Nutrient Dynamics in a Coupled Terrestrial Biosphere and Land Model (ELM-FATES)

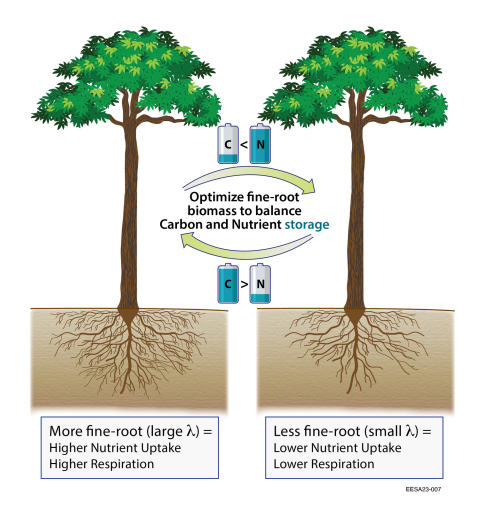
Ryan G Knox¹, Charles D. Koven¹, William J. Riley¹, Anthony P. Walker², Stuart Joseph Wright³, Jennifer A. Holm¹, Xinyuan Wei², Rosie A. Fisher⁴, Qing Zhu¹, Jinyun Tang¹, Daniel M. Ricciuto², Jacquelyn Shuman⁵, Xiaojuan Yang², Lara M Kueppers¹, and Jeffrey Chambers¹

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Abstract

We present a representation of nitrogen and phosphorus cycling in the vegetation demography model the Functionally Assembled Terrestrial Ecosystem Simulator (FATES), within the Energy Exascale Earth System (E3SM) land model. This representation is modular, and designed to allow testing of multiple hypothetical approaches for carbon-nutrient coupling in plants. The model tracks nutrient uptake, losses via turnover from both live plants and mortality into soil decomposition, and allocation during tissue growth for a large number of size- and functional-type-resolved plant cohorts within a time-since-disturbance-resolved ecosystem. Root uptake is governed by fine root biomass, and plants vary in their fine root carbon allocation in order to balance carbon and nutrient limitations to growth. We test the sensitivity of the model to a wide range of parameter variations and structural representations, and in the context of observations at Barro Colorado Island, Panama. A key model prediction is that plants in the high-light-availability canopy positions allocate more carbon to fine roots than plants in low-light understory environments, given the widely different carbon versus nutrient constraints of these two niches within a given ecosystem. This model provides a basis for exploring carbon-nutrient coupling with vegetation demography within Earth System Models (ESMs).



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12 Abstract

We present a representation of nitrogen and phosphorus cycling in the vegetation 13 demography model the Functionally Assembled Terrestrial Ecosystem Simulator (FATES), 14 within the Energy Exascale Earth System (E3SM) land model. This representation is 15 modular, and designed to allow testing of multiple hypothetical approaches for carbon-16 nutrient coupling in plants. The model tracks nutrient uptake, losses via turnover from 17 both live plants and mortality into soil decomposition, and allocation during tissue growth 18 for a large number of size- and functional-type-resolved plant cohorts within a time-since-19 20 disturbance-resolved ecosystem. Root uptake is governed by fine root biomass, and plants vary in their fine root carbon allocation in order to balance carbon and nutrient limi-21 tations to growth. We test the sensitivity of the model to a wide range of parameter vari-22 ations and structural representations, and in the context of observations at Barro Col-23 orado Island, Panama. A key model prediction is that plants in the high-light-availability 24 canopy positions allocate more carbon to fine roots than plants in low-light understory 25 environments, given the widely different carbon versus nutrient constraints of these two 26 niches within a given ecosystem. This model provides a basis for exploring carbon-nutrient 27 coupling with vegetation demography within Earth System Models (ESMs). 28

²⁹ Plain Language Summary

This work introduces a new set of nutrient cycling hypotheses integrated with a coupled terrestrial biosphere model. This includes the cycling of carbon, nitrogen and phosphorus, and focuses mainly on plant acquisition, allocation, and turnover. An analysis shows the model offers reasonable response to perturbations in parameter constants and boundary conditions, considering its design balance between process complexity and parameterization burden.

36	Key Points:
37	• The nutrient enabled ELM-FATES model represents reasonable pattern responses
38	to nutrient availability and parameter perturbations.
39	• The model has been designed to introduce a reasonably small parameterization
40	burden, considering the total number of newly introduced parameters and the rel-
41	ative ease of finding values (i.e. directly from observations, or through calibration).
42	• The model formulation described here does not make any assertions that it offers
43	a complete representation of nutrient and carbon dynamics, rather that these are
44	a set hypothesis that can capture certain element of carbon-nutrient dynamics and
45	can be further intercompared with other hypotheses.

46 **1** Introduction

Projections of the global climate system response to anthropogenic CO_2 emissions 47 require coupled models of the climate system and carbon cycle. Much of the uncertainty 48 in current climate projections arises from the global terrestrial carbon cycle, and in par-49 ticular the responses of plants to elevated CO_2 (Arora et al., 2020). Many current Earth 50 System Models (ESMs) do not take into account plant size structure, disturbance his-51 tory, and other aspects known to govern ecosystem function and thereby current and fu-52 ture responses to anthropogenic pressures (D. Purves & Pacala, 2008). Additionally, lim-53 itation by nutrients of plant productivity under elevated CO_2 has been shown to strongly 54 affect both the historical and future uptake of carbon (Hungate et al., 2003; P. Thorn-55 ton et al., 2007; Zaehle & Friend, 2010; Wang et al., 2015). The importance of includ-56

ing nutrient dynamics in projecting the global terrestrial carbon budget is evidenced by 57 its expanded role in Earth System land models, such as CASACNP (Wang et al., 2010), 58 GFDL LM4.1-BNF (Kou-Giesbrecht et al., 2021a; Sulman et al., 2019), ED2-MEND-59 NCOM (Medvigy et al., 2019), and Quincy v1.0 (Thum et al., 2019) to name a few. The 60 processes that govern nutrient cycling rates in ecosystems are highly uncertain, since many 61 of them occur belowground where observation is more difficult than in plant canopies. 62 To allow exploration of this epistemic uncertainty, we propose here a modular approach 63 to representing nutrient cycling that facilitates exploration of alternative process hypothe-64 ses and parameter and structural uncertainty quantification. 65

This manuscript describes a modeling methodology for plant acquisition, storage, 66 and allocation of nutrients and carbon within the terrestrial biosphere of an ESM. This 67 manuscript also describes how the plant nutrient dynamics interface with existing soil 68 nutrient hypotheses, but does not introduce new soil modeling hypotheses. This system 69 is an extension of the coupled modeling framework of the Energy Exascale Earth Sys-70 tem Model (E3SM) (Caldwell et al., 2019) and the Functionally Assembled Terrestrial 71 Ecosystem Simulator (FATES). E3SM includes a land model (ELM) among other com-72 ponents such as atmosphere, ocean, ice, and human. The terrestrial vegetation simulated 73 in FATES is based on the plant size and time-since-disturbance structured approach de-74 rived from the Ecosystem Demography model (Moorcroft et al., 2001; R. A. Fisher et 75 al., 2015). 76

The nutrient modeling framework we describe here can be summarized in three model 77 components: 1) a new module that handles on-plant nutrient and carbon allocations to 78 different organs, designated the Plant Allocation, Reactions, and Transport Extensible 79 Hypotheses (PARTEH), 2) a restructuring of the rest of the FATES model to track vari-80 able chemical elements in seed, unfragmented litter, and coarse woody debris pools, and 81 3) a means of acquisition and competition for nutrients by FATES plants amongst mi-82 crobes and mineral surfaces. An evaluation of the model at a tropical test-bed site fol-83 lows. 84

2 Model Description - Plant-Soil Nutrient Dynamics

The Energy Exascale Earth System Model Land Model component (Burrows et al., 86 2020), resolves numerous processes related to the cycling of water, energy, carbon, ni-87 trogen and phosphorus in natural and anthropogenic ecosystems. Soil decomposition is 88 handled via a derivative of the CENTURY approach (C. Koven et al., 2013; Parton et 89 al., 1988). Total ecosystem nitrogen fixation has been represented in ELM by assuming 90 proportionality with either evapotranspiration or net primary productivity (Cleveland 91 et al., 1999; P. Thornton et al., 2007). Symbiotic fixation at the plant level is introduced 92 in the Methods section and will be described in more detail. When symbiotic fixation 93 is represented by FATES, the total ecosystem fixation module in ELM is scaled down 94 to represent on free-living fixation in the soil. Soil biogeochemical dynamics lead nitrogen to cascade from organic pools with different turnover times to ammonium (NH_4) and 96 nitrate (NO_3) pools. ELM contains two alternative representations of competition for 97 these these nutrient species amongst plants, microbes (decomposers for organic pools, 98 as well as nitrifiers and denitrifiers for mineral nitrogen pools), and mineral surfaces (for 99 phosphorus). These are the Relative Demand (RD) approach (P. Thornton et al., 2007; 100 X. Yang et al., 2014, 2019) and a Capacity-Based approach that applies the Equilibrium 101 Chemistry Approximation (CB) (Tang & Riley, 2013; Zhu et al., 2016, 2019). 102

FATES is a vegetation demography model (R. A. Fisher et al., 2015; C. D. Koven et al., 2020) that represents the demographics of vegetation using plant size and timesince-disturbance structured scaling algorithms defined in the Ecosystem Demography (ED) Model (Moorcroft et al., 2001). FATES represents vegetation by grouping plants of similar size and functional type into cohorts, which inhabit patches of the landscape

that are defined by their time since last disturbance. Unlike unstructured vegetation mod-108 els, which treat growth and mortality processes as changes to the size of whole ecosystem-109 level carbon and nutrient pools, cohort-based vegetation models like FATES explicitly 110 track the growth of plants, the size (volume, height, etc) growth of various components, 111 and the resulting carbon and nutrient pools of their tissues using allometrically-defined 112 scaling relationships with stem diameter. FATES also allows for competition for light 113 between plant types in the same vertical profile, which leads to self-thinning dynamics 114 and other spatio-temporal changes in vegetation composition to emerge as a function of 115 variation in plant functional traits. FATES estimates mortality at the plant cohort scale, 116 based on several factors including carbon starvation, understory impact survival, hydraulic 117 stress (R. A. Fisher et al., 2015), background mortality (i.e. unspecified or unknown effects) (Moorcroft 118 et al., 2001), fire (Thonicke et al., 2010), and relationships with plant age or size (Needham 119 et al., 2020). In the current version of FATES, the time-since-disturbance patch discretiza-120 tion only resolves heterogeneity in the above-ground environment, with all cohorts on 121 all patches drawing water and nutrients from the same soil pools. 122

FATES represents a variety of processes, including but not limited to: photosyn-123 thesis and its coupling with water limitations on stomatal conductance (Farquhar et al., 124 1980; Collatz et al., 1991; Oleson et al., 2013; G. Bonan et al., 2014), respiration (Ryan, 125 1991) of live tissues, vertical distribution of canopy functional trait parameters (G. B. Bo-126 nan et al., 2012), radiation scattering (Norman, 1979), phenology (Botta et al., 2000), 127 and turnover into coarse woody debris and fine litter (R. A. Fisher et al., 2015; Oleson 128 et al., 2013). FATES' radiation scattering module accounts for both the vertical struc-129 ture of vegetation and the variable scattering characteristics of different plant functional 130 groups in parallel (R. A. Fisher et al., 2015). The spatial configuration of the canopy scat-131 tering elements is driven by a modified perfect plasticity approximation (D. W. Purves 132 et al., 2008; R. A. Fisher et al., 2015; C. D. Koven et al., 2020). While FATES can op-133 tionally utilize sophisticated representations of plant hydraulics (Christoffersen et al., 134 2014; Fyllas et al., 2014; Fang et al., 2022) and fire (Thonicke et al., 2010; Buotte et al., 135 2021; Ma et al., 2021), in order to maintain a focus on nutrient dynamics, these options 136 were not active in the modeling exercises described herein. Details on all of these pro-137 cess representations can be found in the FATES technical documentation (FATES-Development-138 Team, 2019). 139

The introduction of nutrients to FATES, via the PARTEH approach to (Plant Re-140 source Allocations, Reactions, and Transport Extensible Hypotheses) described here, fol-141 lows a sequence of operations that are shown in Figure 1. The descriptions of these pro-142 cesses follow the flow-chart order and can be found in the following subsections: sym-143 biotic fixation in 2.1, plant acquisition of aqueous soil nutrients in 2.2, re-absorption dur-144 ing senescent turnover in 2.3, updating the target fine-root biomass (the optimal fine-145 root biomass associated with the plant's nutrient requirements) in 2.6, and allocation 146 to the various plant organs in 2.5. Soil hydrology in ELM is the same as that described 147 in the CLM technical manual (Oleson et al., 2013). 148

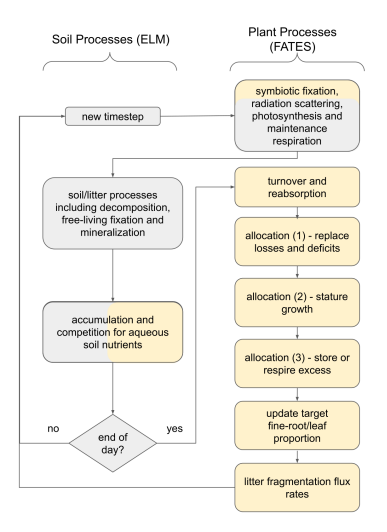


Figure 1. Flow-chart of key processes and order of operations for nutrient cycling in ELM-FATES. This chart places more emphasis on plant-side processes. New processes described in this manuscript are shown in yellow boxes. Grey boxes indicate a pre-existing but relevant processes in the model. For the process of soil nutrient competition, nothing has changed from the original schemes, except how plants present themselves as competitors (shown as with a split grey/yellow color). In the box highlighting symbiotic fixation, scattering, etc., symbiotic fixation is a new process and the others are un-modified.

