Koven et al. (2013). As the vertically resolved soil temperature and soil moisture from CLM are used in MIMICS+, the saprotrophic decomposition 220 rates that are functions of these variables have a depth dependency in the model. The mycorrhizal N uptake are functions of the amount of mycorrhizal biomass and inorganic N (and SOM for EcM) in the soil layer, hence uptake can vary with depth.

## 2.1.5 Parameter sensitivity analysis

To test the soil C sensitivity to different model parameters, we performed a sensitivity analysis on 16 key parameters. For one parameter at a time, we either increased or decreased the value ~~with-by~~ 25 % compared to the default, giving a total of 32 experiments which was performed for each of the 50 sites simulated in this study (see Sect. 2.2).

## 225 2.2 Soil profile database

For comparison, a forest soil database collected in connection with the ICP forest monitoring program level 1 sites was used (Lorenz, 1995). These data have been further analyzed by Strand et al. (2016), and provide a unique source of information about boreal soil conditions. A total of 1040 soil profiles were described, sampled and analyzed between 1988 and 1992 (Esser 230 and Nyborg, 1992). Soil profile descriptions were done according to standardized procedures (Sveistrup, 1984) and classified according to the Canadian System of Soil Classification (CSSC). Relevant information from the database includes C and N stocks, Mean Annual Temperature (MAT) and Mean Annual Precipitation (MAP). Specifically, the database contains C content down to 30 cm, 50 cm and 100 cm, making it possible to compare vertically modeled C stocks to observations in these depth 235 intervals. The dataset also contains separate measurements of C and N in the organic LFH (Litter, Fermented, Humic) layer and mineral soil. The organic layer consists of more or less decomposed litter, and although not directly comparable to modeled litter and SOM pools, the C:N ratio in organic vs. mineral soil is still a useful quantity for model evaluation purposes. A more detailed description of the database is given in Strand et al. (2016). Because podzols are the most common soil category in Norwegian forests, we chose to focus on the podzolic sites in the dataset, giving a total of 578 sites. Due to computational 240 resource limitations, we chose a subset of 50 representative sites (out of the 578) for the site simulations with CLM and MIMICS+. The remaining 528 sites were used for further comparison with the modeled carbon stocks. The 50 sites cover an area from 5° W to 29°W longitude, and from 5° N to 70° N latitude. The MAT varies from -1.3°C to 7°C, while MAP ranges from 356 to 2510 mm · y<sup>-1</sup>.

**Table 1.** MAT and MAP intervals for dividing the sites into climate categories

Data source	cooler [°C]	warmer [°C]	drier [ $\text{mm} \cdot \text{yr}^{-1}$ ]	wetter [ $\text{mm} \cdot \text{yr}^{-1}$ ]
Observed	(-1.3)–2.5	2.6–7.1	355–975	1009–2510
Model forcing	(-1.8)–3.8	3.9–8.1	494–1243	1244–3606

## 2.3 Simulation setup

For the subset of 50 sites, we performed single-site simulations using CLM5.1 in BGC (biogeochemistry) mode. Data from these simulations were used both to force MIMICS+, and to compare the C and N stocks as calculated by the standard decomposition model in CLM. The CLM variables that are used to force MIMICS+ are listed in Table A6. For the simulations we assume that all C allocated to active N uptake by plants in CLM is directed to mycorrhiza (in default CLM this C is assumed to directly respire).

The observations were performed during the years 1988–1992, so we ran the models up to and including 1992, and averaged model values over the five years. Unless otherwise stated, these averages is what is used for the comparisons. The three datasets each containing data from 50 sites will be referred to as OBS (observations from database), CLM (CLM simulations) and MIMICS+ (MIMICS+ simulations with CLM forcing). An overview of yearly mean input of carbon and nitrogen is shown in Fig. C1.

For the CLM simulations, a single site configuration with 100 % natural vegetation was used together with atmospheric forcing from the Global Soil Wetness Project forcing data set (GSWP3, <https://hydro.iis.u-tokyo.ac.jp/GSWP3/>). This is the default atmospheric forcing for CLM and provides 3-hourly data with 0.5° resolution. Following CLM spin-up protocol (Lawrence et al., 2019), all sites were spun up for 500 years in "accelerated-decomposition" mode followed by 700 years of "regular spinup" by recycling atmospheric forcing for 1901–1930. For the period 1850–1900 the atmospheric forcing cycles the years 1901–1920, then historical forcing was used until the end of the simulation.

As with the CLM simulations, MIMICS+ needs to be spun up to equilibrium before running a historical period. The spinup was performed from arbitrary initial concentrations by recycling monthly averages of soil temperature and moisture, N deposition, litter and C input from the CLM history files for the years 1850–1869 (which is using atmospheric forcing from 1901–1920) for 1000 years.

### 2.3.1 Comparison of climate gradient profiles

To examine how well the models capture variation with temperature, the three datasets (OBS, MIMICS+, CLM) were sorted by increasing MAT. The first half (N=25) was labeled "cooler", while the second half (N=25) was labeled "warmer". To capture variation in moisture, the sites were sorted by MAP in the same manner, with the first half labeled "drier" and the second half labeled "wetter". Because the MAP and MAT data from the observations and the model forcing ~~differs, a differ~~, some sites ended up in different categories depending on whether they were sorted by the observed or forcing climate data (12 sites for

270 MAT and 8 sited sites for MAP). Since we are investigating specific sensitivities to temperature and precipitation, we chose to split the datasets based on their local climate forcing (We split the dataset this way—OBS by observed climate and MIMICS+ and CLM by model forcing climate), and not geographic location—because we are investigating sensitivities to temperature and precipitation. (Fig. S1 shows distribution based on geographic location)results of this analysis with all points classified according to their OBS climate.) The MAT and MAP intervals for each category are given in Table 1. For some sites the measured soil depth was shallower than 50 cm or 100 cm. These sites, where the depth to bedrock was less than 50 cm or 100 cm, were removed from both the model and observation datasets before making distribution box plots for these depth intervals.

### 2.3.2 N enrichment experiment

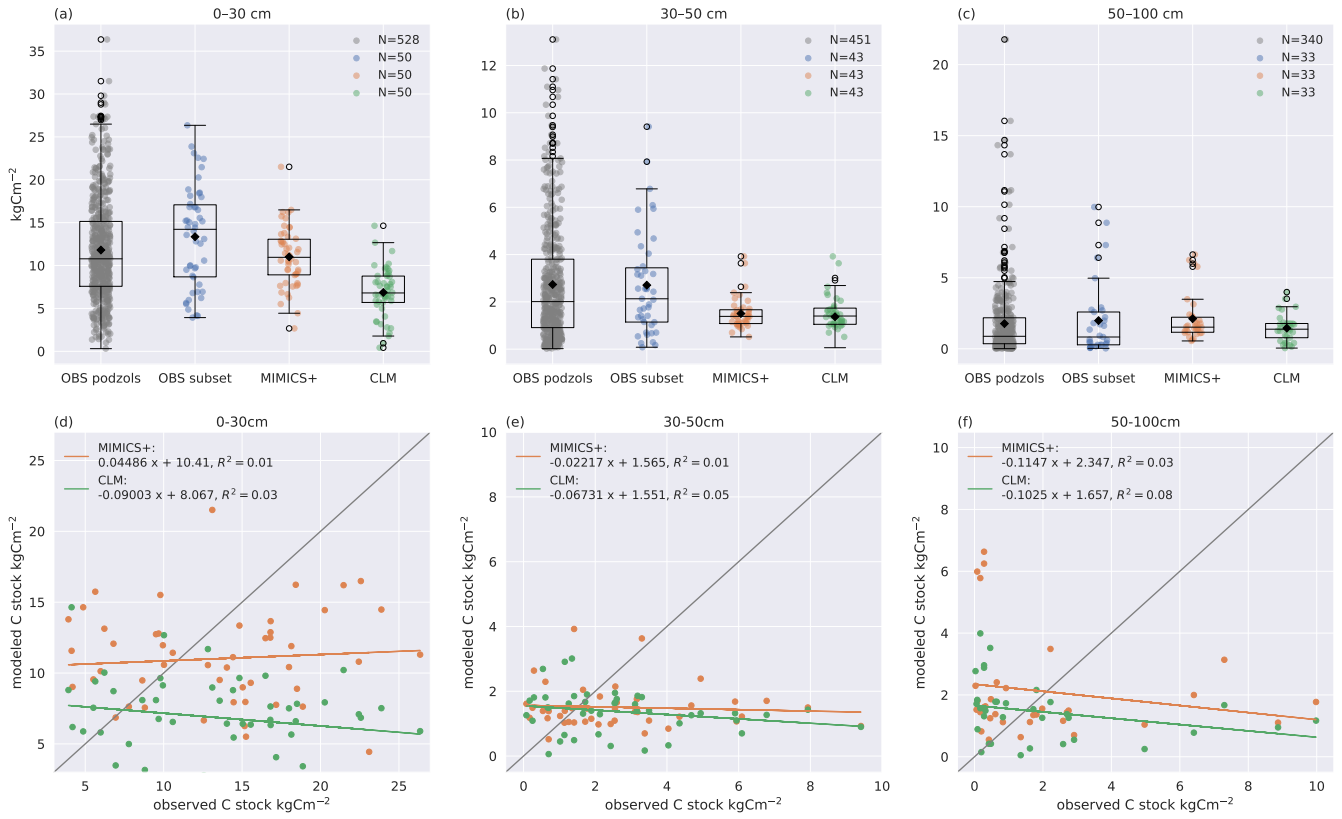
275 To investigate the MIMICS+ modeled response to N enrichment we performed an idealized N addition experiment. Starting from spun-up conditions, we ran two parallel simulations for all 50 sites for 30 years; one "control", using N deposition from the CLM runs, and one "treatment", with an extra amount of  $15 \text{ gNm}^{-2}\text{y}^{-1}$  deposited. This is a common amount used in forest fertilization (Högberg et al., 2017). The additional nitrogen was added equally in each time step throughout the second simulation year to give a total of  $15 \text{ gNm}^{-2}$ . We used these simulations to investigate the temporal response ratios (RR = treatment:control) for different C and N pools, as well as for HR.

## 3 Results

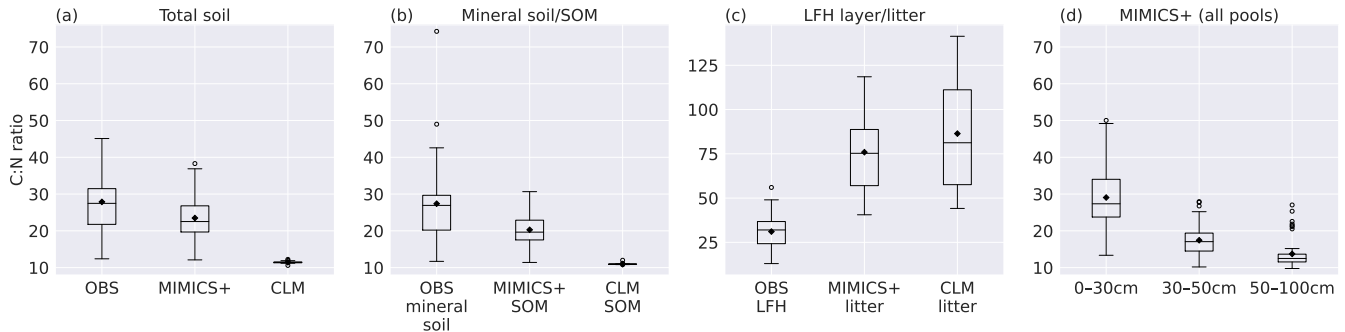
### 3.1 Comparison of modelled and empirical C and N stocks

285 Observed and modelled soil carbon stocks are shown in Fig. 2. Both models capture the general trend of decreasing C concentration with increasing depth. The modeled mean C stocks of MIMICS+ across the 50 sites are closer to observations in the 0–30 cm depth interval while the CLM simulations clearly underestimate C stocks (both models are significantly different from the subset of observations,  $p < 0.05$ ). The models both underestimate carbon at the 30–50 cm interval, while there is no significant difference between the modeled and observed C content in the deepest layer. Due to the heterogeneous nature of 290 real soils and the impact of differences of litter production between the sites, a larger variability in the observations compared to the simulations is not unexpected. However, site-to-site comparisons with observations are poor for both models, although marginally better for MIMICS+ (Fig. 2d–f). This is likely explained by subgrid variability in the observations that are not captured by the models and their forcing. As the model is intended to work on larger spatial scales within an ESM model, a good one-to-one match with specific sites is of less importance than being able to capture larger patterns in temperature and 295 moisture. By looking at the collection of sites together, we remove some of the uncertainty related to the variability between the sites and focus on larger patterns in our analyses. There is no significant difference between the two observational subsets, meaning that the 50 sites chosen for the direct model comparison is representative for the broader region.