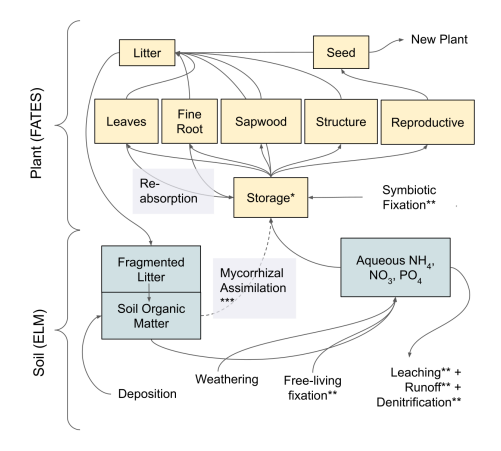


Figure 2. Diagram showing the key pools and fluxes for nitrogen and phosphorus cycling in ELM-FATES. Plant pools are shown with yellow boxes, soil pools are shown with slate colored boxes. Similar to the convention in Figure 1, this conveys new model content, in this case the introduction of nitrogen and phosphorus mass accounting in FATES. *Denotes the special status of nutrient storage, in that storage nutrient pools are distinct from the carbon storage pool. The leaf, fine-root, sapwood, structure and reproductive tissues assume that carbon, nitrogen and phosphorus are bound together in the tissues. **Fixation fluxes only refer to nitrogen. Leaching refers to phosphate and nitrate (not ammonium) and denitrification losses are only from nitrate. ***Myccorhizal uptake is not currently represented in the model, but may be in the future.

Three chemical elements are tracked and conserved by mass within the plant, litter and soil system: carbon (non isotope specific), nitrogen, and phosphorus. The masses of these elements at any given instance in time are designated $C_{(o)}$ for carbon and $M_{(o,s)}$ for the two nutrients elements, where o is the generic subscript for the organs and s is the generic subscript for the two nutrient elements nitrogen and phosphorus.

In the following model description, parameter constants are indicated by lowercase Greek letters. Some variables not associated with a mass use lower-case letters. Unless specified otherwise, fluxes and rates of change use an over-dot (i.e. \dot{X}). All plant states have units of [kg plant⁻¹]. Turnover and allocation within the plant occur at a daily frequency. Thus, they have units of [kg plant⁻¹ day⁻¹]. Nutrient competition between plants and soil competitors (e.g., microbes, mineral surfaces) is resolved at sub-diurnal timescales (typically 30 minutes), and is integrated over the day and presented as a daily uptake $[kg plant^{-1} day^{-1}].$

Each plant cohort is represented by an average individual that maintains discrete mass pools for the following organs (and associated sub-scripts): leaf (lf), sap-wood (sa), dead-wood (de), fine-root (fr), reproductive (re) and storage (so) (see Table 1).

Organ Name	Subscript Symbol
leaf	lf
fine-root	$^{\mathrm{fr}}$
live (sap) wood	sa
dead (structural) wood	de
reproductive	re
$storage^*$	SO

Table 1. The plant organs and their subscripts represented in a FATES cohort. *Storage is not technically an organ, but its sum throughout the plant is tracked.

168

Sapwood refers to all living woody tissues, including organs such as the cambium, 169 phloem, and xylem. Dead-wood refers to all non-living tissues such as heartwood and 170 bark. Both cases (dead and live) include below and above-ground components. The dead 171 pool should not be confused with the coarse woody debris associated with dead trees. 172 Fine-roots are functionally classified as tissues with high turnover and respiration rates, 173 as compared to below-ground sapwood (coarse roots). Reproductive organs encompass 174 all ephemeral tissues associated with reproduction, including seeds, cones, flowers, fruits, 175 etc. For storage, the term "organ" is used loosely because reserves are spatially distributed 176 throughout the plant, often in vacuoles, referring to all forms of C, N, and P that can 177 be re-mobilized for growth or maintenance of other tissues. Carbon storage refers to non-178 structural carbohydrates of starches and sugars. Storage of N is often comprised of pro-179 teins and amino compounds (Millard & Grelet, 2010), whereas phosphate compounds 180 are typical for phosphorus storage (S.-Y. Yang et al., 2017). 181

Plants represented in FATES can acquire nutrients through several means: 1) uptake of mineral nutrients from soil solution, 2) symbiotic nitrogen fixation, and 3) re-absorption preceding litterfall. In PARTEH, each function is designed to be modular and interchangeable with alternative hypotheses. Here we describe the default options for these uptake processes.

187

2.1 Acquisition through Symbiotic Fixation

The carbon cost of symbiotic nitrogen fixation is modeled as an obligate (temper-188 ature dependent) increase in maintenance respiration (Houlton et al., 2008). Plants that 189 are designated as nitrogen fixers generate a fixation rate M_f [kgN plant⁻¹ day⁻¹] by respir-190 ing carbon r_f [kgC plant⁻¹ second⁻¹] at a rate that is a constant fraction $\rho_{f(pft)}$ of all 191 non-fixation fine-root maintenance respiration (Ryan, 1991) (non-growth) costs r_m [kgC 192 $plant^{-1}$ second⁻¹]. This simplification assumes that all resources driving nitrogen fix-193 ation are mediated through respiration, and those costs act as a surrogate for other costs 194 such as building and maintaining nodules and feeding specialized bacteria. This repre-195 sents an obligate strategy because all plants of a Plant Functional Type (PFT) with a 196 nonzero value of $\rho_{f(pft)}$ constantly fix N and incur the respiratory cost of doing so. The 197 representation of facultative nitrogen fixation strategies in FATES is left for future work. 198

¹⁹⁹ The nitrogen fixation flux is accumulated on each sub-daily time-step (of duration $\Delta t =$

1800 seconds) over the total steps for the day t_d .

$$r_{f} = r_{m} \cdot \rho_{f(pft)}$$
$$\dot{M}_{f} = \sum_{t}^{t_{d}} r_{f} n_{f} \Delta t$$
(1)

The rate of nitrogen fixed per unit carbon respired, n_f [gN gC⁻¹], follows the functional form by (Houlton et al., 2008). Calibrated constants are taken from (J. Fisher et al., 2010) ($a_{f1} = -6.25$) and (Houlton et al., 2008) ($a_{f2} = -3.62$, $a_{f3} = 0.27$, $a_{f4} =$ 204 25.15) and the soil temperature T_{soil} is prognostic variable of ELM.

$$n_f = a_{f1} \left(e^{a_{f2} + a_{f3} \cdot T_{soil} \left(1 - 0.5 \cdot \frac{T_{soil}}{a_{f4}} \right)} - 2 \right)$$
(2)

2.2 Plant Acquisition of Aqueous Soil Nutrients

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In both supported nutrient competition schemes (CB and RD, see below for details), FATES cohorts compete with other cohorts, as well as microbes and mineral surfaces (for phosphorus) for aqueous nutrients in each discrete soil layer j. (Note that the CB scheme does allow for occlusion of ammonium and nitrate in clay soils) Plants compete for 1) ammonium (NH₄) with decomposer and nitrifier microbes, 2) nitrate (NO₃) with decomposer and denitrifer microbes, and 3) phosphate (PO₄) with decomposer microbes and mineral surfaces.

In the native "big-leaf" vegetation representation in ELM, each functional type com-213 petes for nutrients as a group. In contrast, FATES enables many cohorts of different sizes 214 and functional types, all to compete independently for resources with soil competitors 215 (typical cohort counts on a site can number anywhere from tens to more than a thou-216 sand, depending on local biodiversity and modeler decisions on how to delineate func-217 tional groups and size-similarity). Both CB and RD schemes require each cohort to pro-218 vide a potential uptake rate, or uptake capacity, for each mineral nutrient species $(\hat{M}_{u,NH_{d}(i)})$ 219 $\hat{M}_{u,NO_3(j)}$, or $\hat{M}_{u,PO_4(j)}$ units [kg m⁻² s⁻¹]). The actual net daily uptake flux $\dot{M}_{u(s,j)}$ [kg m⁻² day⁻¹] results from the competition schemes, which we denote with a generic 220 221 "competition function" $\Gamma_{c(j,t)}$. The Relative Demand (RD) (P. Thornton et al., 2007) 222 method distributes nutrient uptake to competitors in proportion to their demands. It 223 also provides controls to scale up or down the relative competitiveness of each entity when 224 total mineralized nutrients are less than total demand. If the total demand exceeds avail-225 ability, all uptake rates are down-scaled to ensure that the scheme does not generate neg-226 ative soil N and P concentrations. The Capacity Based (CB) (Zhu et al., 2019) method 227 utilizes a Michaelis-Menten approach to estimate the simultaneous uptake of compet-228 ing entities with half-saturation parameters, and is therefore influenced by the soil aque-229 ous nutrient concentrations. The CB model also accommodates phosphatase dynamics 230 following (Wang et al., 2010). The ELM-FATES modeling coupling does enable these 231 phosphatase dynamics, as well as resulting preferential phosphorus availability to the plants. 232 The exact form of the competition functions and details about the schemes are described 233 in Zhu et al. (2019). 234

FATES calculates plant growth and allocation on a daily basis, hence, the total daily uptake for each cohort includes the sum of the uptake over each of the total number of j_s soil layers and sub-daily time-steps (of duration Δt seconds) over the total for the day t_d .

$$\dot{M}_{u(N)} = \sum_{j}^{j_s} \sum_{t}^{t_d} \left(\hat{M}_{u,NH_4(j)} \cdot \Gamma_{c,NH_4(j,t)} + \hat{M}_{u,NO_3(j)} \cdot \Gamma_{c,NO_3(j,t)} \right) \Delta t$$
$$\dot{M}_{u(P)} = \sum_{j}^{j_s} \sum_{t}^{t_d} \left(\hat{M}_{u,PO_4(j)} \cdot \Gamma_{c,PO_4(j,t)} \right) \Delta t$$
(3)

The nutrient uptake capacity of a FATES cohort is defined by the per-plant fineroot biomass $C_{(fr)}$ [kg plant⁻¹], the plant density n_p [plants m⁻²], the fraction of fineroot biomass in each soil layer $f_{fr(j)}$ (see Section Appendix C), and the maximum uptake rate per unit fine-root biomass $\nu_{max(pft)}$. This parameter is unique to each mineral nutrient chemical species (NH_4, NO_3, PO_4) for each PFT represented by FATES [kg kg⁻¹ s⁻¹]. Cohort density and fine-root biomass are prognostic variables in FATES.

$$\hat{M}_{u,NH_4(j)} = \nu_{max,NH_4} \cdot C_{(fr)} \cdot n_p \cdot f_{fr(j)}$$

$$\hat{M}_{u,NO_3(j)} = \nu_{max,NO_3} \cdot C_{(fr)} \cdot n_p \cdot f_{fr(j)}$$

$$\hat{M}_{u,PO_4(j)} = \nu_{max,PO_4} \cdot C_{(fr)} \cdot n_p \cdot f_{fr(j)}$$
(4)

Note that for the RD approach, the nitrate uptake capacity $M_{u,NO_3(j)}$ is handled slightly different than equation 4. With RD, uptake for nitrogen happens sequentially. The uptake capacity for ammonium and nitrate are combined, and used to drive uptake first from the ammonium pool. This will fulfill some of the plant's needs, and reduce the joint uptake capacity. The remaining joint uptake capacity is then applied to draw down the nitrate pool.

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2.3 Losses and Re-acquisition During Turnover

FATES tracks the daily turnover from senescent tissues on live plants with a car-252 bon loss rate $\dot{C}_{t(o)}$ and nutrient loss rates $\dot{M}_{t(o,s)}$ [kg plant⁻¹ day⁻¹] for all non-reproductive 253 plant tissue pools: leaf, fine-root, sapwood, storage, and structural wood, for each co-254 hort. These turnover fluxes are non-episodic, and the rates are controlled by the turnover 255 period parameter $\tau_{(o,pft)}$ [years] associated with the plant's phenological dynamics, which 256 are PFT dependent. The storage, sapwood, and structural wood all share the same turnover 257 rate which is associated with branch-fall. A module that explicitly tracks damage lega-258 cies and represents degraded crowns exists (Needham et al., 2022), but is not used here 259 to reduce confounding model factors during analysis. 260

$$\dot{C}_{t(o)} = C_{(o)} / (365 \cdot \tau_{(o,pft)})
\dot{M}_{t(o,s)} = M_{(o,s)} / (365 \cdot \tau_{(o,pft)})$$
(5)

Plants re-absorb a portion of nutrients before leaf and fine-root tissues are shed during senescent turnover. This rate $\dot{M}_{ra(o,s)}$ [kg plant⁻¹ day⁻¹] is drawn from the turnover rate, is directed towards plant storage $M_{(so,s)}$, and is removed from the litter mass flux. There is no re-absorption during fire, and no re-absorption from wood tissues. This re-

absorption happens at a constant proportion for leaves $\omega_{lf(s,pft)}$ and fine-roots $\omega_{fr(s,pft)}$

 $[kg kg^{-1}]$ specific to each nutrient (N or P) and PFT. Plants with high re-absorption rates

will require less nutrient acquired from other sources, and will generate litter with lower

nutrient density per unit carbon. A description of how re-absorption rates are estimated is described in Section 3.1.

$$\dot{M}_{ra(lf,s)} = \dot{M}_{t(lf,s)} \cdot \omega_{lf(s,pft)}
\dot{M}_{ra(fr,s)} = \dot{M}_{t(fr,s)} \cdot \omega_{fr(s,pft)}$$
(6)

Litter mass nutrient fluxes from senescent turnover of live plants follow the same proportion rules and constants as carbon for how they are proportioned into the labile, lignin and cellulose litter pools. For more details, see the FATES technical manual (FATES-Development-Team, 2019).

2.4 Definition of Plant Organ Targets

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In FATES-PARTEH, plants grow (to the extent possible, as described below) their 275 organs to preserve observationally-constrained allometric relationships. As the plant grows 276 and increases in stature (defined by stem diameter at reference height d), these allomet-277 ric functions define a target carbon mass for each organ, $C_{(o)}$. The plant will always at-278 tempt to allocate resources such that mass in an organ matches the target (i.e. replace 279 what has been lost), before it attempts to further grow in stature. The method of defin-280 ing carbon targets in FATES remains unchanged from (R. A. Fisher et al., 2015; C. D. Koven 281 et al., 2020) for all organs except fine-root. Fine-root mass targets now vary as a func-282 tion of carbon and nutrient storage. This is a key new model mechanism introduced in 283 this work and is explained in Section 2.6. Details on carbon targets and allometry func-284 tions for the other organs can be found in the FATES technical manual (FATES-Development-285 Team, 2019). 286

Nutrient targets $\dot{M}_{(o,s)}$ for leaf, fine-root, sapwood, and structural wood seek to maintain a constant stoichiometry (i.e., constant P:C and N:C ratios), defined as parameter constants $\alpha_{(o,s,pft)}$ specific to each plant functional type, element (N or P) and organ.

$$\dot{M}_{(o,s)} = \alpha_{(o,s,pft)} \cdot C_{(o)} \qquad (o = lf, fr, sa, de)$$
(7)

Unlike the other organs, the nutrient to carbon ratio of the reproductive tissues and 290 storage (o = re, so) are not defined directly by parameter constants. FATES, like many 291 vegetation demography models, does not mechanistically resolve germination or other 292 processes of plants below a minimum recruitment size (Hanbury-Brown et al., 2022); 293 instead it assumes that a fraction of carbon flux allocated to reproduction emerges as 294 new recruits at some time later. We extend this approach to nutrients as well. The sto-295 ichiometry of reproductive tissues is set to match the nutrient to carbon ratios of a newly 296 recruited plant (i.e. a plant with the smallest trackable stem diameter $d = d_{min}$). This 297 approach means that only the nutrients that are needed to produce recruits with a known 298 stoichiometry are allocated to reproduction, and represents the optimal reproductive al-299 location stoichiometry that also satisfies mass conservation. FATES initializes newly re-300 cruited plants with no reproductive tissues, and they start "on-allometry" (i.e. when their 301 actual mass matches the allometrically defined target). 302

$$\alpha_{(re,s,pft)} = \frac{\sum \dot{M}_{(o,s)}}{\sum \dot{C}_{(o)}} \qquad (d = d_{min}, \quad o = lf, fr, sa, de, so)$$
(8)

The target nutrient storage $M_{(so,s)}$ is a special case, because it is not associated 303 with a specific tissue. Therefore the target is scaled $(\mu_{(s,pft)})$ based on the target nu-304 trient content of the leaves when "on-allometry". Alternative hypotheses are available 305 for users to test, allowing for storage capacity to scale off of any combination of other 306 organs (e.g., sapwood, fine roots). 307

$$\dot{M}_{(so,s)} = \mu_{(s,pft)} \cdot \dot{M}_{(lf,s)}$$
(9)

2.5 Plant Allocation and Mass Balance Accounting

Both the carbon and nutrient fluxes in the plant and soil systems are mass con-309 servative (i.e. all mass fluxes are accounted for and nothing is created or destroyed). Sup-310 plemental nitrogen and phosphorus are often added to the soil in the early years of a spin-311 up simulation. These effective 'accelerate' the accumulation of nutrient pools in the soil, 312 and the unresolved processes pf primary succession. These fluxes are tracked in the over-313 all balance as well. The FATES code performs mass-balance checks at both plant and 314 landscape (i.e. contains all disturbance history patches in each FATES site) scale every 315 day. The following rules are stated explicitly for nutrients M, but are also valid for car-316 bon C. The sum of daily allocated nutrient $M_{a(o,s)}$ over all organs, should equal the dif-317 ference between the plant's total gains for the day $\dot{M}_{g(s)}$ and losses due to exudation $\dot{M}_{e(s)}$. 318 The total change in mass over the course of the day $M_{(o,s)}$ is therefore the difference be-319 tween what is allocated $M_{a(o,s)}$ and lost in turnover $M_{t(o,s)}$. 320

$$\sum_{o} \dot{M}_{a(o,s)} = \dot{M}_{g(s)} - \dot{M}_{e(s)}$$
$$\dot{M}_{(o,s)} = \dot{M}_{a(o,s)} - \dot{M}_{t(o,s)}$$
(10)

(11)

For nitrogen, the daily gain $\dot{M}_{g(s=N)}$ includes a queous uptake $\dot{M}_{u(s=N)}$ and sym-321 biotic fixation M_f . As per the ELM soil biogeochemistry model, any nitrogen made avail-322 able by free-living fixers in the soil are assumed to be added directly to the aqueous NH_4 323 pool, and does not need to be explicitly included in this calculation. 324

$$\dot{M}_{g(s=N)} = \dot{M}_{u(N)} + \dot{M}_{f}$$

$$\dot{M}_{q(s=P)} = \dot{M}_{u(P)}$$

308

The model considers three phases for allocation. In each phase, the mass pool for 328 the daily gain M_q is reduced as portions of this mass are transferred into plant organs 329 $M_{(o,s)}$. This phase proceeds sequentially in this order, as described next: 330

- Phase 1: Replacement of Turnover 331
- Phase 2: Stature Growth 332
- Phase 3: Remainder and Overflow 333

2.5.1 Allocation Phase 1: Replacement of Turnover

In the first phase, replacement of tissues lost to turnover is controlled by a prior-335 itization scheme, whereby a user controlled parameter indexed by plant organ $\delta_{(\rho)}$ is used 336 to assign priority. Organs with the highest priority have a $\delta_{(o)}$ of 1, organs with the low-337 est priorities will have larger values of $\delta_{(o)}$. A priority of 1 indicates the organ of inter-338 est has the first opportunity, along with other organs with that priority, to replace losses 339 and thus increase the mass of the organ toward the allometric target (turnover losses shift 340 organ masses "off-allometry"). Other organs with incrementally increasing $\delta_{(q)}$ are then 341 allowed to replace losses while there is still mass in the daily gain pool M_q . The high-342 est priority organs ($\delta_{(o)} = 1$) have some special considerations to how they are applied 343 (see Appendix Appendix B). This flexible scheme reflects persistent uncertainty over the 344 prioritization of allocation by plants and in principle allows rapid hypothesis testing. 345

The amount of nutrient $M_{a(o,s)}$ (or carbon) sent to each organ is driven by the deficit between the actual element mass of the organ and its target mass $\dot{M}_{(s,o)}$. We define a set of organs $\Theta_{1(p)}$ (subscript 1 is for "phase-1" allocation") at priority level p, and the fraction of the total allocation demand that can be filled $f_{a(s)}$ for all organs in this priority level (bounded between 0 and 1). The allocation for each organ is simply its portion of the total replacement demand, scaled by the total replaceable fraction $f_{a(s,p)}$.

$$f_{a(s,p)} = \min\left(1, \dot{M}_{g(s)} / \sum_{\Theta_{1(p)}} \max\left(0, \dot{M}_{(s,o)} - M_{(s,o)}\right)\right)$$
$$\dot{M}_{a(o,s)} = f_{a(s,p)} \cdot \max\left(0, \dot{M}_{(s,o)} - M_{(s,o)}\right) / \sum_{\Theta_{1(p)}} \max\left(0, \dot{M}_{(s,o)} - M_{(s,o)}\right)$$
(12)

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With each successive allocation, mass is removed from the daily gain $M_{a(s)}$.

2.5.2 Allocation Phase 2: Stature Growth

If mass remains in all of the daily gain pools $(\dot{M}_{g(s=N)}, \dot{M}_{g(s=P)}, \text{ and } \dot{C}_g)$ following the replacement phase, the plant will grow in stature. The stem diameter will increase, the target masses of the plant organs (which are tied allometrically to diameter) will increase, and the mass of each organ will increase. All organs grow together as a group, but exceptions can occur for numerical reasons (this is discussed in Section Appendix D). This set of organs that are "on-allometry" are defined as set Θ_2 .

The first task is to determine which of the three elements (C, N or P) is in short-360 est supply and will limit growth. We do this by estimating the mean stoichiometric ra-361 tios of the potential new plant growth. Note that to get the relative proportions of new 362 mass allocated to the organs, we evaluate the derivative of the target carbon with re-363 spect to change in diameter d, $\frac{d\dot{C}_{(s,o)}}{dd}$, for the organs in set Θ_2 , which can be retrieved 364 the from allometric functions at the current stem diameter. The estimated mean stoi-365 chiometries (represented by the two summation terms in the brackets to the right of the 366 $M_{q(s)}$ term in Equation 13) are then used to transform the daily nutrient gain into an 367 equivalent carbon $C_{q,eq(s)}$ gain. 368

$$\dot{C}_{g,eq(s)} = \dot{M}_{g(s)} \left[\sum_{\Theta_2} \frac{d\dot{C}_{(o)}}{dd} / \left(\sum_{\Theta_2} \alpha_{(ft,s,o)} \frac{d\dot{C}_{(o)}}{dd} \right) \right]$$
(13)

The minimum of the actual carbon gain \dot{C}_g and the two equivalent carbon gain pools $\dot{C}_{q,eq(s)}$ then defines the carbon that is available for stature growth \dot{C}_{sg} .

$$\dot{C}_{sg} = \min(\dot{C}_g, \dot{C}_{g,eq(s=N)}, \dot{C}_{g,eq(s=P)})$$
(14)

The carbon fluxes into each plant organ are solved via a set of coupled differential 371 equations, conducted via numerical integration from bounds zero to C_{sg} for each organ 372 in set Θ_2 . The proportion of carbon gain directed to each organ is defined by the deriva-373 tive of the diameter-to-mass allometry functions, as a fraction of the sum of all deriva-374 tives in set Θ_2 . The derivatives of the target masses with respect to diameter $\left(\frac{d\dot{C}_{(o)}}{dd}\right)$ 375 are readily available by differentiating the allometry functions. These are coupled equa-376 tions because they are all simultaneously drawing down \dot{C}_{sg} together, and the deriva-377 tives are continuously changing as they grow. 378

$$\dot{C}_{a(o)} = \dot{C}_{a(o)} + \int_{0}^{\dot{C}_{sg}} \left[\frac{d\dot{C}_{(o)}}{dd} / \left(\sum_{\Theta_2} \frac{d\dot{C}_{(o)}}{dd} \right) \right] d\dot{C}_{sg}$$
(15)

To handle the allocation of nutrient gains, the same allocation rules from Phase 1 are applied here in Phase 2, using the updated carbon biomass of each organ just explained. Refer to Equations 7 and 12. It should be noted that this modelling hypothesis holds the stoichiometries of plant organs (aside from storage) to the values provided by the parameter constants $\alpha_{(ft,s,o)}$. Small deviations may periodically occur, but they will be corrected automatically by the nature of the algorithm.