Looking at C:N ratios, the overall picture with a higher ratio in the forest floor (observations) and litter pools (models), than in the total soil is captured by both models, again MIMICS+ being closer to the observed values (Fig. 3a and b). Both



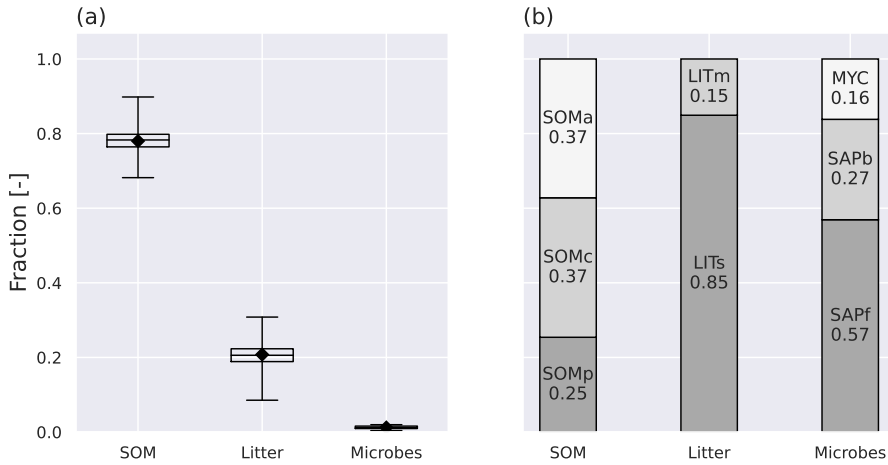
**Figure 2.** Modeled and observed C stocks. (a)-(c): box plots of C stocks in (a) 0–30 cm, (b) 30–50 cm, (c) 50–100 cm soil depths for; all observed podzols except the 50 modeled ones (left), the 50 modeled sites (center-left) from Strand et al. (2016), simulated with MIMICS+ (center-right) and with CLM (right). The line in each box is the median, while the diamonds mark the mean values. The box upper and lower edges are the 75th and 25th percentiles, respectively. The whiskers extend from the box by 1.5 times the inter-quartile range. Note the different scales on the y-axes. As not all observed soil profiles are reaching a depth of 30–50 cm or 50–100 cm, these sites are omitted in all boxplots for these depths, hence N=43 for (b) and N=33 for (c). (d)-(f): scatterplots of observed (x-axis) and modeled (y-axis) C stocks in (d) 0–30 cm, (e) 30–50 cm, (f) 50–100 cm soil depths. The legend shows the slope, intercept, and  $R^2$  for the linear regression line fitted to the scatter points. The 1:1 line is added in grey for reference.



**Figure 3.** Box plots of C:N ratios for observed values from Strand et al. (2016), MIMICS+ and CLM simulations of (a) Total soil, (b) mineral soil (observations) sum of SOM pools (models) (c) the observed forest floor compared to the C:N ratio of simulated litter pools, (d) Total soil at different depths as simulated by MIMICS+. Inorganic N is not considered in any of the plots. The line in each box is the median, while the diamonds mark the mean values. The box upper and lower edges are the 75th and 25th percentiles, respectively. The whiskers extend from the box by 1.5 times the inter-quartile range. N=50 sites.

300 models have significantly lower C:N ratios in the total as well as in the mineral soil, but MIMICS+ has significantly higher values than CLM ( $p < 0.05$ ). For the litter pools, the pattern is opposite, and the models have significantly higher C:N litter ratios than observed in the LFH layer. The modeled litter pools are not directly comparable to the LFH layer, but we get an indication of how the modeled C:N ratio compares to the partly decomposed matter. Both models have higher mean values and greater variability than the observations (Figure 3c). This is expected as the observed LFH layer is partly decomposed, and 305 will therefore have lost some C compared to the simulated litter pools which have not yet been affected by the decomposition processes. In addition, the modeled litter pools contain some low-quality (high C:N ratio) CWD, which is not included in the LFH samples.

The observed total C:N ratio ranges from 12-45 with a mean value of 28, while MIMICS+ and CLM have mean values of 23 and 11, respectively. The range of C:N values from the models are narrower than the observed, with MIMICS+ values 310 ranging from 12-38 and CLM varies only between 11-12. The large variability among the observations indicates the influence of local conditions on a subgrid-scale. The fact that MIMICS+ has a larger variability than CLM indicates that differences in soil quality are captured better with the improved modeling framework. Microbial competition for N and a higher fraction of directly plant-derived SOM are factors contributing to this difference between the modeled C:N ratios. Figure 3d shows the MIMICS+ simulated C:N ratios at three different depth layers. As expected, the top layer with more litter has the highest 315 ratio, while in the middle and lowest layers the ratios are significantly lower. For the CLM simulations the C:N ratio is constant around 11 for all three depth intervals. Since we do not have access to observed vertical N stocks, it was not possible to produce this plot for the observed sites.



**Figure 4.** Annual mean pool fractions as simulated by MIMICS+. (a) The fractions of total C stored in the main pool categories, soil organic matter (SOM), litter and microbes. The box upper and lower edges are the 75th and 25th percentiles, respectively. The whiskers extend from the box by 1.5 times the inter-quartile range, and N=50 sites. (b) The fraction of C in each pool within each main pool category. MYC covers both EcM and AM, as the AM contribution is so small that it would not be visible on its own.

### 3.2 Modelled C pools in MIMICS+

In this section we look more in detail into model properties of MIMICS+. The sensitivity analysis showed that total soil C change using perturbed parameters was mostly within  $\pm 10\%$  of the default values (Fig. B1). Modeled soil C was most sensitive to the fraction of structural litter going directly to protected SOM, as well as mycorrhiza-related parameters (maximum CUE and mining decay rate  $K_{MO}$ ). The sensitivity of total C to parameter values related to inorganic N was small.

With the current model parameterization, the SOM pools contain about 78 % of the total soil C (all nine pools, ref. Fig. 1), and 62 % of that is in the protected pools (SOMc and SOMp in Fig. 1). The litter pools contain most of the remaining C, while 1.2 % is microbial biomass (Fig. 4). The modeled percentage of microbes ranges from 0.3–2 %, and is in agreement with the 1–3 % microbial biomass C typically reported for soils (Frey, 2019). The microbial respiration (HR) shows a clear seasonal pattern, with a stronger summer peak and winter limitation with MIMICS+ than with CLM (Fig. B4). Figure 4b shows the relative magnitude of each pool within a pool category. Mainly due to the relatively high CWD contribution to the input, the structural pool is the largest litter C pool (18 % of total C, 85 % of total litter C), while metabolic litter consisting of leaf and fine root litter is accounting for ca. 3 % of the total C, and 15 % of total litter. The saprotrophic microbial biomass C is dominating over the mycorrhizal fungi biomass C, and the saprotrophic fungi dominate over saprotrophic bacteria (mean saprotrophic F:B biomass ratio of 2 and above 1 for all sites). This is largely a consequence of the parameter choices in the model, and are further discussed in Sect. 4.

For the focus region of this study (boreal sites in Norway), total C (TOTC) is strongly correlated with both MAT and C\_input (+0.49 and +0.65, respectively) indicating that higher plant productivity at warmer sites is an important control on total soil

C in the MIMICS+ simulations (Fig. 5). The CUE presented in Fig. 5 is calculated as the ratio of total microbial C uptake in biomass over the C total uptake (including respiration). CUE is positively correlated with available N, pointing to higher microbial efficiencies at sites with higher nutrient content. This is also illustrated by the positive relationship between the percentage of microbial biomass (pct\_microbes) and available inorganic N (+0.41 for  $N_{NO_3}$  and +0.62 for  $N_{NH_4, sol}$ ). The 340 negative correlation between CUE and MAT is likely explained by lower quality litter input at warmer sites, as there is a positive relationship between the C:N ratio of the litter input and temperature (+0.46 p < 0.001, not shown). The lower litter quality causes reduced CUE and hence a negative relationship between temperature and CUE. The strong correlation between 345 production (C\_input) and HR (+0.81) indicates that most sites are close to equilibrium. Lower litter quality at high production (and respiration) sites can explain the negative relationship between CUE and HR. There is a negative correlation (-0.64) between CUE and total C.

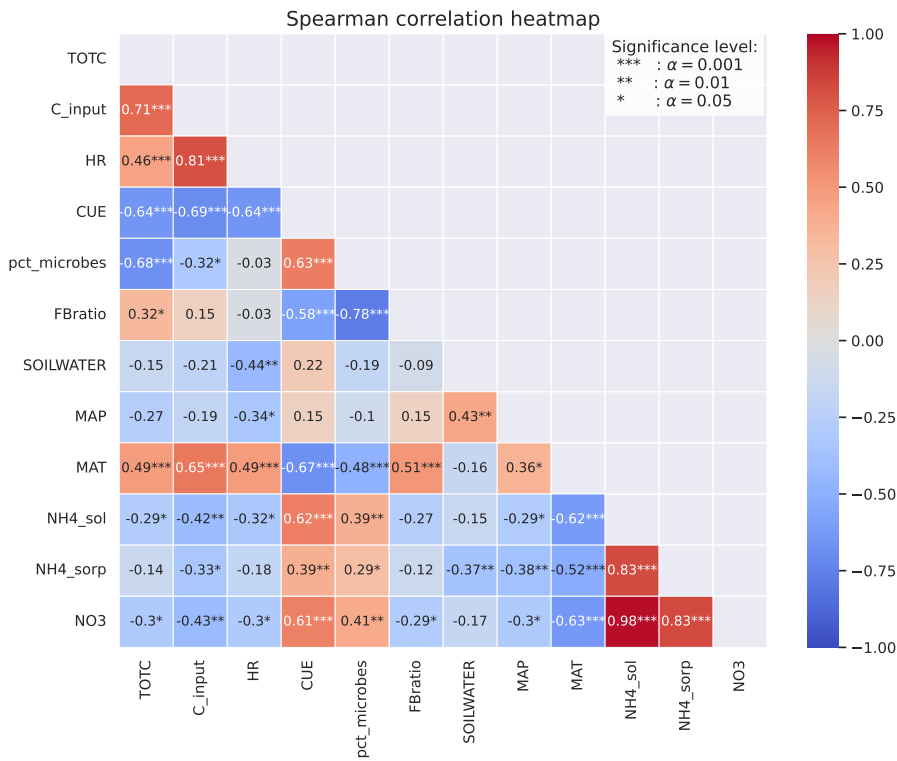
The fungal:bacterial saprotrophic biomass ratio (FBratio) is negatively correlated to available inorganic N (-0.29 for  $N_{NO_3}$  and -0.27 for  $N_{NH_4, sol}$ ), reflecting the stricter stoichiometrical constrain on bacteria. There is a strong negative correlation between the percentage of microbes and the fungal:bacterial ratio (-0.78), reflecting that sites with more available N are more favorable for microbial growth in both pools, but most beneficial for bacteria.

350 All three inorganic N pools are negatively correlated with MAP (-0.30 for  $N_{NO_3}$ , -0.29 for  $N_{NH_4, sol}$  and -0.38 for  $N_{NH_4, sorp}$ ),  $N_{NH_4, sorp}$  also with soil water (-0.37). This indicates that the modeled microbes also respond to moisture conditions through the effects of moisture on inorganic N processes (Leaching, runoff and sorption of  $NH_4$ ) which contribute to making N unavailable, and not only through the modifications of the reverse Michaelis-Menten kinetics.

### 3.3 Comparison of climate gradient profiles

355 In Fig. 6 the 50 sites have been divided into two subsets of 25 sites based on climate categories described in Sect. 2.3.1. (Fig. S1 shows the result of the division of sites based only on the on the observed climate.) Figures 6a–c show lower carbon stocks for colder than for warmer sites for both models and observations for all three depth intervals, indicating that the models are broadly able to capture the temperature-dependent processes that govern the C storage in soils. As shown in Fig. 5 the modeled 360 C input is positively correlated with MAT and total soil C, indicating that the difference is mainly caused by differences in litter input. The MIMICS+ simulations show a significant difference between the cold and warm mean C content (p < 0.05) for all depth intervals, while the cold and warm means from the CLM simulations are not significantly different (0.14 < p < 0.29). This indicates that MIMICS+ temperature dependencies have a larger impact on soil C sequestration than the standard CLM formulation since the C inputs and soil temperatures are the same for the two models.

Figures 6d–f show that in the observations, the drier sites have a lower mean C stock than the wetter sites (but not significant). 365 This is opposite of the modeled results; both models show higher mean C content for the drier sites than for the wetter sites. For MIMICS+ this discrepancy is only evident in the top layer, whereas for the lower layers, there are no significant differences between the drier and wetter sites. For the CLM simulations, the pattern is consistent and significant for all three depth intervals (p < 0.05). The influence of moisture on decomposition is represented differently in the two models, which can explain some of the difference between the modeled values. This is further discussed in Sect. 4.3.

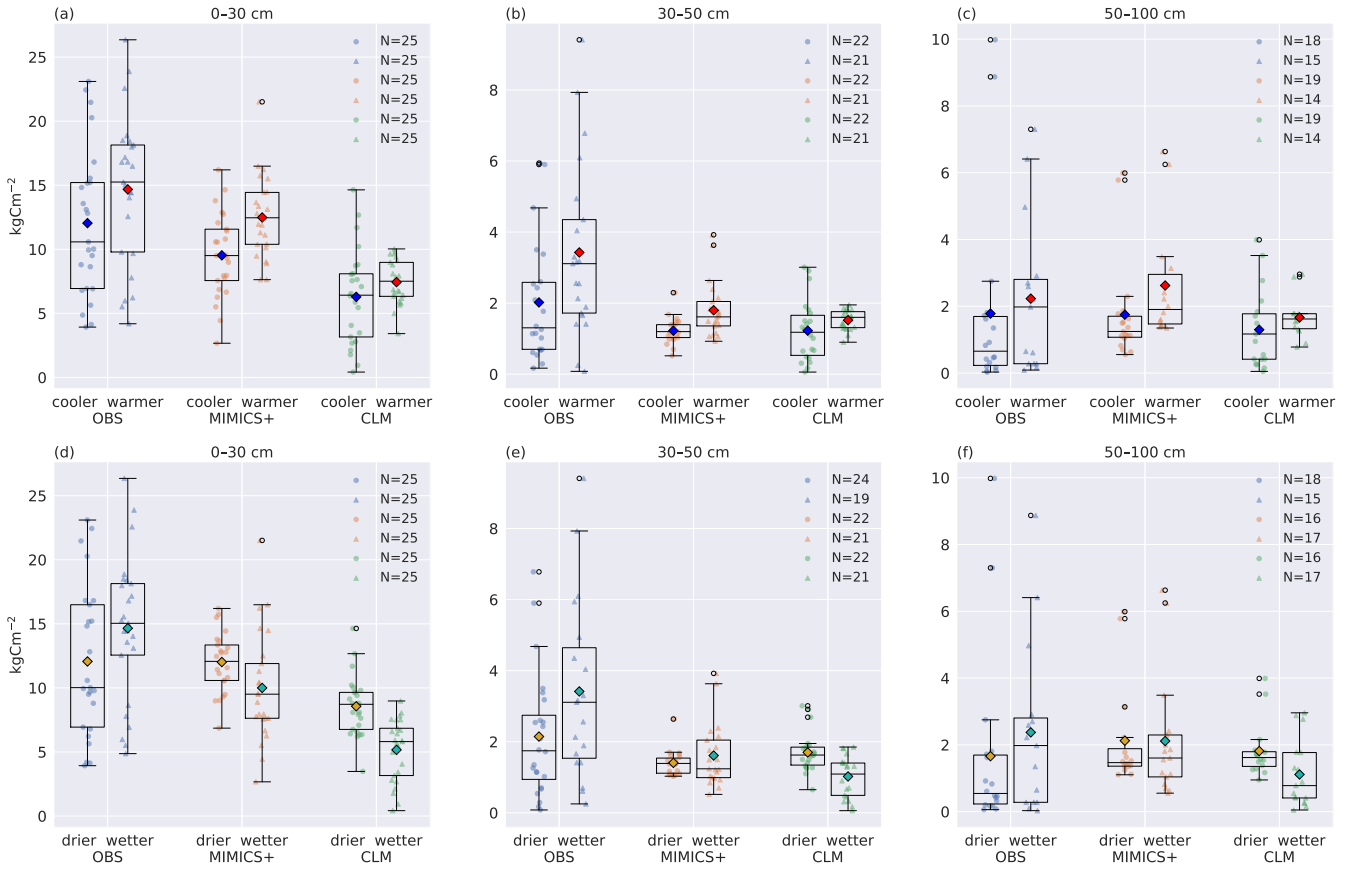


**Figure 5.** Spearman correlation coefficients between different variables calculated from MIMICS+ simulations of the 50 sites. The stars represent significance level of the correlation. Numbers without stars are not significant ( $p > 0.05$ ). The color indicate whether the correlation is positive (red) or negative (blue), and the shade indicate the strength of the correlation.

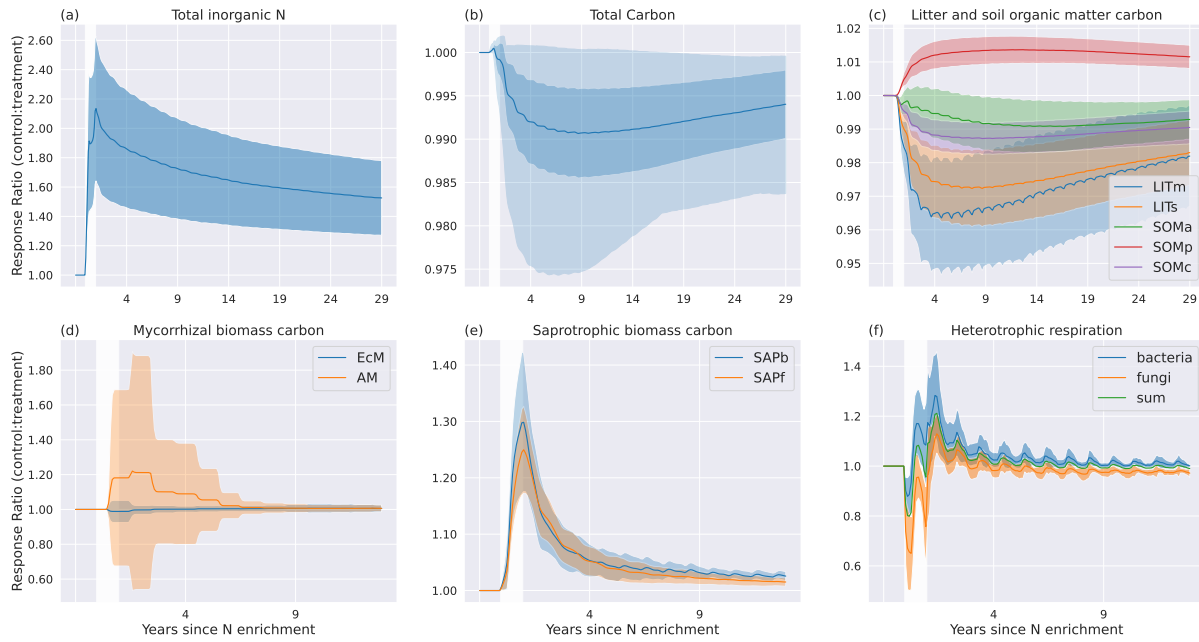
### 370 3.4 N enrichment experiment

The responses to the N enrichment experiment are a result of how the extra reactive N ( $15 \text{ gNm}^{-2}$  distributed evenly during one year) is distributed between the inorganic nitrogen pools after addition (Fig. 7a). All extra N is added to the  $N_{\text{NH4,sol}}$  pool, which had the largest response ratio of the three inorganic N pools. Some of this N will move gradually to  $N_{\text{NO3}}$  via nitrification or to  $N_{\text{NH4,sorp}}$  through sorption. While N is lost from  $N_{\text{NO3}}$  relatively fast via plant and microbial uptake and leaching, the extra sorbed N serves as a long-term supply of inorganic N, slowly releasing N back to the dissolved pool. This sustains the higher CUE of the microbes and leads to increased saprotrophic biomass for the duration of the 30-year simulation. 375 Although  $N_{\text{NH4,sol}}$  has the largest relative response to N addition, the change in mass of N is largest in the  $N_{\text{NH4,sorp}}$  pool.

Looking at each C pool response separately, we see the largest responses in the microbial pools (Fig. 7d–e). The extra inorganic N gives a relatively higher return of investment (ROI, Eq. (9)) for AM, resulting in more C allocated to AM and less 380 to EcM. The initial large response declines gradually but remains positive throughout the simulation period. Although there is a shift to more AM, the EcM carbon pool is always larger than the AM pool.



**Figure 6.** Total carbon stocks for cooler/warmer (a–c) and drier/wetter (d–f) parts of the dataset. Box plots of carbon stocks in the (a), (d) top 30 cm, (b), (e) 30–50 cm, (c), (f) 50–100 cm soil depths for observed profiles from Strand et al. (2016) (left), simulated with MIMICS+ (center) and with CLM (right). In (a–c) the leftmost quartiles represent the coldest 50 % of the dataset, while the rightmost represent the warmest 50 % of the dataset. In (d–f) the leftmost boxes represent the drier 50 % of the total subset, while the rightmost represent the wetter 50 %. The line in each box is the median, while the diamonds mark the mean values. The diamond color represent the climate category; yellow: drier, turquoise: wetter, blue: cooler, red: warmer. The box upper and lower edges are the 75th and 25th percentiles, respectively. The whiskers extend from the box by 1.5 times the inter-quartile range.



**Figure 7.** Temporal mean (N=50 sites) response ratios (treatment:control) to experimental N enrichment for (a)  $N_{NH_4, sol}$ , (b)  $N_{NH_4, sorp}$ , (c)  $N_{NO_3}$ , (d) mychorrizal fungi pools, (e) saprotrophic pools, (f) heterotrophic respiration, (g) total soil C (h) Litter pools, (i) soil organic matter pools,. The white area marks the year of N enrichment and the shading indicates the standard deviation. In (g) the lighter shading indicate the total spread of the values.

Both saprotrophic C pools respond instantly and positively to the N enrichment, with a maximum increase of about 25 % for fungi, and 30 % for bacteria at the end of the N addition year. The increase in saprotrophic biomass is a result of higher CUE made possible by more available N. After the N enrichment year, the response gradually decreases until it stabilizes at around 1 % after ca. 5 years. The long-term response is marginally higher for bacteria than for fungi.

The initial response in HR (Fig. 7f) is a result of a lower respiration fraction, (1-CUE), leading to increased saprotrophic biomass and thus gradually increased rates of litter decomposition. After the initial negative response in HR in the N enrichment year, there is a positive response due to the higher decomposition rate. For bacterial HR, the response ratio stabilizes at a low positive value, while for fungi it stabilizes at a slightly negative value. Combined, the response ratio is close to zero for HR after approximately four years.

The positive microbial biomass responses result in initial decreases in the substrate pools, LITm, LITs and SOMa (Fig. 7c). Most microbial necromass ends up in either the physically protected pool (SOMp) or the available pool (SOMa), leading to a positive response for SOMp, while the increased decomposition of SOMa keeps the response ratio below 1. The chemically protected pool experience a small negative response because increased microbial biomass increases the rate of the depolymerization process that moves chemically recalcitrant SOM to the available SOM pool (C11). The responses in SOM and litter

pools are weak, and following one year of N enrichment the mean response of total C is a marginal decrease compared to the control. It is worth noting that some sites experience markedly larger responses in total C than others (shading in Fig. 7b).

The overall response of the model illustrates that shifts in N availability have consequences for microbial C and N dynamics, although not necessarily for the total C storage and respiration. It should be noted that in this experiment we did not increase 400 plant productivity and thus carbon input to the soil, which is expected after N enrichment. This also means that possible changes in plant-microbe competition for N ~~was-were~~ not captured. The added value of this experiment is that we isolate the in-soil processes and quantify the effects of added nutrients available to microbes and how this affects the soil carbon pools.

## 4 Discussion

This study aimed to introduce a microbially explicit soil decomposition model, MIMICS+, designed to represent key soil 405 processes that control carbon and nitrogen processing in boreal ecosystems, but still be general enough to be used within an ESM. The model was applied to investigate responses to an N enrichment experiment. The results show that modeled mean C stocks with MIMICS+ matches observations reasonably well, and for Norwegian forested podzolic sites the model is performing on par with or better than the state-of-the-art land model CLM using a traditional decomposition formulation (Fig. 2a–c). However, ~~due to local heterogeneity not captured by the models~~, both models showed poor one-to-one agreement 410 with the observations (Fig. 2d–f), possibly due to local heterogeneity that is not captured by the models and/or inaccuracies in the model climate forcings. The C:N ratios from MIMICS+ are closer to observations than CLM, and the predicted fraction of microbial biomass matches well with values reported in the literature (Fig. 3 and Fig. 4). Several noteworthy correlations between variables were found from the MIMICS+ simulations (Fig. 5). Both models capture the climatic temperature pattern from the observed soil profiles, although they both struggle to represent the observed pattern in C concentrations emerging from 415 the MAP categories (Fig. 6). The N enrichment experiment demonstrates the implications of adding the Langmuir algorithm for inorganic N, as the sorbed NH<sub>4</sub> works as a long-term supply of N for the microbes. The overall effect of the idealized enrichment experiment on soil C storage and respiration was minor but had interesting effects on the relative distribution of the microbial groups, and shows the need for further investigation into the role of sorption-desorption processes of inorganic N, especially in N-limited areas like boreal forests (Fig. 7).

### 420 4.1 Comparison of modelled and empirical C and N stocks

Looking at the total distribution for the 50 sites, MIMICS+ is closer than CLM to the observations for the top layer (0–30 cm), models are similar in the middle layer (30–50 cm), while none of the modeled means are significantly different from the observations in the bottom layer (50–100 cm). The site-to-site comparisons with observations were poor for both models, showing that there is a discrepancy between observed and modeled stocks at local scales. This challenge of local factors 425 was illustrated by Pierson et al. (2022) who used the C-only version of MIMICS with optimized parameters based on local observations and showed reduced error of C stocks on smaller scales (catchments < 50 km<sup>2</sup>). Such methods would likely also reduce the errors of MIMICS+ at smaller scales. However, it is important to keep in mind that the intention with MIMICS+

is to develop a module that is simple and fast enough to be used in an ESM to simulate the soil carbon dynamics at a grid cell average scale. When forced with grid cell average input, it is not intended to and should not be expected to accurately 430 describe local variation in soil carbon stocks. Upscaling of point observations of soil C stocks to landscape level in our study area (Norwegian boreal forests) would be useful for comparison of ESM simulations with empirically-based estimates of soil C stocks.