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2.5.3 Allocation Phase 3: Remainder and Overflow

³⁸⁶ Daily gain pools $(\dot{M}_{g(s=N)}, \dot{M}_{g(s=P)})$, and $\dot{C}_g)$ that were not limiting stature growth ³⁸⁷ or replacement may still be available and must be allocated to storage or removed from ³⁸⁸ the plant. Unlike other pools, we allow storage to exceed the target, up to a maximum ³⁸⁹ "overflow" capacity that is based on the target and a user-defined PFT-level parame-³⁹⁰ ter constant $\mu_{ov(pft)}$. An overflow flux $\dot{M}_{ov(s)}$ captures this transfer from gains to stor-³⁹¹ age; carbon fluxes follow the same rules as nutrients and are omitted for simplicity.

$$\begin{split} \dot{M}_{ov(s)} &= \max(0, \min(\dot{M}_{(so,s)}(1 + \mu_{ov(ft)}) - M_{(so,s)}, \dot{M}_{g(s)})) \\ \dot{M}_{g(s)} &= \dot{M}_{g(s)} - \dot{M}_{ov(s)} \\ \dot{M}_{a(so,s)} &= \dot{M}_{a(so,s)} + \dot{M}_{ov(s)} \end{split}$$
(16)

If the storage overflow capacity is full and there are still gains (i.e. $M_{g(s=N)} > 0$) 392 that have not been allocated, the plant will exude residual nutrient $M_{e(s)}$ into the metabolic 393 (i.e. from labile sources, contains no lignin or cellulose) litter pool with vertical profile 394 fluxes proportional to the fine-root density of each soil layer. If excess carbon remains, 395 there are two options to get rid of the excess \dot{C}_e . The default method is to burn it off 396 as autotrophic respiration. Alternatively, users can also opt to exude the carbon with 397 the same partitioning rules along with the nutrients. As will be described in the next 398 section, this model features optimization process that seeks to balance uptake of carbon 399 and nutrients, which will also serve to minimize these excesses. This is evaluated in the 400 analysis. 401

2.6 Dynamic Fine-root Biomass Response

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Along with symbiotic relationships with nitrogen-fixing bacteria, plants also mod-403 ify their network of fine-roots to regulate uptake of mineralized nutrient (Forde & Lorenzo, 404 2001). Some plant and ecosystem models have utilized this behavior for some time (Thornley, 405 1995; de Kauwe et al., 2014), and a dynamic fine-root optimization scheme is detailed 406 here as well. This scheme seeks to adjust resource allocation above- and below-ground 407 in order for plant growth to be equally limited by carbon, nitrogen and phosphorus (Bloom 408 et al., 1985). If the resource limitations on growth are balanced, the relative amount of 409 carbon in storage (i.e. mass of carbon currently in storage $C_{(so)}$ divided by the target 410 amount of carbon storage $\dot{C}_{(so)}$) will match the relative amount of nutrient in storage 411 (i.e. mass of nutrient currently in storage $M_{(so,s)}$ divided by target amount of nutrient 412 storage $M_{(so,s)}$). Given the high complexity of within-plant signalling mechanisms that 413 govern allocation and growth of leaf and root tissues, we do not try to mechanistically 414 represent these processes, and instead aim to tractably represent with as few parame-415 ters as possible the net effects of these mechanisms via the optimality-based approach 416 developed here. 417

We quantify a plant's carbon to nutrient balance with the term f_{cn} , see Equation 17. The term is calculated for the two nutrient elements and takes the maximum, which represents the nutrient with lowest relative storage. A natural log transform is applied for several reasons: 1) the metric becomes centered on zero, where carbon limited plant is less than and a nutrient limited plant is greater than zero, 2) by not being a ratio, it can be averaged and/or smoothed, and 3) enables additive properties in the algorithm that optimizes fine-root (explained below).

$$f_{cn} = \ln \left(\max_{s}^{N,P} \left[\frac{C_{(so)}/\dot{C}_{(so)}}{M_{(so,s)}/\dot{M}_{(so,s)}} \right] \right)$$
(17)

Fine-root biomass is living tissue that both respires (where maintenance respiration r_m is a function of mass, nitrogen concentration in the tissue, and temperature, following (Ryan, 1991)) and requires continual replacement of losses. An increase in fineroot mass will therefore result in more respiration and lower carbon use efficiency per unit leaf area, but it will also increase capacity for acquiring mineralized nutrients in the soil (recall Equation 4). This is visualized in Figure 3. Thus, positive values of f_{cn} will drive fine-root growth, and negative values of f_{cn} will drive fine-root reductions.

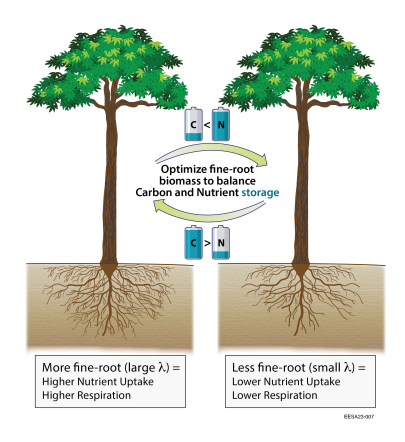


Figure 3. Visualization of the dynamic interaction between differential carbon-nutrient storage and fine-root growth. A plant (left) with proportionally more fine-root will tend to have decreased carbon allocation and increased nutrient allocation, than a plant (right) with proportionally less fine-root. The algorithm presented here seeks to balance these allocations through modifying fine-root growth. Illustration by Diana Swantek, Lawrence Berkeley National Laboratory.

⁴³³ In the FATES allometric model, the fine root target is defined by its proportion-⁴³⁴ ality λ with target leaf biomass $\hat{C}_{(lf)}$. The target leaf biomass is defined by the plant's ⁴³⁵ allometry (function of diameter), for details see the FATES technical manual (FATES-⁴³⁶ Development-Team, 2019).

$$\dot{C}_{(fr)} = \lambda \cdot \dot{C}_{(l)} \tag{18}$$

In previous versions of FATES, the proportionality λ between leaf and fine-root was a constant user specified parameter. Here, we create an algorithm where it is allowed to be dynamic, and it's value is optimized to result in a differential carbon to nutrient storage f_{cn} that tends towards zero. This system of carbon and nutrient regulation is summarized in Table 2

442 marized in Table 2.

437

condition		root response		plant response
high/positive f_{cn}	\rightarrow	$\lambda \uparrow$, $C_{(fr)} \uparrow$	\rightarrow	$r_m \uparrow$, $\dot{M}_u \uparrow$, CUE \downarrow
low/negative f_{cn}	\rightarrow	$\lambda \downarrow$, $C_{(fr)} \downarrow$	\rightarrow	$r_m \downarrow , \dot{M}_u \downarrow, \text{CUE} \uparrow$

Table 2. Table describing the plant's response to the f_{cn} storage metric. Relatively high and positive f_{cn} drives increases in the fine-root bimoass target, which drives increases in fine root biomass, which results in higher respiration (lower carbon use efficiency CUE) yet increased uptake. The reverse is true for low and negative f_{cn} status (ie proportionally high nutrient).

In early iterations of developing this hypothesis, we found that a linear model be-443 tween λ and f_{CN} was prone to over and undershooting an optimal solution, leading to 444 oscillations of λ in steady state climate conditions. To suppress the oscillatory behav-445 ior, we included the temporal derivative of f_{CN} . With this, the methodology became a 446 reduced form of a Proportional Integral Derivative (PID) "control-loop" system. In this 447 particular example, f_{cn} is the "process variable" which is driven by a "set-point" (λ). 448 PID controllers also contain an integral term along with the proportion and derivative 449 term. Each of the three terms is given a scaling coefficient, see Equation 19. The cal-450 ibration of the controller is discussed further in Section 3.3. 451

$$\lambda_t = \lambda_{t-1} + K_{p(pft)} f_{cn} + K_{i(pft)} \int f_{cn} dt + K_{d(pft)} \overline{\frac{df_{cn}}{dt}}$$
(19)

452

2.7 Software Features

The processes described here are encoded in a modular and extensible software struc-453 ture. It is modular because the software for the plant algorithms do not reference data 454 structures from the FATES (or other) model and uses a lightweight coupler to commu-455 nicate with FATES. This approach allows the plant model to be ported to any terres-456 trial biosphere model that uses a cohort or individual plant type of scaling approach. It 457 is extensible because the software is written so that other configurations of plant organs 458 (e.g. leaf spatial layering, storage pools with different functions, mycorrhizae, etc) and 459 chemical elements (e.g. Potassium, Magnesium, etc) can be readily adapted, if the user 460 can provide relevant parameter constants and the surrounding terrestrial biosphere model 461 can accommodate the boundary fluxes. Further, the FATES model code that processes 462 litter fluxes has been written to loop over the self-describing data structures for the chem-463 ical elements present (instead of explicitly defining new variable primitives for each mass 464 pool or flux associated with a chemical species). 465

466 **3** Model Calibration and Evaluation

An evaluation of the new model mechanics is performed via simulations at the Smith-467 sonian Tropical Research Institute's Barro Colorado Island (BCI) site in Panama. The 468 BCI site is conducive to evaluating a nutrient enabled terrestrial biosphere model be-469 cause it has an extensive 100 year history of ecological monitoring and analysis in ar-470 eas including forest demography and census (Condit et al., 2017), growth and mortal-471 ity (Wright et al., 2010), plant allometry (Martínez Cano et al., 2019; Cushman et al., 472 2021), nitrogen fixation (Wieder & Wright, 1995; Batterman et al., 2013), litter and soil 473 biogeochemistry (Mirabello et al., 2013; Yavitt et al., 2011; Yavitt & Wright, 2001; Pow-474 ers et al., 2005) and many more. It also stands out among tropical monitoring sites for 475

the long (> 30 year) and quality controlled meteorological data that is used to drive the FATES model (Patton, 2019a, 2019b, 2019c, 2019d; Faybishenko et al., 2018).

Over the course of our analysis, we found that evaluating both nitrogen and phos-478 phorus dynamics simultaneously expanded our scope beyond what can be covered in one 479 manuscript. We therefore decided to focus solely on evaluating nitrogen limitations, given 480 the following considerations: 1) the model mechanics for phosphorus within the plant 481 are almost exactly the same as with nitrogen with the exception of different parameter 482 constants; 2) nitrogen has a more complicated representation in the model because it has 483 two soil mineral pools and can be fixed by the plant; and 3) there is a companion paper evaluating phosphorus dynamics at a different site (Wei et al. in prep). While pre-485 vious research at Barro Colorado Island has found phosphorus and potassium limitations 486 to vegetation growth (Wright et al., 2011), we still find this an excellent site to evalu-487 ate the model mechanics in a carbon-nitrogen framework. This is because of wealth of 488 previously mentioned observations at the site, and also that previous experiments with 489 FATES are available to provide a solid calibration basis (C. D. Koven et al., 2020). Fur-490 ther, the objective here is not to make predictions, but rather determine if the model can 491 capture pattern responses in a test-bed that has realistic parameter constraints and bound-492 ary conditions. 493

Turning off phosphorus limitations is straightforward and achieved by 1) providing a supplementation term that feeds phosphorus directly to soil decomposers, plants, and mineral surfaces so that their nutrient demands are completely met and 2) using a plant uptake affinity parameter $\nu_{max(s=P)}$ that is excessively efficient (large). This results in the plants ignoring phosphorus effects on the fine-root biomass optimization, acquiring more than enough phosphorus for growth requirements and therefore releasing the excess back to the soil and litter.

501

3.1 Initial Parameter Calibration

A set of model parameter constants derived from previous research were used as 502 a basis for investigating the sensitivity and function of newly introduced parameters. C. D. Koven 503 et al. (2020) performed a parameter sensitivity analysis of the pre-existing carbon-only 504 ELM-FATES at BCI, where they generated an ensemble of 576 parameter combinations 505 to explore model response to twelve plant traits. Their model output was retrieved and 506 compared at different size classes to measurements of growth increment (centered at 7.5, 507 12.5 and 40cm), mortality rate (centered at 5.5 and 30 cm), and integrated total basal 508 area (< 30 cm, < 70 cm and all) (Condit et al., 2017). Only two size classes were used 509 for mortality (compared with 3 for basal area and growth increment), to compensate for 510 fewer data points (observations) associated with mortality. Scalar values of leaf area and 511 gross primary productivity (GPP) were also compared (Ely et al., 2019). This totals 10 512 values that can be compared: 10 = 2 size classes of mortality + 3 size classes of basal 513 area +3 size classes of growth increment +1 for leaf area and +1 for GPP. 514

For each ensemble member i and each of these 10 comparison points (subscript j), a difference between the observed and modeled $x_{(i,j)}$ were aggregated to a single fitness metric ϵ_i for each ensemble member, by summing the difference squared between the modeled and observed variables, divided by the variance of the difference across across ensembles. The parameter set associated with the simulation that minimized the fitness metric was used as a basis for simulations described here, a comparison of that parameter set with data is provided in Figure 4.

$$\Delta x_{(i,j)} = x_{obs(j)} - x_{(i,j)}$$

$$\epsilon_{(i)} = \sum_{j} \left(\Delta x_{(i,j)} \right)^2 / \sigma_{\Delta x_{(j)}}$$
(20)

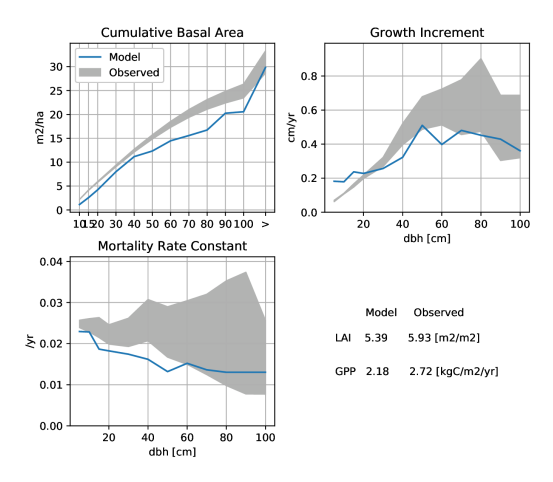


Figure 4. Comparison of observations with the most optimal ELM-FATES parameter set from the ensemble of simulations generated in (C. D. Koven et al., 2020).

Some of the parameters described in Table A1, organ stoichiometries $\alpha_{o,s,pft}$ and organ turnover rates $\tau_{o,pft}$, are also derived from the optimization of output (C. D. Koven et al., 2020). For nutrient enabled simulations, there are several new parameter constants that must be estimated. The methods for estimating the parameter constants for this study are explained here:

 $\omega_{fr(N,pft)}$ Evidence of root re-absorption of N and P is sparse, but has been observed in extratropical sites(Nambiar, 1987; Freschet et al., 2010). It is also believed that

if roots do not actively retranslocate nutrients before senescent turnover, some por-532 tion of nutrient in the newly made root litter will be made available for plant up-533 take by mycorrhizae. We assume a 0.25 fine-root re-absorption fraction of nitro-534 gen on senescence. Given this uncertainty, this modeling framework could be used 535 in further studies to better understand the sensitivity of different re-absorption 536 assumptions on ecosystem response. 537 $\delta_{(o)}$ We assign leaves and fine-roots the highest replacement priority, followed by stor-538 age and then sapwood and structural wood. We view other prioritization group-539 ings as alternative modeling hypotheses that can be explored in further study. 540 $\mu_{(N,pft)}, \mu_{ov}$ The parameter that controls the size of the nitrogen storage target $\mu_{(N,pft)}, \mu_{ov}$ 541 and how much storage overflow is allowed μ_{ov} is explored in experiment IV (see 542 Table 3 and Section F_2). Understanding plant nutrient storage is a difficult, but 543 there is some thought that plants store enough nutrient for seasonal use (Millard 544 & Grelet, 2010). As a base assumption, for all other experiments, we assume the 545 overflow is 100% of the target, and target nitrogen storage is 1-times the size of 546 total nitrogen bound in leaf tissues. 547 $\rho_{f(pft)}$ The maintenance respiration surcharge fraction for obligate symbiotic dinitro-548 gen fixation is explored in experiment VII (See Table 3 and Section 3.5. Symbi-549 otic fixation is turned off in all other experiments, and total ecosystem fixation 550 is used as a surrogate in those cases. 551 $\nu_{max(N)}$ Nitrogen uptake efficiency was viewed as a model calibration parameter, and 552

not something directly determined via field measurements. Although calculations 553 based on field inventory data provided a rough starting point. A spectrum of up-554 take efficiencies were tested and model output was evaluated for basal area com-555 parable to field observations (30 m2 ha^{-1}). For relative demand soil competition 556 mechanics, this resulted in a base value of $\nu_{max(N)} = 5e^{-9}$ [gN gC⁻¹ s⁻¹]. Sen-557 sitivity is evaluated in experiment VIII. The capacitance based competition nutrient scheme has a much smaller actual/potential uptake ratio due to a different 559 algorithm, and thus we arrived at higher values of $\nu_{max(NH4,NO3)} = 1.75e^{-7}$. Fur-560 ther details are provided in Experiment IX. 561

562 563 $K_{p(pft)}, K_{i(pft)}, K_{d(pft)}$ The calibration and sensitivity of the PID scaling parameters are covered in Experiment I and II (see Table 3 and Section 3.3).

564

3.2 Description of Experiments and Simulations

We conduct a series of experiments to elucidate the model's behavior and parameter spaces that focus on nutrient dynamics. A list of the experiments and the simulations used in each is provided in Table 3.

Some simulations are "spin-ups". In these simulations, FATES vegetation is ini-568 tialized with saplings (if more than 1 plant-functional type is present, the abundances 569 are equal). For the first 30 years of the simulation, nitrogen is added to the soils to ac-570 commodate the potential uptake capacity of the plants and microbes that is not met by 571 the existing aqueous nitrogen in the soil (i.e. nitrogen limitations are removed from all 572 competitors). After this 30 year phase, nitrogen then accumulates in the system through 573 the natural mechanisms of deposition and fixation (fixation is the dominant input in the 574 system and evaluated further). The modeled decomposition process in this phase uses 575 increased rate constants (often referred to as Accelerated Decomposition (P. E. Thornton & Rosenbloom, 2005)). The atmospheric CO_2 concentration in spin-ups is fixed at 577 pre-industrial levels (290 PPM). Eventually, the nitrogen and carbon content of the soils 578 reaches an equilibrium, as the vegetation evolves towards a mature demographic (quan-579 tified by a steady basal area distribution across size and functional types) and the lit-580 ter fluxes from the vegetation reaches a steady state. 581

The objective of several spin-up simulations were to provide an initial-condition for industrial-era simulations with transient CO_2 concentrations. In those, we determined that the spin-ups had reached steady-state because the log of the absolute value of Net Biome Productivity had reached very small values (approx 10^{-4} [kgC m⁻² year⁻¹]) (C. Koven et al., 2013).

Industrial-era simulations used normal (un-accelerated) decomposition rate constants, initialized size and age structure of vegetation from preceding spin-up simulations, and likewise initialized organic soil C and N pools using a multiplier of the values passed in from the preceding spin-up simulations. In some experiments where simulations were evaluated into the future, CO₂ concentrations follow from scenario SSP2-4.5.

All simulations utilized the 13-year (2003-2016) meteorological record from Faybishenko 592 et al. (2018) to provide ELM-FATES with rainfall, down-welling solar radiation, down-593 welling thermal radiation, atmospheric pressure, humidity, wind-speed and surface tem-594 perature. The simulations were all much longer than the meteorological record, so the 595 forcing was looped. Most experiments made use of the relative demand competition scheme, 596 solely because this approach is mathematically and conceptually simpler, which is help-597 ful in probing the complexities of the plant dynamics it is coupled with. The final ex-598 periment uses the capacitance based scheme because it was effective at maintaining aque-599 ous soil nutrient pools under high demand from competitors, as well as to see if its dy-600 namics were sensible and coupled correctly. 601

Experiment	Description	Competition	Period	Section
Experiment	Description	Scheme	Period	Section
Ι	Single cohort simulations	RD	150 year spin-up	3.3
	for controller sensitivity			
II	Competition based controller	RD	1000 year spin-up	3.3
	calibration	RD	·	
III	Evaluation of base	RD	500 year spin-up + 300	3.4
	parameterization	RD	year industrial-era	
IV	Sensitivity to storage capacity	RD	500 year spin-up	F2
V	Sensitivity to sub-module	RD	500 year spin-up	F3
V	hypotheses		·	
VI	Sensitivity to free-living	RD	500 year spin-up + 300	F4
	fixation hypotheses		year industrial-era	
VII	Evaluation of competition	RD	500 year spin-up	3.5
	between symbiotic fixers			
	and non-fixers			
VIII	Evaluation of interplay between	RD	500 year spin-up + 300	3.6
,	uptake efficiency $(\nu_{max(N)})$	10D	year industrial-era	0.0
	and soil N availability	CD	year industrial-era	~ -
IX	Evaluation of fine-root biomass	CB	500 year spin-up + 300	3.7
	estimates against field data		year industrial-era	

Table 3. List experiments, a brief description, the ELM nutrient competition scheme used (Relative Demand RD or Capacitance Based CB), the simulations periods used for each, and the section.

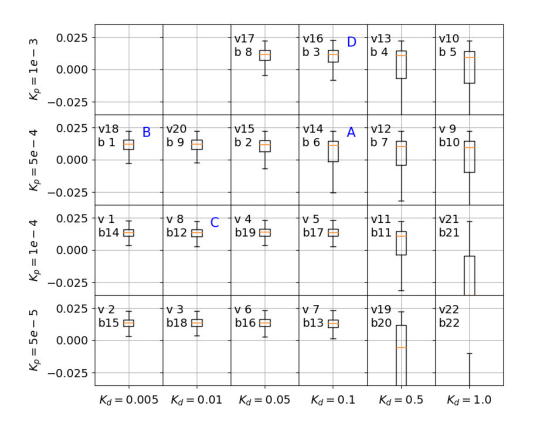
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3.3 Experiments I and II: Controller Calibration

The three terms in the PID control system serve distinctly different functions. The proportion term serves to push the process variable back towards its target value ($f_{cn} =$ 0) when there is a large difference between the current and target value. The derivative term serves to promote stability by suppressing rapid change in the process variable. The ⁶⁰⁷ integral term is most useful in reducing small and/or persistent biases between the pro-⁶⁰⁸ cess variable and its target; as the bias grows over time it will exert greater influence to ⁶⁰⁹ change the set variable. In testing the PID controller and exploring all three terms, we ⁶¹⁰ were able to achieve stable results without the integral term, so we decided to set it's ⁶¹¹ scaling constant K_i to zero for this study.

This system of nutrient cycling is fairly complex, with many plant and soil actors 612 competing for and cycling resources, all amongst changing meteorological conditions. The 613 plants are not experiencing a steady availability of nutrients for acquisition, and thus the 614 relationship between controller set point (root proportion λ) and the process variable 615 (storage ratio f_{CN}) are continually experiencing perturbations. To reduce the impact 616 of these perturbations in destabilizing the control system, we apply a multi-day smoother 617 to the derivative term. We use simple exponential smoothing where the future smoothed 618 value X_{t+1} is updated by the instantaneous value Y, the previous smoothed value X_t 619 and a weighting factor D synonymous with the number of time-points (days) over which 620 to weight the instantaneous variable: $X_{t+1} = X_t \cdot (1-1/D) + Y \cdot (1/D)$. In the experi-621 ments described here, we apply an D = 10 day smoother. We also tested 5 and 20 day 622 smoothing windows. Ultimately, reasonable controller response was found with all win-623 dows depending on the strength of the scaling constants. 624

In Experiment I, special reduced complexity simulations were conducted over a two dimensional log-scale grid search of the K_p and K_d terms from Equation 19. These reduced complexity simulations turned off recruitment and disturbance, which resulted in a simulation of a single plant cohort over a 100 year life-cycle trajectory. Each simulation was assessed for the variance and mean of the process variable $f_{CN(N)}$ (Figure 5), leaf to fine-root biomass multiplier λ (set point, Figure F1) and growth increment (an indicator of optimization of resource use, Figure F2).



 f_{CN}

Figure 5. Mean and variance of the relative storage of carbon to nitrogen variable (and Proportion Integral Derivative controller process variable) $f_{CN(N)}$, over a range of proportion and derivative controller settings. The values next to "v" and "b" indicate the ascending rank of each parameter couplet for bias and variance. Test v1 and b1 had the lowest variance and biases. Couplets designated A,B,C and D are used in the follow-up simulation.

The grid search shows that the model simulations are stable and viable (i.e. the 632 plants survive and can adequately adapt their root sizes to become productive) over a 633 large range of parameters (> 2 orders of magnitude each). However in Experiment II, 634 to determine which parameters offer a solution that is most in line with the optimality-635 based idea that underlies this approach, we create a simulation with four different func-636 tional types of plants that compete against each other for resources. A comparison of 637 their basal area trajectories, and the root proportion λ of newly recruited plants in open 638 and exposed sunlight are shown in Figure 6. The plant functional types have the same 639 parameters (i.e., traits) and initial seedling density (this is a spin-up style simulation) 640 with the exception of different PID controller constants. These parameter couplets are 641 labeled A, B, C, and D in Figure 5. 642

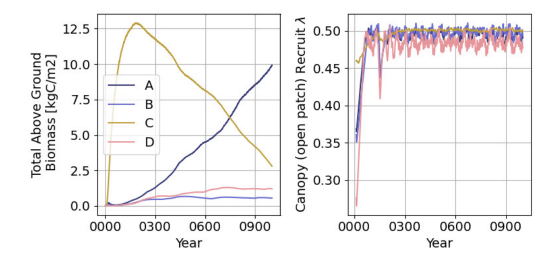


Figure 6. ELM-FATES spin-up simulation output containing four competing plant functional types, labeled A-D. These four PFTs have identical traits and parameters, with the exception of their proportion and derivative PID parameters. The values of their PID parameters are provided in 5. The right hand panel shows the mean root proportionality λ term for the different PFTs for open patch recruits. Patches that are open, do not have closed canopies and thus light availability to recruits. Recruit λ values are shown to avoid any differences associated with differences in size structure.

The result of Experiment II shows that the "C" PID parameter couplet was most 643 effective at rapidly adjusting to the competitive resource environment and homing in on 644 a reasonable λ value fairly quickly (see right panel of F1), thus initially occupying the 645 canopy. The "B" and "D" parameter couplets, while not dominating during any phase, 646 persisted through the simulation. The "A" PID parameter couplet eventually out-competes 647 "C" to dominate the canopy, and was ultimately chosen as a default set of PID constants 648 moving forward. The complex competition and coexistence dynamics of the four differ-649 ent parameter couplets prompts the question as to whether the responsiveness of a plant 650 to adjust to nutrient and carbon needs is a part of how plants navigate and find niches 651 in the multidimensional coexistence space, and this responsiveness might align with other 652 traits that determine successional dynamics. 653

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3.4 Experiment III: Towards an Optimality-Based Root Allocation Model

In experiment III, a single spin-up style simulation is conducted, using a single plant functional type, and the base set of parameters. The purpose of this experiment is to view the emergent stand structure of the simulated vegetation under the new nitrogen constraints and fine-root biomass optimization algorithm.

In Figure 7, size-structured estimates of Basal Area, Above Ground Biomass, and the leaf to fine-root proportion term λ partitioned into canopy and under-story plants, are projected across time. All plots show the signature of the spin-up simulation, where over the course of 400 years the trees grow into the larger size classes from saplings.