With the MIMICS-CN version Kyker-Snowman et al. (2020) obtained soil C:N ratios that, although within observed ranges, had much lower maximums than the observed ratios. They suggested increasing the fraction of litter going directly to SOM, 435 as forest soils (compared to agricultural and grassland soils) have been shown to contain a high fraction of C in plant residues (Grandy and Robertson, 2007). Our focus area is forested ecosystems, so we increased the fraction of litter going directly to protected SOM without going through microbial decomposition to 50 % for both structural and metabolic litter (these fractions also affect the total C, see Fig. B1). This leads to a longer lifetime of soil C (stored in protected pools) before it becomes available for microbial decomposition and respiration. The higher directly plant-derived fraction in the SOM pools increases 440 the soil C:N ratio, although it is still lower than observed for total and mineral soil (Fig. 3). A recent study by Angst et al. (2021) indicates that the fraction of directly plant-derived SOM may be much higher than previously assumed, especially for forested sites and podzols. The high C:N ratios in our observational dataset point in the same direction, suggesting that the direct plant-derived fraction is an important factor to consider when modeling boreal soils. Our results demonstrate that we get closer to observed C:N ratios with MIMICS+ compared to the CLM formulation, a main reason being the high direct plant- 445 derived fraction. In the CENTURY-based decomposition cascade in CLM, the C:N ratios of the SOM pools are fixed, which gives limited options to account for high C:N ratios, and the implications that may have on soil C dynamics.

## 4.2 Modelled C pools in MIMICS+

The division of C between the different pools in MIMICS+ shows that most soil C is in the SOM pools (78 %), whereof 62 % is protected. This again reflects the relatively high fraction of litter going directly to protected SOM, but also the lifetime of C 450 in the protected pools before it is either depolymerized or desorbed into the available SOM pool. Compared to MIMICS-CN we doubled the desorption coefficient (see Table A5), but this is still one order of magnitude lower than the value used in the C-only version of MIMICS (Wieder et al., 2015). In the above-mentioned studies and the present study, this parameter has been adjusted to match the observed data. In the model formulation, the desorption coefficient is a function of soil clay content, and more observational studies constraining this parameter as a function of clay content and/or other observable variables would 455 benefit further model development.

Saprotrophic fungi are the dominant microbial group in our simulations. Fungi are assumed to have a higher maximum CUE than bacteria in the model (0.7 vs. 0.4, respectively), and are more efficient at decomposing structural litter than the bacterial pool (higher  $V_{max}$  for decomposition of LITs by SAPf than SAPb). This is based on the assumption that fungal decomposers are more specialized towards recalcitrant substrates, while bacteria thrive on labile, easy-access metabolic litter (Wardle et al., 460 2004). The fraction of CWD litter provided from CLM is relatively large at these forested sites, giving more substrate that is preferable for fungi. The Norwegian podzols we are looking at are nutrient poor, and fungal dominance is expected under

N-limited conditions (Strickland and Rousk, 2010). Figure 5 indicate a negative relationship between available inorganic N and F:B ratio, meaning a higher fraction of bacteria in more nutrient rich conditions, in line with observations. Further, the N enrichment experiment showed that bacteria had a larger positive response to the added N in the long term, which indicates  
465 that the model can capture shifts in microbial communities in response to N conditions.

The modeled saprotrophic fungal biomass C dominates over the mycorrhizal fungi biomass C. This contrasts an observational study on boreal forests that indicate that EcM can account for as much as 47–84 % of fungal biomass (Bååth et al., 2004). Also, Clemmensen et al. (2013) challenged the traditional view that C sequestration is mainly driven by the decomposition of above-ground litter by saprotrophs with their study that showed a dominance of root-associated fungi in deeper parts  
470 of the LFH in boreal forests. Few studies exist to inform models about fungal dominance in boreal systems, so parameters determining mycorrhizal growth and turnover are poorly constrained, and not particularly adjusted for boreal conditions in this model iteration. The sensitivity analysis showed that particularly the EcM mining rate ( $K_{MO}$ ) and maximum mycorrhizal CUE impacts total modeled C (Fig. B1), highlighting the need to inform these parameters with representative observations. The C supply to mycorrhizal pools is prescribed directly from CLM output, and the growth of these pools is therefore governed by  
475 this input rate. Coupling MIMICS+ to the above-ground vegetation will allow the plant C supply to react to nutrient conditions in the soil, and is a priority in future model development.

Regarding the correlations presented in this study (Fig. 5), one should always keep in mind that correlation does not imply causation, especially in a coupled non-linear system like this model. The analysis should be regarded as a broad investigation into possible relationships within the soil dynamics. Recently, Tao et al. (2023) presented CUE as a strong predictor of SOC  
480 globally, and argued for a positive correlation between CUE and soil carbon storage (SOC) based on a combination of global-scale datasets, a microbial-process explicit model, data assimilation, deep learning and meta-analysis. In contrast, our analysis showed a negative correlation between microbial CUE and soil carbon storage, in addition to a strong correlation between total carbon and plant litter input. A relatively large fraction of the litter input in MIMICS+ (50 %) initially omits the microbial pathway (affected by CUE) as directly plant-derived organic matter into protected SOM pools, which weakens the relationship  
485 between microbial CUE and TOTC. A high fraction of microbial necromass ends up in SOMa (Eq. C13-C18 in Table A3 and parameters in Table A5). This leads to a relatively rapid recycling of the C that initially goes through the microbial pathway, which can also contribute to a weaker relationship between CUE and C storage than if larger fractions of the necromass ended up in the protected SOM pools. However, more microbially derived mass in the protected SOM pools will decrease the C:N ratio, bringing modeled values further away from the observed C:N ratios in Strand et al. (2016). Tao et al. (2023) used a  
490 process-guided deep learning and data-driven modeling (PRODA) approach to optimize parameters in a microbially explicit model (Allison et al., 2010) using observations. Default model parameters prior to optimization gave a negative relationship between CUE and SOC, illustrating how model estimates rely on parameter choices. Using a similar approach to inform MIMICS+ can lead to more robust parameter values in future model iterations.

In MIMICS+ the availability of inorganic N is highly dependent on soil water processes because both N leaching from  
495  $\text{NO}_3^-$  and the Langmuir isotherm algorithm is dependent on soil moisture. This is evident from Fig. 5, where we see a negative correlation between inorganic N pools and moisture-related variables (MAP and SOILWATER). The available inorganic N

500 pools are again positively correlated with the percentage of microbes, giving an indirect dependence of microbes on soil moisture. The total C is negatively correlated with the percentage of microbes, and has a high correlation with the incoming C. With higher temperatures, the model gives a higher turnover rate and thereby more release of soil C to the atmosphere. However, increased temperatures also stimulate plant production, especially in boreal and Arctic regions, which can exceed or offset the effect of higher decomposition rates (Hobbie et al., 2002). The correlation patterns from our simulation indicate that the effect of temperature on plant production dominates the effect of temperature on decomposition rates in the model. Pierson et al. (2022) found that increased temperature sensitivity of the decomposition kinetics compared to the original MIMICS/MIMICS+ parameter values reduced error compared to their observational data, indicating that the temperature sensitivity in MIMICS and MIMICS+ may be too weak. However, the agreement between models and observations in Fig. 6a–c indicates that more plant production is the dominating effect of higher temperatures in Norwegian forests.

### 4.3 Comparison of climate gradient profiles

510 Although simple, dividing sites into different climatic categories serves as an idealized "space-for-time" investigation of climate change responses. Assuming that the climate in boreal forests in general, and Norwegian forests specifically will get warmer and wetter in the future (Hanssen-Bauer et al., 2017), the observations indicate higher soil C content at sites with higher MAP and MAT. The models indicate higher C content for warmer sites, but lower C content for the wetter sites, especially in the 0–30 cm layer. There is a positive correlation between MAT and MAP, particularly for the observed climate forcing (Fig. B3). When dividing the observed sites into the climate categories, a large fraction end up as either "cold and dry" or "warm and wet". We therefore did a simple "temperature-dependence removal" on the total podzol dataset (N=578) by dividing the sites 515 into narrow temperature intervals of 0.5 °C (Fig. S2). This did not reveal a clear pattern between the wetter and drier sites, and it is therefore difficult to disentangle the effects of moisture from the effects of temperature in the observed data. Since the models use soil moisture, not MAP to define parameters, we also analyzed the results using a soil moisture variable from the CLM simulations (*SOILWATER\_10CM*) instead of MAP to discriminate between "drier" and "wetter" sites to investigate any effects on the climatic pattern (Fig. S3). This showed the same pattern as in Fig. 6d-f (more C in drier soils for the models, 520 and less C in drier soils for observations) for all three distributions, except for the deepest layer, where the trend shifted for the observations, but not significantly. The CLM simulations show a negative correlation between MAP and total C (-0.63,  $p<0.001$ , Fig. B2), while this is not evident for MIMICS+, indicating that it is different factors that determine the pattern from the two models. In MIMICS+, the moisture modifier on decomposition works on the fluxes from substrate to the microbial pools. The modeled microbes are most abundant in the top 0–30 cm, which can explain why we observe a difference between 525 drier and wetter sites only in this layer. In CLM, the moisture modification on decomposition rates works on every step in the decomposition cascade from litter to SOM pools. Since the SOM pools have more C in deeper layers, it can explain why we see the pattern in all three depth intervals for the CLM simulations. The moisture modifier used in MIMICS+ (see  $r_{moist}$ , Table A5, and Wieder et al. (2017); Sulman et al. (2014)) is a bell-shaped function of soil moisture, limiting decomposition both in the case of very wet and very dry soil conditions. If the optimal soil moisture conditions according to this function are not 530 representing the optimal soil moisture value of the real soils, this could explain why MIMICS+ predicts the opposite pattern

between the drier and wetter soils. Also, soil moisture can vary with subgrid features like slope and aspect, variations not expected to be captured by CLM. Therefore, discrepancy between actual and modeled soil moisture can also be a contributing factor.

#### 4.4 N enrichment experiment

535 Meta-analyses of observational N enrichment studies show that microbial biomass tends to decrease after enrichment, while the response in total soil C is relatively modest (Treseder, 2008; Janssens et al., 2010). The small modeled response of total soil C to N enrichment (Fig. 7b) is in line with these observations, but the modeled microbial biomass showed a marginal long-term increase after an initial high response (Fig. 7d-e). Treseder (2008) proposed several mechanisms for N effects on microbial growth (Fig. 1 in her study), some leading to an increase while others leading to a decrease in microbial biomass.

540 The sites studied in our model simulations are mainly N-limited (N immobilization via mechanism 1 in Sect. 2.1.2 dominates), and we see an accumulation of microbial biomass as a direct consequence of the increased N availability, which is one of the mechanisms suggested by Treseder (2008) for an increase in microbial biomass. Mechanisms proposed to reduce microbial biomass in response to N enrichment are decreases in soil pH, decrease in ligninase activity, increase in melanoidins and a decrease in below-ground NPP. In this offline iteration of MIMICS+ we are unable to capture potential decreases in below-545 ground NPP allocation. Coupling to a vegetation model will enable this possibility, and might affect the modeled response N enrichment. When dividing results into separate biomes, Treseder's analyses indicate that for boreal forests the response for bacteria is positive (RR = 1.061), while for fungi negative (RR = 0.717) but with a confidence interval covering both positive and negative responses (0.0402–1.434). This points to uncertainties also in observations of responses of N enrichment. To cover more of the possible mechanisms for microbial biomass decline in the model, one or more of the other mechanisms mentioned 550 above could be included.

The strong N-limitation in the model is partly a consequence of using low, constant C:N requirements for the saprotrophic pools ( $CN_b = 5$  and  $CN_f = 8$  ref. Table A5). A less strict C:N requirement, or a dynamic C:N approach, as presented in the ORCHIMIC model (Huang et al., 2018, 2021), could lead to a weaker modeled N-limitation and/or more microbial N mineralization, which can affect the response to N enrichment. This could also improve the modeled underestimation of soil 555 C:N ratio, as N in inorganic forms are subject to loss through direct plant uptake and leaching.

In the simulations, the largest loss of soil N is through the ectomycorrhizal pathway (N29). The parameter sensitivity analysis also show a stronger sensitivity of total C to mycorrhizal parameters than to the plant uptake (Fig. B1). The high microbial immobilization of N, together with the simplified representations of direct plant uptake (constant loss rate of available inorganic N) might cause an overestimation of the loss of organic N through mycorrhiza at the expense of direct plant uptake of inorganic 560 N (N33). To model a more realistic scenario, with increased plant production and changes in plant N acquisition strategies as a response to the extra N, it is therefore necessary to couple MIMICS+ to a vegetation model. In such an experiment, both the increase in litter production and shifts in plant C allocation will affect the soil dynamics. The enrichment experiment presented in this study showed that the model is able to capture microbial responses, and in a coupled system it can be used to further study plant-microbe competition for nutrients.

565 **4.5 Limitations and Future Improvements of the MIMICS+ framework**

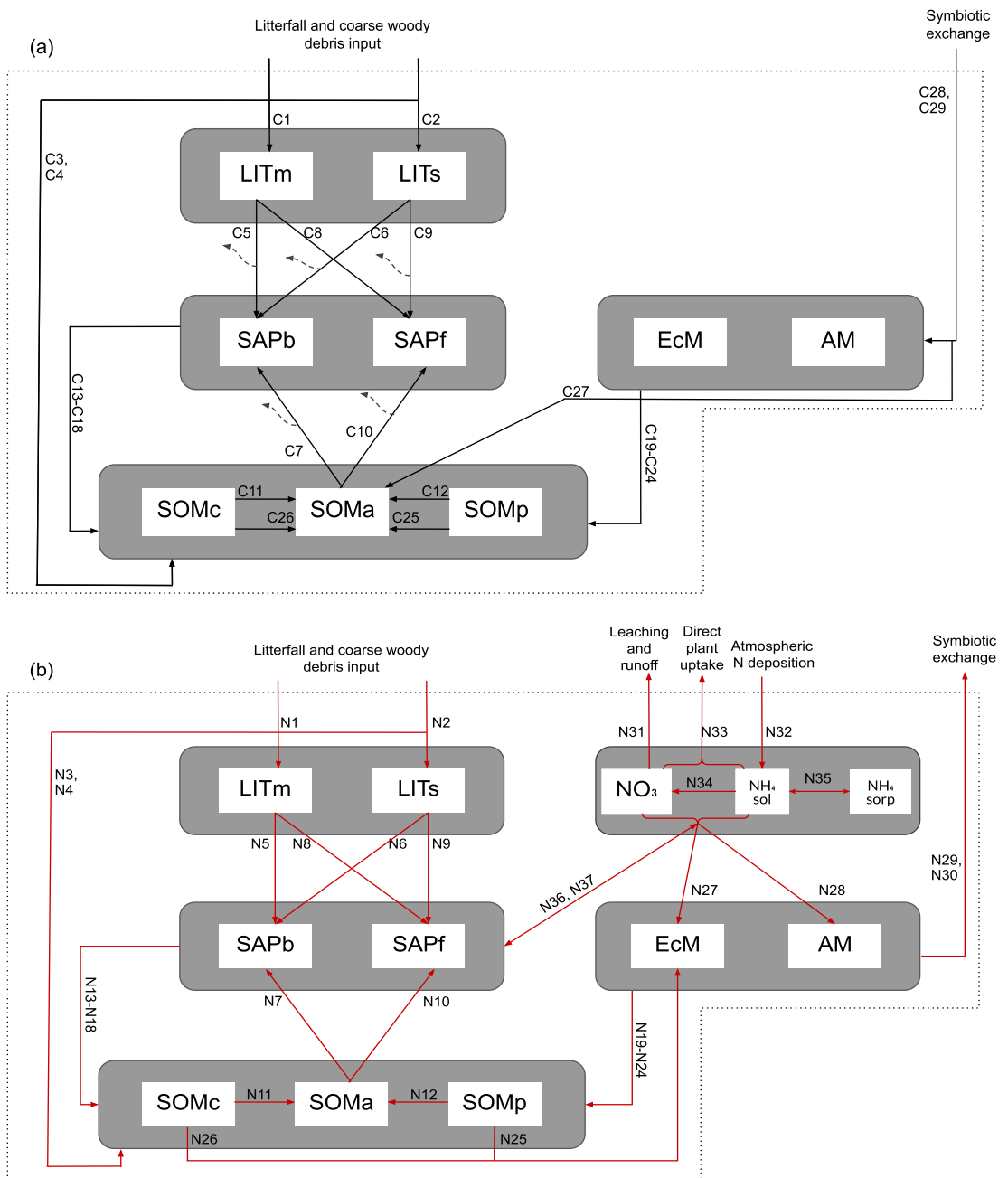
By expanding the MIMICS framework with extra microbial groups and an N cycle, we increase possibilities to capture microbe-microbe interactions and, after coupling also plant-microbe interactions. However, we also introduce additional parameters and a more complex model structure that makes the model more prone to overfitting and equifinality issues. While acknowledging this possible drawback, we believe valuable insights can be gathered through a more detailed process representation, especially  
570 as new technologies allow measurements suitable to constrain model parameters. Although the model produces results comparable to the observations from Strand et al. (2016), there are still poorly constrained parameters in the model, especially related to mycorrhizal C and N transfer. Recent insights about the mycorrhizal role in soil C dynamics are valuable contributions for future model development (Huang et al., 2022a, b). A more robust parameter optimization procedure like the PRODA approach  
575 (Tao et al., 2023) or a Monte Carlo approach (Pierson et al., 2022) will contribute to constraining model parameters. The model should also be evaluated against observations from other ecosystems, which will increase confidence in model structure and parameter choices. This offline version of MIMICS+ does not capture plant-microbe interactions and feedbacks, which is essential to capture terrestrial responses to climate change. Therefore, coupling with a vegetation model is a priority in future model development.

## 5 Conclusions

580 The soil model MIMICS+ provides a tool for investigating soil C processes and interactions with the N cycle, particularly relevant for boreal areas. Furthermore, the model framework will serve as a valuable soil module in ESMs as it is general enough to work on larger scales. The model produces soil C and N stocks comparable to observed values in Norwegian forest podzols. The explicit representation of microbial groups enhances performance compared to the traditional CLM, and enables detection of soil dynamics not possible with a conventional model. In particular, the novel representation of sorbed inorganic  
585 N can be further developed to examine climate responses in N-limited systems like boreal forests, but also in possible impacts on other ecosystems not limited by N. In this study the MIMICS+ model is decoupled from vegetation, so we cannot directly detect feedbacks between nutrient availability and plant productivity. Coupling MIMICS+ to a dynamical vegetation model like FATES will further enable investigation of the interplay between soil microbes and changing above-ground vegetation.

*Code availability.* MIMICS+ (v1.0) is written in Fortran90. Figures and analyses was produced with Python and Jupyter notebook. The  
590 model code and Jupyter notebook is available online at <https://doi.org/10.5281/zenodo.10610814>.

## Appendix A: Model Description Details



**Figure A1.** Illustration of all carbon (a) and nitrogen (b) pools and fluxes in the system. Expressions for each flux is found with the corresponding numbers in Table A3 and A4.

**Table A1.** Mass balance equations for the carbon pools in the model, calculated for each vertical layer (subscript dropped for readability).  $FC'_{donor,receiver}$ :  $\text{gCm}^{-3}\text{h}^{-1}$ . Details about the fluxes are found in Table A3.

Eqn	Stores	Growth rates	Fluxes
(a)	Metabolic litter	$dC_{LITm}/dt =$	$FC_{Veg,LITm} - FC_{LITm,SAPb} - FC_{LITm,SAPf}$
(b)	Structural litter	$dC_{LITs}/dt =$	$FC_{Veg,LITs} - FC_{LITs,SAPb} - FC_{LITs,SAPf}$
(c)	Saprotrophic bacteria	$dC_{SAPb}/dt =$	$CUE_b \cdot FC_{uptake,SAPb}^a - FC_{SAPb,SOMp} - FC_{SAPb,SOMa} - FC_{SAPb,SOMc}$
(d)	Saprotrophic fungi	$dC_{SAPf}/dt =$	$CUE_f \cdot FC_{uptake,SAPb}^a - FC_{SAPf,SOMp} - FC_{SAPf,SOMa} - FC_{SAPf,SOMc}$
(e)	Ecotomycorrhiza	$dC_{EcM}/dt =$	$CUE_{EcM} \cdot FC_{Veg,EcM} - FC_{EcM,SOMp} - FC_{EcM,SOMa}$
			$-FC_{EcM,SOMc} - FC_{enzEcM,SOMa}$
(f)	Arbuscular mycorrhiza	$dC_{AM}/dt =$	$CUE_{AM} \cdot FC_{Veg,AM} - FC_{AM,SOMp} - FC_{AM,SOMa} - FC_{AM,SOMc}$
(g)	Phys. protected SOM	$dC_{SOMp}/dt =$	$FC_{Veg,SOMp} + FC_{SAPb,SOMp} + FC_{SAPf,SOMp}$
			$+FC_{EcM,SOMp} + FC_{AM,SOMp} - FC_{SOMp,SOMa} - FC_{EcMdecompSOMp}$
(h)	Chem. protected SOM	$dC_{SOMc}/dt =$	$FC_{Veg,SOMc} + FC_{SAPb,SOMc} + FC_{SAPf,SOMc}$
			$+FC_{EcM,SOMc} + FC_{AM,SOMc} - FC_{SOMc,SOMa} - FC_{EcMdecompSOMc}$
(i)	SOM available	$dC_{SOMa}/dt =$	$FC_{SAPb,SOMa} + FC_{SAPf,SOMa} + FC_{EcM,SOMa} + FC_{AM,SOMa}$
			$+FC_{SOMp,SOMa} + FC_{SOMc,SOMa} + FC_{enzEcM,SOMa}$
			$+FC_{EcMdecompSOMc} + FC_{EcMdecompSOMp} - FC_{SOMa,SAPb} - FC_{SOMa,SAPf}$
			$+FC_{Veg,LITn} + FC_{Veg,LITs} + FC_{Veg,SOMp} + FC_{Veg,SOMc}$
			$+CUE_{EcM} \cdot FC_{Veg,EcM} + CUE_{AM} \cdot FC_{Veg,AM} - (1 - CUE_b) \cdot FC_{uptake,SAPb}$
			$-(1 - CUE_f) \cdot FC_{uptake,SAPf}^a$
Net Carbon change:		$dC'/dt =$	$FC_{Cuptake,s} = FC_{LITm,s} + FC_{LITs,s} + FC_{SOMa,s}$

<sup>a</sup>  $FC_{Cuptake,s} = FC_{LITm,s} + FC_{LITs,s} + FC_{SOMa,s}$

**Table A2.** Mass balance equations for the nitrogen pools in the model, calculated for each vertical layer (subscript dropped for readability).  $FN_{donor,receiver}$ :  $\text{gNm}^{-3}\text{h}^{-1}$ . Details about the fluxes are found in Table A4.

Eqn	Stores	Growth rates	Fluxes
(j)	Metabolic litter	$dN_{LITm}/dt =$	$FN_{Vegetation,LITm} - FN_{LITm,SAPb} - FN_{LITm,SAPf}$
(k)	Structural litter	$dN_{LITs}/dt =$	$FN_{Vegetation,LITs} - FN_{LITs,SAPb} - FN_{LITs,SAPf}$
(l)	Saprotrophic bacteria	$dN_{SAPb}/dt =$	$NUE \cdot FN_{uptake,SAPb}^a - FN_{SAPb,SOMp} - FN_{SAPb,SOMa} - FN_{SAPb,SOMc} + FN_{IN,SAPb}^c$
(m)	Saprotrophic fungi	$dN_{SAPf}/dt =$	$NUE \cdot FN_{uptake,SAPf}^a - FN_{SAPf,SOMp} - FN_{SAPf,SOMa} - FN_{SAPf,SOMc} + FN_{IN,SAPf}^c$
(n)	Ectomycorrhiza	$dN_{EcM}/dt =$	$FN_{IN,EcM}^b + FN_{SOMp,EcM} + FN_{SOMc,EcM}$ $- FN_{EcM,SOMp} - FN_{EcM,SOMa} - FN_{EcM,SOMc} - FN_{EcM,Veg}$
(o)	Arbuscular mycorrhiza	$dN_{AM}/dt =$	$FN_{IN,AM}^b - FN_{AM,SOMp} - FN_{AM,SOMa} - FN_{AM,SOMc} - FN_{AM,Veg}$
(p)	Phys. protected SOM	$dN_{SOMp}/dt =$	$FN_{SAPb,SOMp} + FN_{SAPf,SOMp} + FN_{EcM,SOMp} + FN_{AM,SOMp} + FN_{Veg,SOMp}$
(q)	Chem. protected SOM	$dN_{SOMc}/dt =$	$- FN_{SOMp,SOMa} - FN_{SOMp,EcM}$ $FN_{SAPb,SOMc} + FN_{SAPf,SOMc} + FN_{EcM,SOMc} + FN_{AM,SOMc}$ $+ FN_{Veg,SOMp} - FN_{SOMc,SOMa} - FN_{SOMc,EcM}$
(r)	SOM available	$dN_{SOMa}/dt =$	$FN_{SAPb,SOMa} + FN_{SAPf,SOMa} + FN_{SOMp,SOMa} + FN_{SOMc,SOMa}$ $+ FN_{EcM,SOMa} + FN_{AM,SOMa} - FN_{SOMa,SAPb} - FN_{SOMa,SAPf}$
(s)	Ammonium, solved	$dN_{NH4,sol}/dt =$	$FN_{DEP} + (1 - NUE)(FN_{uptake,SAPb} + FN_{uptake,SAPf})^a -$ $f_{NH4}(FN_{IN,EcM} + FN_{IN,AM} + FN_{IN,Veg}) - f_{NH4}(FN_{IN,SAPb} + FN_{IN,SAPf})$ $- FN_{NH4,NO3} + FN_{sol,sorp}$
(t)	Ammonium, sorbed	$dN_{NH4,sorp}/dt =$	$- FN_{sol,sorp}$
(u)	Nitrate	$dN_{NO3}/dt =$	$FN_{NH4,NO3} - FN_{run+leach} -$ $(1 - f_{NH4})(FN_{IN,EcM} + FN_{IN,AM} + FN_{IN,Veg}) - (1 - f_{NH4})(FN_{IN,SAPb} + FN_{IN,SAPf})$ $- FN_{IN,Veg} - FN_{EcM,Veg} - FN_{AM,Veg}$
Net Nitrogen change:		$dN/dt =$	

<sup>a</sup>  $FN_{uptake,s} = FN_{LITm,s} + FN_{LITs,s} + FN_{SOMa,s}$

<sup>b</sup>  $FN_{IN,receiver} = FN_{NO3+NH4sol,receiver}$

<sup>c</sup> Can be either positive or negative.