By the completion of the simulation, there is a fairly uniform distribution of basal area across size classes. There is also a signature of inter-annual variability at any given size. This feature emerges due to the internal dynamics of the cohorts, as similar cohorts fuse together, and grow from one size-classification to a larger one. Other simulations (not shown) were conducted that removed the inter-annual meteorological signal by loop ing a single year of data, and similar patterns emerged there as well. AGB dynamics are
 similar but with more strongly weighted towards larger trees, as they contain a greater
 proportion of the ecosystem-level biomass than basal diameter.

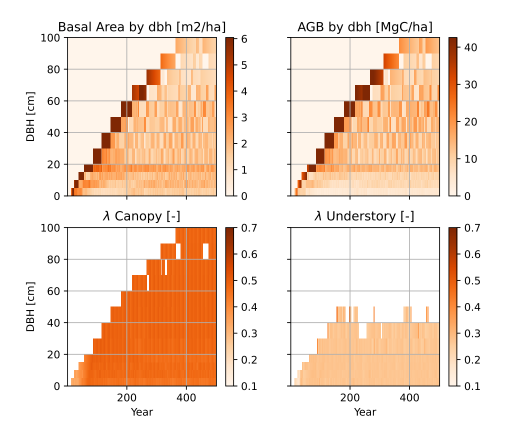


Figure 7. Evaluation of the size structure of vegetated biomass accumulation, and the size and canopy position structure of the fine-root proportion term λ [gC fine-root gC-1 leaf]. Canopy plants display distinctly larger λ values than understory plants at similar size classes.

Notably, the λ values in the canopy plants are distinctly and consistently larger than 671 those in the understory. Canopy plants have greater access to light and have increased 672 primary productivity compared to their understory neighbors. This increased produc-673 tivity places a greater demand on nitrogen acquisition to keep pace with more rapid con-674 struction of plant tissues. Moreover, the increased productivity of the canopy plants pro-675 vides adequate carbon reserves to pay for the increased respiration associated with more 676 fine-root biomass. Plants that are in the understory have limited access to light and sub-677 sequently lower productivity. Without the carbon to build new tissues there is a corre-678 spondingly lower demand on nutrients to match the construction costs of the carbon. This 679 potentially triggers a response in the plant to decrease investments (respiration costs of 680 fine-roots) in the acquisition of nutrients it doesn't need. 681

The massive biological diversity of tropical forests is often associated with a multidimensional competition space. Different species allocate their resources into different organs and traits, features that maximize their success in different niches. A classic example is the growth versus mortality trade-off (Wright et al., 2010). There are examples of species at BCI that bide their time in the understory for many years without growth, and then accelerate growth when light conditions change. This difference in strategy between understory and canopy plants is captured in ELM-FATES, and demonstrated by decreased proportions of fine-root biomass λ in understory plants.

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3.5 Experiment VII: Incorporating Cohort-Scale Symbiotic Fixers

In Experiment VII, we test if the symbiotic nitrogen fixation module can generate an expected ecosystem response. One way to test this is to see if a nitrogen-fixing plant functional type can coexist with a non-fixing functional type in simulation. In this experiment, the two plant functional types have the same traits with the exception of the symbiotic fixing parameters themselves. This test is a simple proof of concept, and ignores the multifaceted trait space that fixers and non-fixers may occupy.

By introducing symbiotic fixation to the simulation, the pre-existing total ecosystem fixation scheme (Cleveland et al., 1999) must be modified to only represent free-living fixation. This approach is similar to the approach used by CLM5 (Lawrence et al., 2020), which identifies that the original total ecosystem fixation rates estimated in Cleveland et al. (1999) projected low and high ranges of fixation. Here, we downscale the NPP-derived total ecosystem nitrogen fixation rate by a multiplicative scaling factor of 0.2.

For plants to achieve co-existence in this model configuration, the unit cost of sym-703 biotic fixation must be higher than the uptake of aqueous nitrogen at its potential rate 704 (i.e. uptake when aqueous nitrogen is abundant and no source side limitation exists), yet 705 must be lower than aqueous uptake under some amount of limitation. Otherwise, sym-706 biotic fixers would always be more efficient and out-compete non-fixers, not only when 707 nitrogen from the mineralized soil pool is limited. Menge et al. (n.d.) has made this type 708 of argument, pointing out that the energetic costs of breaking the triple bonds in N_2 and 709 supporting the nodules in symbiotic fixers are thought to be costly. The expectation is 710 that the two functional types will reach an equilibrium in their relative proportion, where 711 the symbiotic fixers will support a mineralized nitrogen pool to an amount where the car-712 bon costs of actual plant nitrogen uptake balance with that of fixation. 713

In ELM-FATES, we assess the unit carbon efficiency $[gN gC^{-1}]$ (inverse of cost) 714 for potential mineralized nutrient acquisition, as the sum of the potential uptake rate 715 of the plant $\nu_{max(NH4,pft)} + \nu_{max(NO3,pft)}$ [gN gC⁻¹ s⁻¹] (for RD based competition), 716 divided by the maintenance respiration and replacement costs of the roots $[gC gC^{-1} s^{-1}]$. 717 Note this is the steady state rate, and ignores the initial cost to grow the root. The unit 718 cost of fixation is directly quantified by Equation 2, (Houlton et al., 2008; J. Fisher et 719 al., 2010). In the base parameterization, the carbon efficiency for potential mineralized 720 uptake was lower than the fixation efficiency. We found that by increasing the total po-721 tential uptake rate $\nu_{max(NO3)} + \nu_{max(NH4)}$ and increasing the fine-root lifespan $\tau_{(fr)}$ 722 from 1 to 4 years, the potential mineralized uptake efficiency exceeded that of fixation, 723 see Figure 8. 724

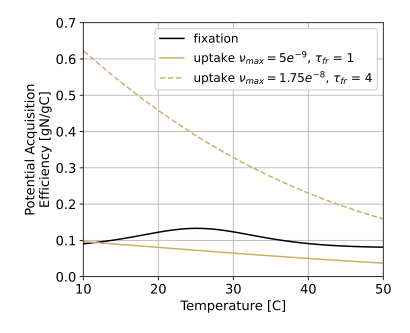


Figure 8. Comparison of the unit carbon efficiency for obligate symbiotic fixation (Houlton et al., 2008; J. Fisher et al., 2010) versus potential aqueous nitrogen uptake under the base parameterization and one with increased efficiency. Carbon costs for uptake efficiency consider maintenance respiration and replacement carbon costs, but not the initial investment.

The result of the test simulation is shown in Figure 9. The symbiotic-fixing PFTs were set to apply a 10% surcharge on fine-root maintenance respiration to fuel symbiotic fixation. This experiment also reduced the external N supplementation period from 30 to 5 years, to ensure that the symbiotic fixers had more control over the system N supply.

Fixers and non-fixers do show coexistence as demonstrated by their total aboveground biomass. The fixers have lower fine-root biomass fractions λ , which indicates their decreased need for mineralized nitrate and ammonium. Very early in the simulation, the non-fixers are more efficient due to the plentiful mineralized soil nitrogen. But after the supplementation period, plant mineralized nitrogen uptake becomes more limited (see bottom right panel), which then creates a competitive opportunity for the fixer PFT to emerge.

As a whole (considering both PFTs), symbiotic fixation accounted for about 5-10%737 of total plant acquisition, and slightly more than 50% of the total nitrogen fixed by the 738 ecosystem (including free-living). The latter is roughly close to what is expected, Batterman 739 et al. (2013) suggested that symbiotic fixation was the dominant mode of introducing 740 nitrogen to the site at BCNM, but within the same order of magnitude. The proportion 741 in the simulation could be increased by further regulating uptake efficiency parameters 742 or scaling down the free-living fixation rate. There are also different symbiotic fixation 743 temperature response functions available (Bytnerowicz et al., 2022), and future ELM-744 FATES testing may use these. 745

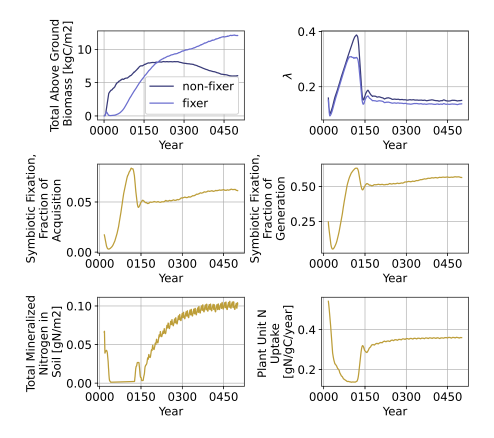


Figure 9. ELM-FATES simulation with coexistence between a fixer and non-fixer plant functional type. The symbiotic fraction of acquisition, refers to the fraction of plant acquisition that is from symbiotic-fixation, along with mineralized nutrient uptake. The fraction of generation looks at the relative contribution of symbiotic fixation to total nitrogen fixed by the system, also considering free-living fixation.

It takes about 500 years in this simulation for the fixer and non-fixer types to reach 746 an equilibrium. Yet, the results are an average of the whole landscape, and not a rep-747 resentation of how a single plot of land recovers from a disturbance. Batterman et al. 748 (2013) estimated that symbiotic fixation peaked near the first decade following a distur-749 bance, but after several decades total fixation flux dwindled considerably. This suggested 750 that symbiotic fixers play an important role in developing the nutrient environment in 751 newly disturbed lands, but perhaps they became less competitive as the nitrogen built 752 up in the soil and vegetation over time. The ELM-FATES model does have the ability 753 to simulate disturbance and discretely track land of different ages (called "patches") and 754 the plants that inhabit them. However in the current version of the model, the soil col-755 umn and its biogeochemistry (i.e. nitrogen and phosphorus concentrations in all forms) 756 are the same across patches of all ages, as the patch structure only represents hetero-757 geneity in the aboveground environment. In future versions of the model it would be in-758 teresting to see if the competitive dynamic of symbiotic fixers and non-fixers can be achieved 759 over the time-scales of decades where each patch has a dedicated nutrient environment 760 in the soil. 761

3.6 Experiment VIII: Sensitivity to Parameter Constants that Control Nutrient Availability and Affinity

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Experiment VIII tests model response to the intersection of three forces: nitrogen 764 availability, plant nitrogen use efficiency and increasing atmospheric CO₂ concentration. 765 Nitrogen use efficiency is modified by perturbing the plant functional parameter constant: 766 unit potential nitrogen uptake rate per mass of fine root (efficiency) $\nu_{max(pft,N)}$ [gN gC⁻¹]. 767 High values of $\nu_{max(pft.N)}$ (highly efficient) will acquire more nitrogen for less fine-root 768 carbon. Nitrogen availability is controlled by applying a constant multiplicative scaling 769 coefficient β to the NPP-based total ecosystem fixation function. The β parameter is not 770 listed in Table A1, and is not considered a component of this model because existing to-771 tal ecosystem fixation schemes were used in this manuscript and the scaling coefficient 772 is only used here to test sensitivity. Four parameter combinations are provided in Ta-773 ble 4. 774

	low plant	high plant	
	uptake affinity	uptake affinity	
low fixation	$\nu_{max(N)} = 5e^{-9}, \beta = 0.5$	$\nu_{max(N)} = 2.5e^{-8}, \beta = 0.5$	
high fixation	$\nu_{max(N)} = 5e^{-9}, \beta = 2.0$	$\nu_{max(N)} = 2.5e^{-8}, \beta = 2.0$	

Table 4. Parameter combinations for the four simulations in Experiment VIII. Each simulations used perturbations to only nitrogen acquisition efficiency $\nu_{max(pft,N)}$ and a scaling coefficient on total ecosystem fixation β . The red (compared to blue) indicates a higher nitrogen availability in the system. The darker shade (compared to lighter) indicates a higher plant uptake efficiency.

In all four simulations, there are several patterns that offer straightforward expla-775 nations. Increased productivity associated with higher CO_2 concentrations drive higher 776 basal area (biomass) and leaf area, as well as increased demand on nitrogen acquisition 777 to meet greater organ construction costs. Mineralized (aqueous) nitrogen depletes over 778 time, as the new additions to the system (fixation) can not keep pace with the increased 779 uptake. Faced with a greater need for (higher production) and a reduced supply of aque-780 ous nitrogen, the plants respond by increasing uptake capacity by building more fine-781 roots (λ). With decreased aqueous nitrogen and higher root mass, the unit uptake of ni-782 trogen per unit biomass decreases. The increase in CUE experienced by the plants is di-783 minished (and mostly reversed) when the mineralized nitrogen pools fully deplete, and 784 the plants are forced to respire newly assimilated carbon that cannot be used to build 785 tissues. 786

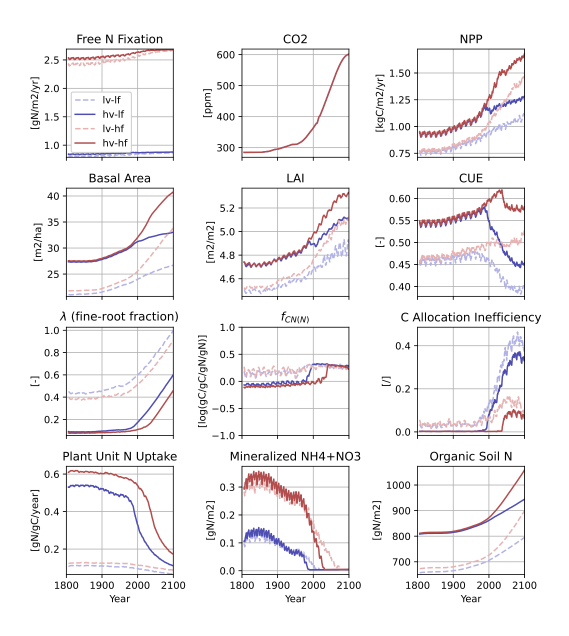


Figure 10. Time series model output for the four simulations described in Experiment VIII. Line colors match simulations described in Table 4, where: "lv-lf" is low-affinity low-fixation, "hv-lf" is high-affinity low-fixation, "lv-hf" is low-affinity high-fixation and "hv-hf" is high-affinity high-fixation". The CO₂ forcing signal uses observed industrial-era concentrations and follows the SSP2-4.5 scenario prediction to 2100. With no symbiotic fixation, Free N Fixation is synonymous with total fixation and is the primary source of nitrogen input. Carbon use efficiency (CUE) is the ratio of net primary production over gross primary production, where NPP accounts (i.e. subtracts) for any extra respiration of "excess carbon", that which couldn't be allocated due to nitrogen limitations. This extra respiration is also captured by "C Allocation Inefficiency", the fraction of grams of excess carbon burned per grams NPP. Plant Unit N Uptake refers to the grams of nitrogen uptake per gram of fine-root carbon.