**Table A3.** Details about C fluxes in the model. The eq. numbers corresponds to the arrows in Fig. A1a. The letters in the fifth column matches with those given in Table A1. All  $FC_{donor,receiver}$  has units  $\text{gCm}^{-3}\text{h}^{-1}$ . Parameters are described in Table A5.

Eq	Flux Name	Rate Functions	Used in Eqn.	Notes
C1	$FC_{Veg,LITm}$	$f_{met} \cdot I_C \cdot (1 - f_{met,SOM})$	(a)	$I_C$ include litterfall + mortality rates
C2	$FC_{Veg,LITs}$	$((1 - f_{met}) \cdot I_C + CWD_C) \cdot (1 - f_{struct,SOM})$	(b)	
C3	$FC_{Veg,SOMp}$	$f_{met} \cdot I_C \cdot f_{met,SOM}$	(g)	
C4	$FC_{Veg,SOMc}$	$((1 - f_{met}) \cdot I_C + CWD_C) \cdot f_{struct,SOM}$	(h)	
C5	$FC_{LITm,SAPb}$	$C_{SAPb} \cdot V_{max1} \frac{C_{LITm}}{K_{m1} + C_{SAPb}}$	(a)(c)	Reverse MMK
C6	$FC_{LITs,SAPb}$	$C_{SAPb} \cdot V_{max2} \frac{C_{LITs}}{K_{m2} + C_{SAPb}}$	(b)(c)	Reverse MMK
C7	$FC_{SOMa,SAPb}$	$C_{SAPb} \cdot V_{max3} \frac{C_{SOMa}}{K_{m3} + C_{SAPb}}$	(i)(c)	Reverse MMK
C8	$FC_{LITm,SAPf}$	$C_{SAPf} \cdot V_{max4} \frac{C_{LITm}}{K_{m4} + C_{SAPf}}$	(a)(d)	Reverse MMK
C9	$FC_{LITs,SAPf}$	$C_{SAPf} \cdot V_{max5} \frac{C_{LITs}}{K_{m5} + C_{SAPf}}$	(b)(d)	Reverse MMK
C10	$FC_{SOMa,SAPf}$	$C_{SAPf} \cdot V_{max6} \frac{C_{SOMa}}{K_{m6} + C_{SAPf}}$	(i)(d)	Reverse MMK
C11	$FC_{SOMc,SOMa}$	$\frac{C_{SAPf} \cdot V_{max2} \cdot C_{SOMc}}{K_O \cdot K_{m2} + C_{SAPb}} + \frac{C_{SAPb} \cdot V_{max5} \cdot C_{SOMc}}{K_O \cdot K_{m5} + C_{SAPf}}$	(h)(i)	As in MIMICS
C12	$FC_{SOMp,SOMa}$	$C_{SOMp} \cdot k_{desorp}$	(g)(i)	As in MIMICS
C13	$FC_{SAPb,SOMp}$	$C_{SAPb} \cdot k_{SAPb,som} \cdot f_{SAPb,SOMp}$	(c)(g)	
C14	$FC_{SAPb,SOMc}$	$C_{SAPb} \cdot k_{SAPb,som} \cdot f_{SAPb,SOMc}$	(c)(h)	
C15	$FC_{SAPb,SOMa}$	$C_{SAPb} \cdot k_{SAPb,som} \cdot f_{SAPb,SOMa}$	(c)(i)	
C16	$FC_{SAPf,SOMp}$	$C_{SAPf} \cdot k_{SAPf,som} \cdot f_{SAPf,SOMp}$	(d)(g)	
C17	$FC_{SAPf,SOMc}$	$C_{SAPf} \cdot k_{SAPf,som} \cdot f_{SAPf,SOMc}$	(d)(h)	
C18	$FC_{SAPf,SOMa}$	$C_{SAPf} \cdot k_{SAPf,som} \cdot f_{SAPf,SOMa}$	(d)(i)	
C19	$FC_{EcM,SOMp}$	$C_{EcM} \cdot k_{myc,som} \cdot f_{EcM,SOMp}$	(e)(g)	
C20	$FC_{EcM,SOMc}$	$C_{EcM} \cdot k_{myc,som} \cdot f_{EcM,SOMc}$	(e)(h)	
C21	$FC_{EcM,SOMa}$	$C_{EcM} \cdot k_{myc,som} \cdot f_{EcM,SOMa}$	(e)(i)	
C22	$FC_{AM,SOMp}$	$C_{AM} \cdot k_{myc,som} \cdot f_{AM,SOMp}$	(f)(g)	
C23	$FC_{AM,SOMc}$	$C_{AM} \cdot k_{myc,som} \cdot f_{AM,SOMc}$	(f)(h)	
C24	$FC_{AM,SOMa}$	$C_{AM} \cdot k_{myc,som} \cdot f_{AM,SOMa}$	(f)(i)	
C25	$FC_{EcMdecSOMp}$	$K_{MO} \cdot dz \cdot C_{EcM} \cdot C_{SOMp} \cdot r_{myc}$	(g)(i)	(Baskaran et al., 2017) + mod. term
C26	$FC_{EcMdecSOMc}$	$K_{MO} \cdot dz \cdot C_{EcM} \cdot C_{SOMc} \cdot r_{myc}$	(h)(i)	(Baskaran et al., 2017) + mod. term
C27	$FC_{enzEcM,SOMa}$	$f_{enz} \cdot CUE_{EcM} \cdot FC_{Veg,Ecm}$	(e)(i)	
C28	$FC_{Veg,Ecm}$	$f_{alloc,Ecm} \cdot I_{veg,Myc}$	(e)	
C29	$FC_{Veg,AM}$	$f_{alloc,AM} \cdot I_{veg,Myc}$	(f)	

**Table A4.** Details about N fluxes in the model. The eq. numbers corresponds to the arrows in Fig. A1b. The letters in the fifth column matches with those given in Table A2. Parameters are described in Table A5.

Eq.	Flux Name	Rate functions	Used in eqn	Notes
N1	$FN_{Veg,LITm} =$	$f_{met} \cdot I_N \cdot (1 - f_{met,SOM})$	(j)	$I_N$ include litterfall + mortality rates
N2	$FN_{Veg,LITs} =$	$((1 - f_{met}) \cdot I_N + CWD_N) \cdot (1 - f_{struct,SOM})$	(k)	
N3	$FN_{Veg,SOMp} =$	$f_{met} \cdot I_C \cdot f_{met,SOM}$	(p)	
N4	$FN_{Veg,SOMc} =$	$((1 - f_{met}) \cdot I_N + CWD_N) \cdot f_{struct,SOM}$	(q)	
N5	$FN_{LITm,SAPb} =$	$FC_{LITm,SAPb} \cdot \left( \frac{N_{LITm}}{C_{LITm}} \right)$	(j)(l)	as in MIMICS
N6	$FN_{LITs,SAPb} =$	$FC_{LITs,SAPb} \cdot \left( \frac{N_{LITs}}{C_{LITs}} \right)$	(k)(l)	as in MIMICS
N7	$FN_{SOMa,SAPb} =$	$FC_{SOMa,SAPb} \cdot \left( \frac{N_{SOMa}}{C_{SOMa}} \right)$	(r)(l)	as in MIMICS
N8	$FN_{LITm,SAPf} =$	$FC_{LITm,SAPf} \cdot \left( \frac{N_{LITm}}{C_{LITm}} \right)$	(j)(m)	as in MIMICS
N9	$FN_{LITs,SAPf} =$	$FC_{LITs,SAPf} \cdot \left( \frac{N_{LITs}}{C_{LITs}} \right)$	(k)(m)	as in MIMICS
N10	$FN_{SOMa,SAPf} =$	$FC_{SOMa,SAPf} \cdot \left( \frac{N_{SOMa}}{C_{SOMa}} \right)$	(r)(m)	as in MIMICS
N11	$FN_{SOMc,SOMa} =$	$FC_{SOMc,SOMa} \cdot \left( \frac{N_{SOMc}}{C_{SOMc}} \right)$	(q)(r)	
N12	$FN_{SOMp,SOMa} =$	$FC_{SOMp,SOMa} \cdot \left( \frac{N_{SOMp}}{C_{SOMp}} \right)$	(p)(r)	
N13	$FN_{SAPb,SOMp} =$	$FC_{SAPb,SOMp} \cdot \left( \frac{N_{SAPb}}{C_{SAPb}} \right)$	(l)(p)	
N14	$FN_{SAPb,SOMc} =$	$FC_{SAPb,SOMc} \cdot \left( \frac{N_{SAPb}}{C_{SAPb}} \right)$	(l)(q)	
N15	$FN_{SAPb,SOMa} =$	$FC_{SAPb,SOMa} \cdot \left( \frac{N_{SAPb}}{C_{SAPb}} \right)$	(l)(r)	
N16	$FN_{SAPf,SOMp} =$	$FC_{SAPf,SOMp} \cdot \left( \frac{N_{SAPf}}{C_{SAPf}} \right)$	(m)(p)	
N17	$FN_{SAPf,SOMc} =$	$FC_{SAPf,SOMc} \cdot \left( \frac{N_{SAPf}}{C_{SAPf}} \right)$	(m)(q)	
N18	$FN_{SAPf,SOMa} =$	$FC_{SAPf,SOMa} \cdot \left( \frac{N_{SAPf}}{C_{SAPf}} \right)$	(m)(r)	
N19	$FN_{EcM,SOMp} =$	$FC_{EcM,SOMp} \cdot \left( \frac{N_{EcM}}{C_{EcM}} \right)$	(n)(p)	
N20	$FN_{EcM,SOMc} =$	$FC_{EcM,SOMc} \cdot \left( \frac{N_{EcM}}{C_{EcM}} \right)$	(n)(q)	
N21	$FN_{EcM,SOMa} =$	$FC_{EcM,SOMa} \cdot \left( \frac{N_{EcM}}{C_{EcM}} \right)$	(n)(r)	
N22	$FN_{AM,SOMp} =$	$FC_{AM,SOMp} \cdot \left( \frac{N_{AM}}{C_{AM}} \right)$	(o)(p)	
N23	$FN_{AM,SOMc} =$	$FC_{AM,SOMc} \cdot \left( \frac{N_{AM}}{C_{AM}} \right)$	(o)(q)	
N24	$FN_{AM,SOMa} =$	$FC_{AM,SOMa} \cdot \left( \frac{N_{AM}}{C_{AM}} \right)$	(o)(r)	
N25	$FN_{SOMp,EcM} =$	$FC_{EcMdecompSOMp} \cdot \left( \frac{N_{SOMp}}{C_{SOMp}} \right)$	(g)(e)	
N26	$FN_{SOMc,EcM} =$	$FC_{EcMdecompSOMc} \cdot \left( \frac{N_{SOMc}}{C_{SOMc}} \right)$	(h)(e)	
N27	$FN_{IN,EcM} =$	$V_{max,myc} \cdot N_{IN} \cdot \left( \frac{C_{EcM}}{(C_{EcM} + K_{m,myc}/dz)} \right) \cdot r_{myc}$	(s)(u)(n)	Baskaran et al. (2017)+ mod. term,

– Continued on next page

**Table A4** – Continued from previous page

Eq.	Flux Name	Rate functions	Used in eqn	Notes
N28	$FN_{IN,AM} =$	$V_{max,myc} \cdot N_{IN} \cdot \left( \frac{C_{AM}}{(C_{AM} + K_{m,myc}/dz)} \right) \cdot r_{myc}$	(s)(u)(o)	$IN = N_{NO3} + N_{NH4,sol}$ Baskaran et al. (2017)+ mod. term
N29	$FN_{EcM,Veg} =$	$(FN_{IN,EcM} + FN_{SOMc,EcM} + FN_{SOMP,EcM}) - CUE_{EcM} \cdot FC_{Veg,EcM} \cdot (1 - f_{enz})/CN_{EcM}$ or lower, if N limited (reduced CUE)	(n)	$IN = N_{NO3} + N_{NH4,sol}$
N30	$FN_{AM,Veg} =$	$FN_{IN,AM} - CUE_{AM} \cdot FC_{Veg,AM}/CN_{AM}$ or lower, if N limited (reduced CUE)	(o)	$IN = N_{NO3} + N_{NH4,sol}$
N31	$FN_{run+leach} =$	$N_{NO3} \cdot \left( \frac{QDRAI}{H2O_{tot}} + \frac{QRUNOFF}{H2O_{top5cm}} \right)$	(u)	See CTSM doc. 2.22.6
N32	$FN_{DEP} =$	$NDEP\_TO\_SMINN \cdot NDEP\_PROF$	(s)	
N33	$FN_{IN,Veg} =$	$N_{IN} \cdot k_{uptake}$	(s)(u)	$IN = N_{NO3} + N_{NH4,sol}$
N34	$FN_{NH4,NO3} =$	$NH4 \cdot k_{nitr}$ or zero if temp. is below freezing	(s)(u)	based on CTSM doc. chapter 2.22.5
N35	$FN_{sol,sorp} =$			
N36	$FN_{IN,SAPb} =$ or =	$(1 - NUE) \cdot U_{Nb} - CUE_b \cdot U_{Cb}/CN_b$ $f_b \cdot N_{for\_sap}$ if limited N	(l)(s)(u)	$IN = N_{NO3} + N_{NH4,sol}$
N37	$FN_{IN,SAPf} =$ or =	$(1 - NUE) \cdot U_{Nf} - CUE_f \cdot U_{Cf}/CN_f$ $(1 - f_b) \cdot N_{for\_sap}$ if limited N	(m)(s)(u)	

**Table A5.** Description of parameters and other relevant sizes used in the model.

Parameter	Description	Expression/Value	Units	Notes
$f_{met}$	Met. frac. of plant litter	$0.75 \cdot (0.85 - 0.013 \cdot \min(40, \text{lignin} : N))$	-	Wieder et al. (2015)
$f_{clay}$	Clay fraction in soil	0.0-1.0	-	
$T$	Soil temperature	-	°C	Vary with season and depth
Michaelis-Menten kinetics param. for SAP: Wieder et al. (2015), German et al. (2012)				
$V_{max}$	Max reaction velocity	$\exp(V_{slope} \cdot T + V_{int}) \cdot a_V \cdot V_{mod} \cdot r_{moist}$	mg(mg) <sup>-1</sup> h <sup>-1</sup>	
$K_m$	Half saturation constant	$\exp(K_{slope} \cdot T + K_{int}) \cdot a_K \cdot K_{mod}$	mgCcm <sup>-3</sup>	
$K_{slope}$	Regression coefficient	LIT: 0.017, SOMa: 0.027	ln(mgCcm <sup>-3</sup> )°C <sup>-1</sup>	For all 6 fluxes
$V_{slope}$	Regression coefficient	0.063	ln(mg(mg) <sup>-1</sup> h <sup>-1</sup> )°C <sup>-1</sup>	For all 6 fluxes
$K_{int}$	Regression intercept	3.19	ln(mgCcm <sup>-3</sup> )	Directly Wieder et al. (2015)
$V_{int}$	Regression intercept	5.47	ln(mg(mg) <sup>-1</sup> h <sup>-1</sup> )	Directly Wieder et al. (2015)
$a_V$	Tuning coefficient	$1.25 \cdot 10^{-8}$	-	
$P$	Phys. protection scalar used in $K_{mod}$	$1/(2.0 \cdot \exp(-2\sqrt{f_{CLAY}}))$	-	Wieder et al. (2015)
$a_K \cdot K_{mod}$	Tuning coefficients	1.953125, 7.81250, 3.90625 · $P$ , 7.8125, 3.90625, 2.604167 · $P$	-	As in MIMICS imp. in CLM for LITm, LITs, SOMa
$V_{mod}$	Modifies $V_{max}$	10.0, 3.0, 10.0, 3.0, 5.0, 2.0	-	for LITm, LITs, SOMa entering SAPb, SAPf
$KO$	Increase $K_m$ in eq. C11	6	-	Kyker-Snowman et al. (2020)
$k_{myc,som}$	Turnover rate	$1.14 \cdot 10^{-4}$	h <sup>-1</sup>	1y <sup>-1</sup> as Sulman et al. (2019) and (Baskaran et al., 2017)
$k_{SAPb,som}$	Turnover rate of SAPb	$5.2 \cdot 10^{-4} \cdot \exp(0.3 \cdot f_{met}) \cdot \max(p_{mod}, m_{mod})$	h <sup>-1</sup>	Wieder et al. (2015) + mod. term
$k_{SAPf,som}$	Turnover rate of SAPf	$2.4 \cdot 10^{-4} \cdot \exp(0.1 \cdot f_{met}) \cdot \max(p_{mod}, m_{mod})$	h <sup>-1</sup>	Wieder et al. (2015) + mod. term
$p_{mod}$ scales with root profile, $m_{mod} = 0.1$ is the minimum value of the modifier. $m_{mod}$ is used when $T < 0$				
$k_{desorp}$	desorption rate	$2 \cdot 10^{-6} \cdot \exp(-4.5 \cdot f_{clay})$	h <sup>-1</sup>	Modified from Kyker-Snowman et al. (2020)

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**Table A5** – Continued from previous page

Parameter	Description	Expression/Value	Units	Notes
$K_{MO}$	Mycorrhizal decay rate	$3.42 \cdot 10^{-6}$	$\text{m}^2 \text{gC}^{-1} \text{hr}^{-1}$	Baskaran et al. (2017)
$V_{max,myc}$	Max. mycorrhizal uptake of inorg N	$2.05 \cdot 10^{-4}$	$\text{g} \cdot \text{g}^{-1} \text{h}^{-1}$	Baskaran et al. (2017) for EcM, we also use it for AM
$K_{m,myc}$	Half saturation constant of ectomycorrhizal uptake of inorg N	0.08	$\text{gNm}^{-2}$	Baskaran et al. (2017) for EcM, we also use it for AM
$CUE_{EcM}$	Growth efficiency of mycorrhiza	0-0.5	-	Sulman et al. (2019)
$CUE_{AM}$	Growth efficiency of mycorrhiza	0-0.5	-	Sulman et al. (2019)
$CUE_b$	Growth efficiency of sap. bacteria	0-0.4	-	Determined by N availability.
$CUE_f$	Growth efficiency of sap. fungi	0-0.7	-	
$NUE$	Nitrogen use efficiency of saprotrophs	0.8	-	Mooshammer et al. (2014a)
$r_{moist}$	Moisture function:	$\max \left( 0.05, P \cdot \left( \frac{\Theta_{liq}}{\Theta_{sat}} \right)^3 \cdot \left( 1 - \frac{\Theta_{liq}}{\Theta_{sat}} - \frac{\Theta_{frozen}}{\Theta_{sat}} \right)^{2.5} \right)$	-	Wieder et al. (2017), Sulman et al. (2014)
$r_{myc}$	Mycorrhizal modifier	0-1	-	
$f_{SAPb,SOMp}$	Frac. necromass into SOMp	$0.3 \cdot \exp(1.3 \cdot f_{clay})$	-	
$f_{SAPb,SOMc}$	Frac. necromass into SOMc	$0.1 \cdot \exp(-3 \cdot f_{met})$	-	
$f_{SAPb,SOMa}$	Frac. necromass into SOMa:	$1 - (f_{SAPb,SOMp} + f_{SAPb,SOMc})$	-	
$f_{SAPf,SOMp}$	Frac. necromass into SOMp	$0.2 \cdot \exp(0.8 \cdot f_{clay})$	-	
$f_{SAPf,SOMc}$	Frac. necromass into SOMc	$0.3 \cdot \exp(-3 \cdot f_{met})$	-	
$f_{SAPf,SOMa}$	Frac. necromass into SOMa:	$1 - (f_{SAPf,SOMp} + f_{SAPf,SOMc})$	-	
$f_{EcM,SOMp}$	Frac. necromass into SOMp	0.4	-	Assumed
$f_{EcM,SOMc}$	Frac. necromass into SOMc	0.2	-	Assumed
$f_{EcM,SOMa}$	Frac. necromass into SOMa	0.4	-	Assumed
$f_{AM,SOMp}$	Frac. necromass into SOMp	0.3	-	Assumed
$f_{AM,SOMc}$	Frac. necromass into SOMc	0.4	-	Assumed
$f_{AM,SOMa}$	Frac. necromass into SOMa	0.3	-	Assumed
$f_{enz}$	Frac. of EcM C uptake used for enzyme prod.	0.10	-	Assumed
$f_{use}$	Frac. C released by mining taken up by EcM.	0.10	-	Assumed
$f_{alloc,i}$	Frac. of C from plant alloc. to myc. $i$	0-1	-	See Sect. 2.1.2
$f_{met,SOM}$	Frac. of met. litter prod. going directly to SOMp	0.5	-	
$f_{struct,SOM}$	Frac. of struct. litter prod. going directly to SOMc	0.5	-	
$dz$	Soil layer thickness		m	Correspond to layer thickness in CLM

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**Table A5** – Continued from previous page

Parameter	Description	Expression/Value	Units	Notes
$D$	Diffusion coefficient	$1.14 \cdot 10^{-8}$	$\text{m}^2 \text{hr}^{-1}$	Koven et al. (2013): $1 \text{cm}^2 \text{yr}^{-1}$ 1/3 of this value for $N_{NH4,sorp}$
$CN_b$	Optimal CN ratio for bacteria	5	-	Mouginot et al. (2014)
$CN_f$	Optimal CN ratio for sap. fungi	8	-	Mouginot et al. (2014)
$CN_m$	Optimal CN ratio for myc. fungi	20	-	Baskaran et al. (2017), Wallander et al. (2003)
$BD_{soil}$	Soil Bulk density	$1.6 \cdot 10^6$	$\text{g} \cdot \text{m}^{-3}$	Sieczka and Koda (2016)
$NH4_{sorp,max}$	Max. adsorption capacity	$0.09 \cdot BD_{soil} \cdot 10^{-3} = 144$	$\text{gNH4} \cdot \text{m}^{-3}$	converted from Sieczka and Koda (2016)
$K'_L$	Modified Langmuir constant	$0.4 \cdot soil\_water\_frac^{-1}$	$\text{m}^3 \cdot \text{gNH4}^{-1}$	converted from Sieczka and Koda (2016)
$k$	Rate constant ammonium sorption	$0.0167 \cdot 60 \cdot 10^3 \cdot BD_{soil}^{-1}$	$\text{m}^3 \text{g}^{-1} \text{hr}^{-1}$	converted from Sieczka and Koda (2016)

**Table A6.** CLM variables used in MIMICS+

CLM-BGC variable	Units	Long name	Notes
LEAFC_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	leaf C litterfall	
FROOTC_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	fine root C litterfall	
CWDC_TO_LITR2C_vr	$\text{gCm}^{-3}\text{s}^{-1}$	decomp. of coarse woody debris C to litter 2 C	
CWDC_TO_LITR3C_vr	$\text{gCm}^{-3}\text{s}^{-1}$	decomp. of coarse woody debris C to litter 3 C	
M_LEAFC_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	leaf C mortality	
M_FROOTC_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	fine root C mortality	
M_LEAFC_STORAGE_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	leaf C storage mortality	Input to met. lit. (LITm)
M_LEAFC_XFER_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	leaf C transfer mortality	Input to met. lit. (LITm)
M_GRESP_STORAGE_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	growth respiration storage mortality	Input to met. lit. (LITm)
M_GRESP_XFER_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	growth respiration transfer mortality	Input to met. lit. (LITm)
M_FROOTC_STORAGE_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	fine root C storage mortality	Input to met. lit. (LITm)
M_FROOTC_XFER_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	fine root C transfer mortality	Input to met. lit. (LITm)
M_LIVECROOTC_XFER_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	live coarse root C transfer mortality	Input to met. lit. (LITm)
M_DEADCROOTC_XFER_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	dead coarse root C transfer mortality	Input to met. lit. (LITm)
M_LIVECROOTC_STORAGE_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	live coarse root C fire mortality to litter	Input to met. lit. (LITm)
M_LIVESTEMC_STORAGE_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	live stem C storage mortality	Input to met. lit. (LITm)
M_LIVESTEMC_XFER_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	live stem C transfer mortality	Input to met. lit. (LITm)
M_DEADSTEMC_STORAGE_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	dead stem C storage mortality	Input to met. lit. (LITm)
M_DEADSTEMC_XFER_TO_LITTER	$\text{gCm}^{-2}\text{s}^{-1}$	dead stem C transfer mortality	Input to met. lit. (LITm)
LEAFN_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$	leaf N litterfall	Partitioned based on $f_{MET}$
FROOTN_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$	fine root N litterfall	Partitioned based on $f_{MET}$
CWDN_TO_LITR2N_vr	$\text{gNm}^{-3}\text{s}^{-1}$	decomp. of coarse woody debris N to litter 2 C	Input to structural litter (LITs)
CWDN_TO_LITR3N_vr	$\text{gNm}^{-3}\text{s}^{-1}$	decomp. of coarse woody debris C to litter 3 C	Input to structural litter (LITs)
M_LEAFN_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$	leaf N mortality	Partitioned based on $f_{MET}$ .
M_FROOTN_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$	fine root N mortality	Partitioned based on $f_{MET}$ .
M_LEAFN_STORAGE_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$	leaf C storage mortality	Input to met. lit. (LITm)
M_LEAFN_XFER_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)
M_FROOTN_STORAGE_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)