Some patterns in the response are explainable, but less straightforward. Mineralized nitrogen depleted first in the low availability simulations, and also slightly earlier

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in the high versus low affinity simulations. This suggested that plants can be overly competitive for nitrogen, ultimately to their detriment. In simulations with higher uptake
affinity, the plants out-competed decomposers, which prevented the decomposers from
mineralizing nitrogen into the system, which leads to a slightly earlier collapse.

Another interesting pattern was the shift in the process variable $f_{CN(N)}$, from un-793 biased (i.e. closer to 0) to a bias indicative of a perpetually nutrient limited state. In the 794 high affinity case, the algorithm attempts to rectify the bias by increasing the λ value, 795 however the slowly dwindling supply of aqueous nitrogen continually counteracts the af-796 fect that increased fine-root fraction (set variable) should have on the process variable. 797 This raises the question: what is the appropriate time-scale of response for investing in 798 plant nutrient uptake? In these simulations, the PID constants that control λ were cho-799 sen for competitiveness in a constant CO_2 environment, but perhaps a parameterization 800 that favors more rapid adaptability would be more competitive in a changing CO_2 en-801 vironment. 802

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3.7 Experiment IX: Comparison of Fine-root Biomass Estimates with Field Observations

The objective of this experiment is to see if the new model formulation generates quantities of fine-root biomass comparable with observations in the field. In preliminary model simulations, estimated fine-root biomass was relatively low compared to observations. Decreasing nutrient uptake efficiency ν_{max} did generate larger fine-root proportions, however we learned from Experiment VIII that the increased respiration costs that come with higher root proportions also suppress net productivity and generated forests with less biomass.

There are various plant physiological model parameters that can be perturbed to increase carbon availability, in coordination with the decrease in nitrogen uptake efficiency. We chose to increase a parameter that directly impacts carbon assimilation, $V_{c,max}$ (catalytic capacity of Rubisco), from 30.9 to 55. And in increased a parameter that directly impacts carbon use efficiency, $\tau_{(fr)}$ (fine-root turnover timescale) from 1 to 3 years. We chose these because they were simple, powerful and directly related to net carbon acquisition.

Model output is compared to observed profiles by Yavitt et al. (2011), Yavitt and Wright (2001) and Powers et al. (2005) at BCNM (fig. 11). While the model captures the relative shape of fine-root biomass as a function of soil depth, the comparison shows us that the tendency of the model in this parameterization is to underestimate fine-root biomass. We discuss potential causes of the observed differences, and the interplay between carbon productivity, nutrient uptake efficiency and total nutrient availability in the following discussion.

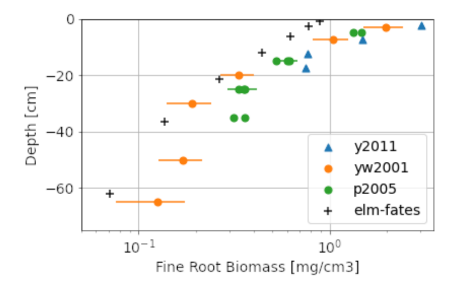


Figure 11. Comparison of ELM-FATES predicted fine-root biomass against observations at BCNM by Yavitt et al. (2011) (y2011), Yavitt and Wright (2001) (yw2001) and Powers et al. (2005) (p2005). All data was filtered to report only estimates of live roots from 0-2 mm diameter. Studies that did not differentiate between live and dead roots were corrected using the necromass to livemass ratio of 0.0806 (Yavitt & Wright, 2001). Data points with circles represent measurements on Barro Colorado Island, data points with triangles are on the other side of the river on the Gigante Peninsula.

4 Discussion

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4.1 Regulation of Nutrient Uptake

There are multiple avenues plants employ to regulate nutrient acquisition, which 828 include the fine-root growth response and obligate symbiotic fixation described in this 829 manuscript. Yet the uptake of nutrient by plants harbours complexity, including mod-830 ification of not just fine-root biomass, but morphology and structure (e.g., Taylor et al. 831 (2014)), considerations of advection and diffusion (e.g., McMurtrie and Näsholm (2018)), 832 symplastic and apoplastic transport (Steudle & Peterson, 1998), enzyme kinetics, sym-833 biotic relationships with facultative nitrogen fixers, algal nitrogen fixation on leaves, my-834 corrhizal associations and moreover acquisition of other nutrients not considered here 835 (K, Ca Mg, etc). 836

Employing dynamic fine-root response to nutrient gradients has both an established history in observation (Forde & Lorenzo, 2001) as well as model development (Thornley, 1995; Farrior et al., 2013; de Kauwe et al., 2014). This method differs from the Thornley models in how fine-root growth is controlled, where they tracked the substrate (C and N) concentrations in the roots and shoots dynamically. So while it is clear that certain plants do modify fine-root growth in response to nutrient availability, the signaling and what underlies these controls is less understood.

Several models also down-regulate mineral nutrient uptake in a more facultative form, outside of signaling increases or decreases in fine-root biomass (Thornley, 1995; Kou-Giesbrecht et al., 2021b; Thum et al., 2019; Zhu et al., 2019). Thornley (1995) called this downregulation product inhibition, and based it on N concentrations in the roots. Kou-Giesbrecht et al. (2021b) identified it as nitrogen stress, and tied it to the deficit of actual to target non-structural nitrogen content in the plant. Thum et al. (2019) also downregulates uptake based on "internal-demand" which also assesses the labile (nonstructural) nutrient content in the plant against the nutrient content of roots and leaves.
The ELM-FATES nutrient cycling model does not include this facultative downregulation, for two reasons. The first is that it was not clear how to represent this economically by associating a cost to this regulation. Secondly, an imperative was placed on minimizing the accumulation of parameters with the interest of using nutrient limitations
in global simulations and their calibrations.

Looking towards future model development and hypothesis testing of active acqui-857 sition, uptake capacity could be associated with dynamic activity levels of the fine roots, 858 as well as the amount of fine-root biomass or their surface area. While many of these 859 processes about to be mentioned are implicitly captured in the work presented here, through 860 maximum uptake capacity of roots and the Michaeles Menten half-saturation constants, 861 their explicit representation could be illuminating. For instance, dynamic activity could 862 be described as enzymatic activity rates, which are tied to nutrient content, the produc-863 tion of exudates to prime decomposition (via mycorrhizae for instance) or to chelate nu-864 trients, and/or the construction of specific infrastructure necessary for transferring nu-865 trient across the root surface. This activity could be controlled explicitly by resource in-866 vestments from the plant (e.g. respiration, carbon and nutrient allocations) and constrained 867 (albeit not explicitly governed or proportional to) by fine-root surface area. Root nutri-868 ent uptake could be made more realistic by considering the diffusion and mass transport of nutrients in the soil, root surface area and root architecture using the model of McMurtrie 870 and Näsholm (2018). 871

A plant-hydraulics model would likely make water transport of nutrients to the root 872 surface more accurate. Further, the fine root allocation approach presented here does 873 not consider the dual need for roots to provide both nutrients and water to plants, and 874 thus future work will develop this allocation model to optimize both water and nutri-875 ent delivery. For instance, the FATES-HYDRO model evaluates fine-root surface area, 876 hydraulic gradients between root surfaces and the soil and the conductivity across those 877 gradients to drive water fluxes to and across the fine-root surfaces. FATES-HYDRO also 878 has a basic representation of root structure that could support developments of the Mc-879 Murtrie model. 880

This model formulation has not explicitly incorporated mycorrhizal activity or its 881 affects on nutrient availability. In a sense, the effects of mycorrhizae are implicit or sub-882 sumed in the soil decomposition and nutrient competition schemes. However, without 883 explicitly representing mycorrhizae, it is impossible to capture the symbiotic benefits of 884 the association with the plants alone, and not just the broader affects of releasing min-885 eralized nutrient to the soil system. It would be interesting to incorporate and test hy-886 potheses of explicit mycorrhizal interactions, and their effects on nutrient cycling. In sum-887 mary, ELM-FATES-CNP uses the processes of fine-root biomass growth and maintenance as a surrogate for representing a broader and more complex set of functions related to 889 nutrient acquisition and regulation. Simulating a more complex plant response to nu-890 trient uptake needs may also be more consistent with phenomena observed at BCNM, 891 as fine-root biomass response to nitrogen fertilization is insignificant compared to that 892 of potassium (Yavitt et al., 2011). 893

However calibration and uncertainty in earth system simulators has become an ever 894 increasing challenge as all facets of the models steadily become more complex, with more 895 tune-able parameter constants and greater process uncertainty. It was imperative that 896 this formulation struck a balance between the desire to represent numerous complex pro-807 cesses and the desire to have a stable model that uses a reasonably small number of pa-898 rameter constants (particularly those that cannot be directly retrieved from measure-899 ments). Without these concessions, terrestrial biosphere models cannot be extended be-900 yond a handful of measurement-rich testbed sites. 901

In light of this, field experiments that uncover the when and why's associated with different modes of nutrient uptake regulation, their relative impacts on plant nutrient budgets, as well as the signaling from the plants that govern would be valuable.

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4.2 Fine-roots at the Nexus of Productivity, Resource Availability and Acquisition Efficiency

Potential nutrient uptake in ELM-FATES is dictated by the amount of fine-root 907 biomass and the maximum uptake rate parameter family ν_{max} , recall Equation 4. As 908 demonstrated in Experiment VIII, small (inefficient) values of ν_{max} drive larger fine-root 909 proportions (λ), and large (efficient) values of ν_{max} drive smaller fine-root proportions 910 λ . The response of λ was also impacted by total system nutrient availability (tested in 911 Experiment VIII by scaling total N addition to the system via community N fixation), 912 and how carbon productive the plants are (See pertubation to catalytic capacity of Ru-913 bisco $v_{c,max}$ in Experiment IX). This model has shown an interplay between the processes 914 of how carbon productive and efficient plants are and how nutrient productive and ef-915 ficient plants are. The fine-root biomass is a manifestation of these processes, and ide-916 ally this would make a great point of model calibration. 917

In theory we could continue to decrease the uptake efficiency parameter ν_{max} un-918 til we achieve comparable estimates of fine-root biomass. This can not be done in iso-919 lation, as to support more fine-root biomass with a similar stand-structure, the plants 920 would need greater productivity and would require a holistic calibration exercise that 921 also looks at net carbon productivity parameters, such as those controlling organ turnover 922 rates, respiration rates and gross primary productivity. However, we take the compar-923 ison of fine-root biomass with field data (see Figure 11) with a grain of salt, and see it 924 more as a rough guide than a validation point. There are several reasons for this. Firstly, 925 we have made the point that the representation of fine-root biomass in this model, and 926 it's response, is a surrogate for other processes (and their associated carbon costs) that 927 are not represented and would harbour increased complexity and uncertainty. 928

The second point, is that there is a difference between how the FATES model dif-929 ferentiates a fine-root from other tissues, and how fine-roots are differentiated in the field. 930 FATES has a functional definition, and differentiates fine-roots as tissues that respire at 931 a higher rate than coarse root or below-ground sapwood. Also FATES makes no differ-932 entiation of absorbing versus transporting fine-root tissues, or how how those tissues respire, 933 turnover or impact nutrient cycling (McCormack et al., 2015). All fine roots are treated 934 as absorbing tissues for the purposes of nutrient uptake. At the same time, the model 935 applies a single fine-root lifetime for the purposes of calculating the costs and benefits 936 of fine root allocation. Alternatively, fine-roots are typically differentiated in the field 937 by size (diameter) and order (Iversen et al., 2017), and efforts to reconcile fine root biomass 938 and ¹⁴C isotopic ratios have demonstrated two distinct populations of fine roots with widely-939 varying turnover times (Gaudinski et al., 2010; Ahrens et al., 2014). In this respect, there 940 is somewhat of a disconnect between the meaning of fine-root represented in the model 941 and how fine-roots are currently conceptualized to influence plant and ecosystem func-942 tion. However making a stronger better aligned comparison between what the model de-943 fines as fine-root, or what sub-classifications it has, and what is measured in the field, 944 would be useful for verification and calibration. In future work, we intend to represent 945 different fine root functional populations and associated turnover times to better rep-946 resent the joint constraints of nutrient uptake rates, root biomass profiles, and root 14 C 947 isotopic ratios. 948