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**Table A6** – Continued from previous page

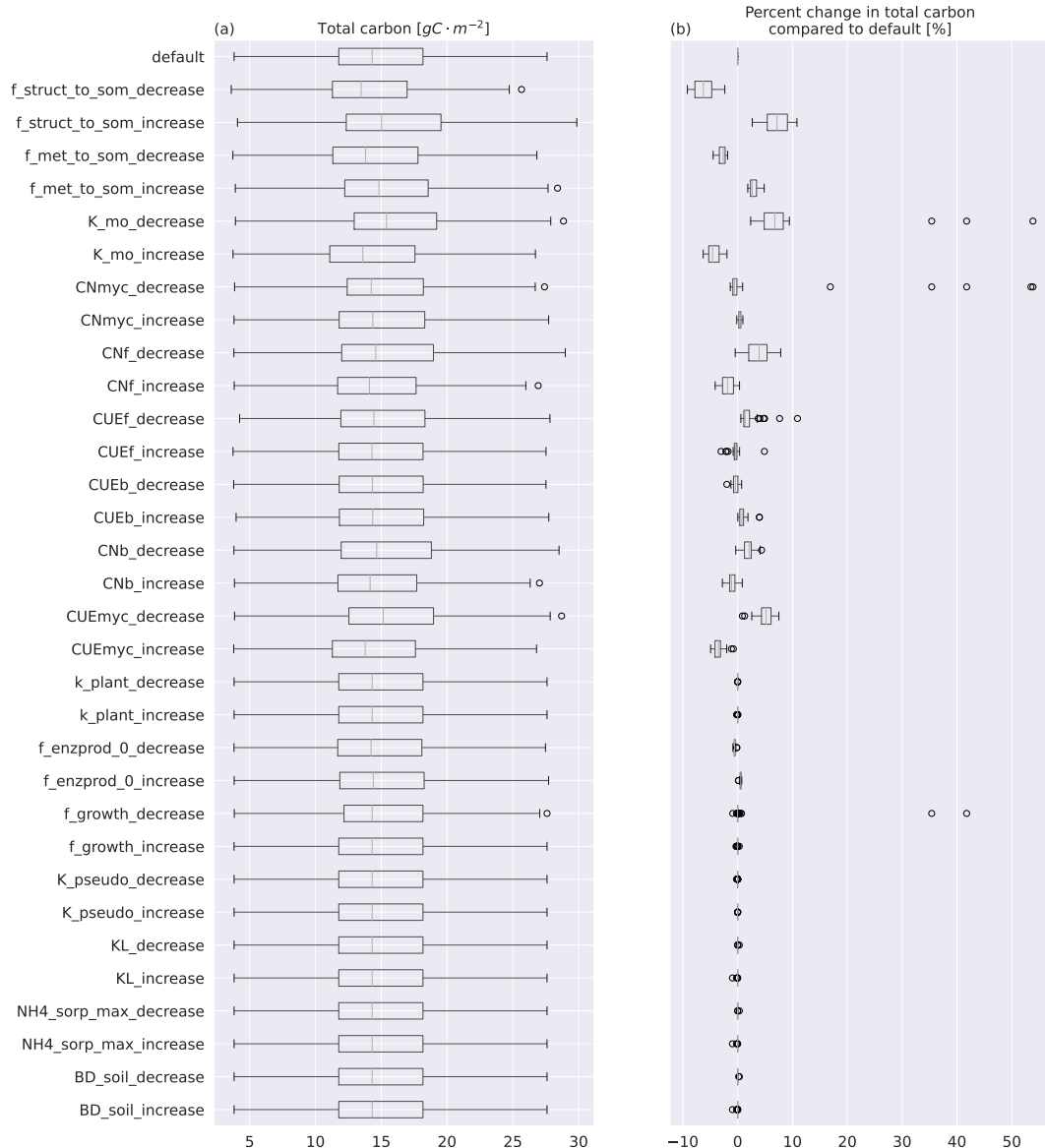
CLM-BGC variable	Units	Long name	Notes
M_FROOTN_XFER_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)
M_LIVECROOTN_XFER_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)
M_DEADCROOTN_XFER_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)
M_LIVECROOTN_STORAGE_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)
M_LIVESTEMN_STORAGE_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)
M_LIVESTEMN_XFER_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)
M_DEADSTEMN_STORAGE_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)
M_DEADSTEMN_XFER_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)
M_RERTRANSN_TO_LITTER	$\text{gNm}^{-2}\text{s}^{-1}$		Input to met. lit. (LITm)
NPP_NACTIVE	$\text{gCm}^{-2}\text{s}^{-1}$		Partitioned between EcM and AM based on $f_{alloc,i}$
NDEP_TO_SMINN	$\text{gNm}^{-2}\text{s}^{-1}$	atmospheric N deposition to soil mineral N	N deposition to NH4 pool
LEAF_PROF	$\text{m}^{-1}$	profile for litter C and N inputs from leaves	Multiplied with LEAF_TO_LITTER to get rates for each layer
FROOT_PROF	$\text{m}^{-1}$	profile for litter C and N inputs from fine roots	Multiplied with FROOT_TO_LITTER to get rates for each layer
CROOT_PROF	$\text{m}^{-1}$	profile for litter C and N inputs from coarse roots	used for input from mortality
STEM_PROF	$\text{m}^{-1}$	profile for litter C and N inputs from stems	used for input from mortality
NDEP_PROF	$\text{m}^{-1}$	profile for atmospheric N deposition	Multiplied with NDEP_TO_SMINN to get deposition for each layer
Environmental variables:			
TSOI	K	soil temperature	Converted to $^{\circ}\text{C}$
WATSAT	$\text{mm}^3\text{mm}^{-3}$	saturated soil water content (porosity)	Used for calculating $r_{moist}$
SOILLIQ	$\text{kg} \cdot \text{m}^{-2}$	soil liquid water	Used for calculating $r_{moist}$
SOILICE	$\text{kg} \cdot \text{m}^{-2}$	soil ice water	Used for calculating $r_{moist}$
W_SCALAR	-	Moisture (dryness) inhibition of decom.	Used in nitrification algorithm
T_SCALAR	-	temperature inhibition of decomposition	Used in nitrification algorithm
QDRAI	$\text{mm} \cdot \text{s}^{-1}$	sub-surface drainage	Used for calculating leaching

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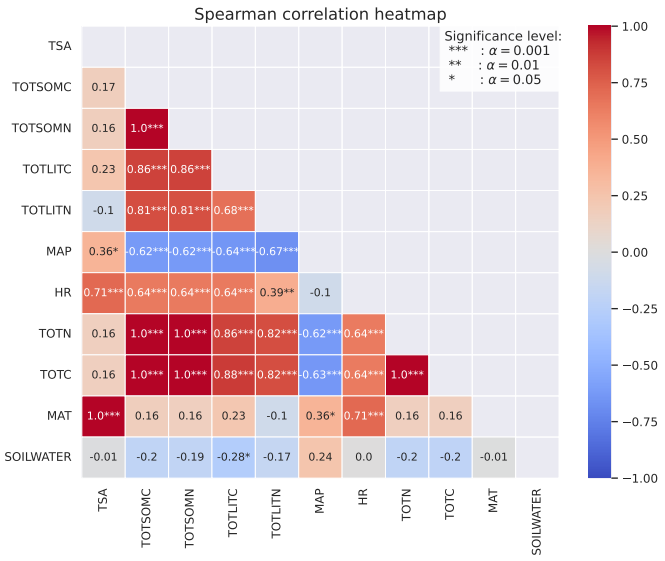
**Table A6** – Continued from previous page

CLM-BGC variable	Units	Long name	Notes
QOVER	$\text{mm} \cdot \text{s}^{-1}$	surface runoff	Used for calculating Runoff
nbedrock	-	index of shallowest bedrock layer	for determining how many layers to use in the simulations
Read from surface data file:			
PCT_CLAY	-	percent CLAY	
PCT_NAT_PFT	-	percent plant functional type on the nat. veg landunit	

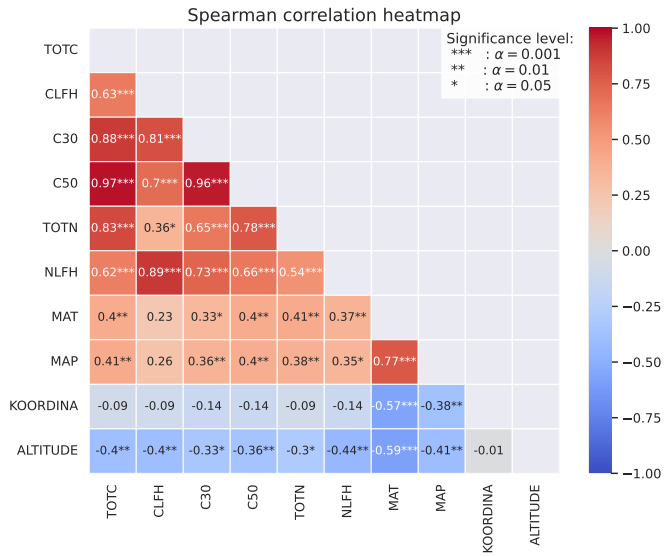
## Appendix B: Additional figures



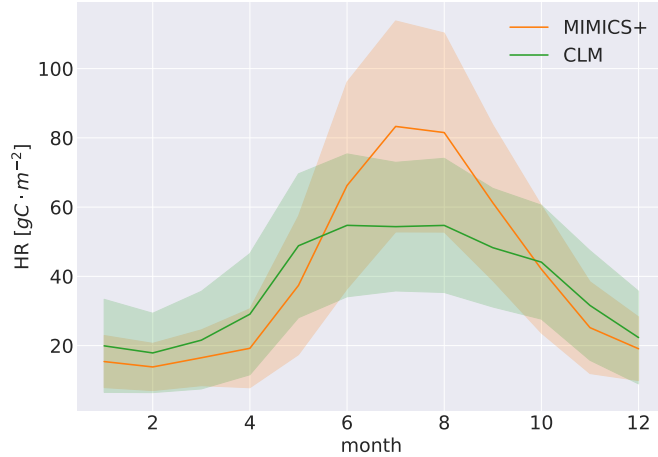
**Figure B1.** (a) Boxplots of total C for the 50 sites modeled with MIMICS+. The top box is default simulations, while the rest are simulations with one parameter perturbed with either a 25 % increase or decrease compared to the default value. (b) Boxplots of the percentage change from default of the same simulations as in (a). The line in each box is the median, the box upper and lower edges are the 75th and 25th percentiles, respectively. The whiskers extend from the box by 1.5 times the inter-quartile range.



**Figure B2.** Spearman correlation coefficients between different variables calculated from the CLM simulations of the 50 sites. The stars represent significance level of the correlation. Numbers without stars are not significant ( $p > .05$ ). The color indicate whether the correlation is positive (red) or negative (blue), and the shade indicate the strength of the correlation.

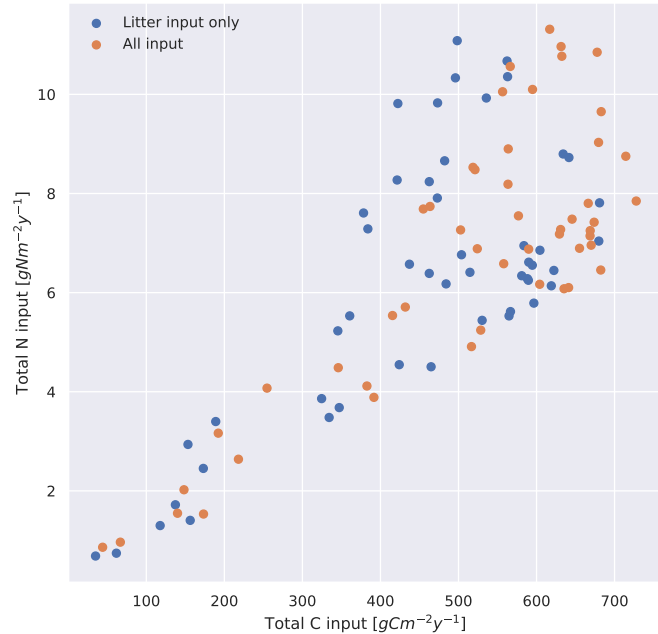


**Figure B3.** Spearman correlation coefficients between different observed variables at 50 sites (Strand et al., 2016). The stars represent significance level of the correlation. Numbers without stars are not significant ( $p > .05$ ). The color indicate whether the correlation is positive (red) or negative (blue), and the shade indicate the strength of the correlation.



**Figure B4.** Modeled mean seasonal heterotrophic respiration over the years 1988–1992 and the 50 modeled sites. The shading indicate the standard deviation among the 50 sites.

### Appendix C: Input plot



**Figure C1.** Yearly mean input of carbon and nitrogen to MIMICS+ from CLM for each of the 50 site simulations (averaged over 1988-1992). The blue dots show litter input only, while the orange dots also include the C allocated to mycorrhizal pools and N deposition.

605 *Author contributions.* ERA and TKB developed the model. ERA ran simulations and wrote the manuscript. All authors contributed to the analyses and editing of the manuscript.

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