949 5 Conclusions

The series of experiments presented here has demonstrated that this model framework can generate sensible patterns of ecosystem response, using a modest parameter

constant calibration effort. To summarize: 1) a small grid search of PID constants K_d 952 and K_p rendered values that enable the model to adapt stable fine-root biomasses with 953 reasonable levels of nutrient and/or carbon efficiency losses, 2) perturbations to param-954 eters that control nutrient storage $\mu_{\mu}\mu_{ov}$ did not exert undue model instability or vari-955 ability, and 3) subtle differences in how the model culls unnecessary roots and removes 956 unused carbon showed modest differences in model output. The number of newly intro-957 duced and salient (to nutrient cycling) model parameters that aren't readily derived from 958 field meausurements (i.e. stoichiometry α , and leaf reabsorption ω_{lf}) that exerted strong 959 control on model response is small (namely, ν_{max}). Balancing model complexity with 960 model robustness and preventing over-calibration is of critical importance and has been 961 identified as a key need in land-surface modeling endeavors (Prentice et al., 2015). 962

The new model hypothesis captures a few simple yet important concepts. Nutri-963 ent acquisition requires resources and that the construction of plant biomass is limited 964 by the acquisition of nutrients. In this case, the payment is the maintenance respiration, 965 construction and turnover replacement cost of the fine-roots. The current model hypothe-966 ses can also work with existing hypotheses in free-living and symbiotic nutrient fixation. 967 Finally, the dynamicism of fine-root proportion allows for a new competitive niche, where 968 understory plants have a new method to conserve resources when there is low access to 969 light and productivity. 970

Symbol	Description	Units
State Varia	ables	
$C_{(o)}$	carbon mass	[kg]
$\hat{C}_{(o)}$	target carbon mass	[kg]
$M_{(o,s)}$	nutrient mass	[kg]
$\dot{M}_{(o,s)}$	target nutrient mass	[kg]
d	reference stem diameter	[cm]
λ	leaf to fine-root target biomass multiplier	[-]
f_{cn}	the relative storage of carbon over the relative	
	storage of nutrient, for the maximum	
	(more limited) of nitrogen and phosphorus	[-]
External a	nd Diagnostic Variables	
f_{trim}	canopy trim fraction	[-]
n_p	the number of plants in a cohort per square meter	$[\text{plants m}^{-2}]$
$f_{fr(j)}$	the fraction of fine-root biomass in each soil layer	$[\mathrm{kg} \ \mathrm{kg}^{-1}]$
Fluxes		
$\hat{M}_{u,NH4(j)}$	plant ammonium uptake capacity in each soil layer	$[{\rm kg} {\rm m}^{-2} {\rm s}^{-2}]$
$\hat{M}_{u,NO3(j)}$	plant nitrate uptake capacity in each soil layer	$[{\rm kg} {\rm m}^{-2} {\rm s}^{-2}]$
$\hat{M}_{u,POx(j)}$	plant phosphate uptake capacity in each soil layer	$[kg m^{-2} s^{-1}]$
$\dot{M}_{u(s)}$	daily uptake of mineralized soil nutrients in solution	$[kg day^{-1}]$
$\dot{C}_{g}^{u(s)}$	daily carbon gain	$[kg day^{-1}]$
$\dot{M}_{g(s)}$	daily nutrient gain	$[kg day^{-1}]$
\dot{M}_{f}	daily nitrogen gained through symbiotic fixation	$[kg day^{-1}]$
$\dot{M}_{e(s)}$	excess nutrient exuded back to soil	$[kg day^{-1}]$
\dot{C}	daily carbon lost to turnover	$[\text{kg day}^{-1}]$
$\dot{C}_{t(o)}$		$[\text{kg day}]$ $[\text{kg day}^{-1}]$
$\dot{M}_{t(o,s)}$	daily nutrient lost via turnover	
$M_{a(o,s)}$	daily nutrient net allocated	$[\text{kg day}^{-1}]$
\dot{r}_e	excess respiration of unusable carbon	$ [kg day^{-1}] [kg day^{-1}] $
$\frac{\dot{r}_f}{\text{Parameter}}$	respiration cost to fix Nitrogen	[kg day]
	nutrient stoichiometric target for	
$\alpha_{(o,s)}$	non-labile tissue, nutrient mass per carbon mass	$[\mathrm{kg \ kg^{-1}}]$
τ_{i} .	non-mortal turnover timescale of plant organs	[years]
$\tau_{(o,pft)}$	* leaf re-absorption fraction of nutrient on turnover	$[\text{kg kg}^{-1}]$
$\omega_{lf(s,pft)}$	* fine-root re-absorption fraction of nutrient on turnover	$\left[\text{kg kg}^{-1} \right]$
$\omega_{fr(s,pft)}$	* proportion of target nutrient stored	[kg kg]
$\mu_{(s,pft)}$	per target nutrient in tissues	$[\mathrm{kg} \ \mathrm{kg}^{-1}]$
	* fractional overflow of storage (all chemical species)	[ng ng]
μ_{ov}	the plant will hold before exuding or respiring	[-]
	* maintenance respiration surcharge fraction	[]]
$\rho_{f(pft)}$	for obligate symbiotic dinitrogen fixation	[-]
\mathcal{U}	* maximum nutrient uptake demand per fine-root biomass	$[kg kg^{-1} s^{-1}]$
$ \begin{aligned} \nu_{max(s)} \\ \delta_{(o)} \end{aligned} $	* allocation priority	[index]
K (π)	* proportion term scaling parameter in PID controller	[IIIdex] [-]
$K_{p(pft)}$	* integral term scaling parameter in PID controller	[-]
$\frac{K_{i(pft)}}{K_{d(pft)}}$	* derivative term scaling parameter in PID controller	[_]
Table A1.	Non exhaustive list of variables and parameter constants in the FA	

971 Appendix A Table of Variables and Parameters

Table A1. Non exhaustive list of variables and parameter constants in the FATES nutrient cycling model. All mass and mass fluxes are assumed to be "per plant" [plant⁻¹]. External variables refers to those variables that are resolved by FATES processes outside the scope of this manuscript and are described in the FATES technical manual. Parameter constants denoted with * are newly introduced in this study. PID stands for proportion integral derivative, and is the controller used to search for optimal fine-root biomass.

⁹⁷² Appendix B Special Plant Organ Allocation Priority Levels

⁹⁷³ During the first phase of plant nutrient and carbon allocation, where allocation seeks ⁹⁷⁴ to replace losses due to continuous turnover (i.e. maintenance replacement), different or-⁹⁷⁵ gans can have higher replacement priority $\delta_{(o)}$ than others.

Priority level $\delta_{(o)} = 1$ has special status. Similar to previous carbon-only versions of FATES (C. D. Koven et al., 2020; R. A. Fisher et al., 2015). Organs at this level are allowed to draw from storage to replace a fraction of their replacement needs. The model will first attempt to use the daily gains and will resort to using storage if daily gains are not available. This allows for different species of plants to have more or less aggressive strategies.

Priority level $\delta_{(o)} = 2$ also has special status, and is reserved solely for replacement of storage. Please refer to the FATES technical manual.

In this study, we follow an organ prioritization similar to previous FATES model studies, where leaves and fine-roots are given the highest priority level (level 1), storage is given second priority, sapwood is the third, and structural wood is the fourth priority

⁹⁸⁸ Appendix C Vertical Fine-root Profile

The vertical attenuation of the fine-root fraction follows a two parameter exponential scaling model based on the depth of the upper (z^+) and lower (z^-) edge of the layer (Oleson et al., 2013). This fraction sums to unity and has no bearing on how much fineroot is present. The mass of fine-root is defined by allometric equations and the growth model (both described later).

$$f_{fr(j)} = 1/2 \left(e^{-7z_{(j)}^+} + e^{-z_{(j)}^+} - e^{-7\bar{z}_{(j)}^-} - e^{-\bar{z}_{(j)}^-} \right)$$
(C1)

Appendix D A Note About Growth Allocation and Numerical Integration

As mentioned earlier, it is typical that during the second phase of growth and al-996 location "stature growth", all organs will be "on-allometry" as indicated by their masses 997 being equivalent to the allometrically derived target mass for the plant's diameter. It is possible that some organs, due either to the process of cohort fusion (see FATES tech-999 nical manual) or numerical integration truncation, will not have masses that exactly match 1000 the target. In other words, the scheme will slightly overshoot or undershoot the target 1001 mass. These differences are both very small, and immediately corrected in this scheme, 1002 preventing the mass of the organs from drifting away from the allometric targets (in sit-1003 uations where the plant is not in carbon deficit). This self correcting methodology is part 1004 of the existing carbon-only FATES model as well. Organs that have less mass than their 1005 target will be corrected on the next day's Phase 1 allocation, see the "max" functions 1006 in Equation 12. Organs that have masses larger than their target are exempt from Phase 1007 2 allocation, and will be re-added to the list of organs in later iterations when the other 1008 organ masses have caught-up. That is why the set of organs $Theta_2$ does not always match 1009 the total set of non-reproductive organs. 1010

The numerical integration can be handled by either an adaptive Euler or an adaptive Runge-Kutta-Fehlerg 4/5 integration scheme. Because the numerical integration errors (either overshooting or undershooting the target mass of an organ) have relatively small consequence due to the self correcting nature of the scheme, we therefore default to the Euler integration scheme in all FATES allocation integrations, and retain the RKF4/5 for experimental purposes.

Appendix E Complementary analysis of the carbon-only model calibration phase

Measurements published by Ely et al. (2019) on leaf N:C ratio and SLA were used to filter the set of model simulations in C. D. Koven et al. (2020), by imposing that only simulations with parameter sets that fell within the 15-85 percentile bounds, see Figure E1. The histograms of other key carbon-only model parameters in C. D. Koven et al. (2020) ensembles are also shown, see Figure E2.

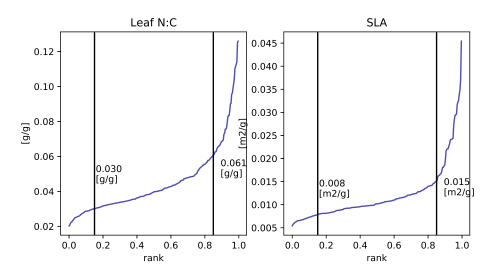


Figure E1. Rank plots of leaf nitrogen:carbon ratios and specific leaf area measured in Panama. Measurements by Ely et al. (2019). Vertical lines indicate the bounds of the 15-85 percentile.

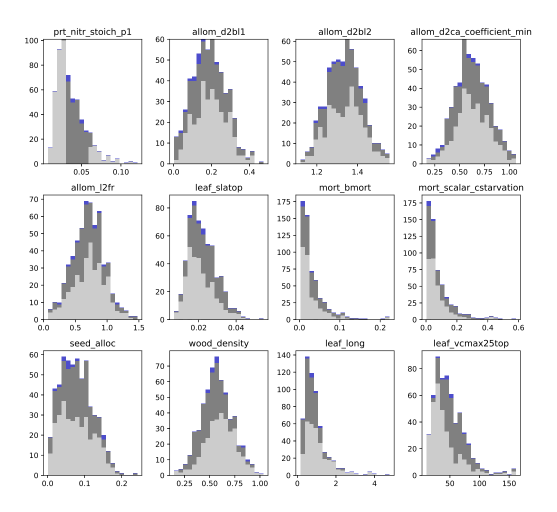
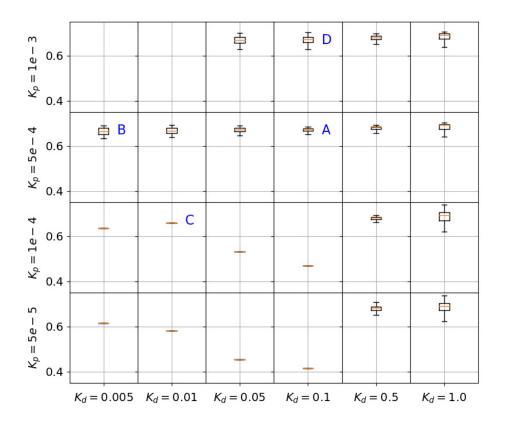


Figure E2. Histograms of model parameters from the parameter perturbation study of (C. D. Koven et al., 2020). Dark grey points represent parameters drawn from ensembles with leaf C:N ratios that fall within central 15 - 85% ranks in Figure E1. Blue points represent a small selection of ensemble members with best agreement with the census data.

- ¹⁰²⁴ Appendix F Experiment Analysis Expanded
- ¹⁰²⁵ F1 Supplemental Proportion Integral Derivative Controller Analysis



λ

Figure F1. Mean and variance of the leaf to fine-root biomass multiplier variable (and PID set point) λ , over a range of PID proportion and derivative controller settings. Settings with letter designations are used in a follow-up simulation designed to identify a more optimal parameter couplet. PID stands for proportion integral derivative and is the fineroot biomass optimization controller.



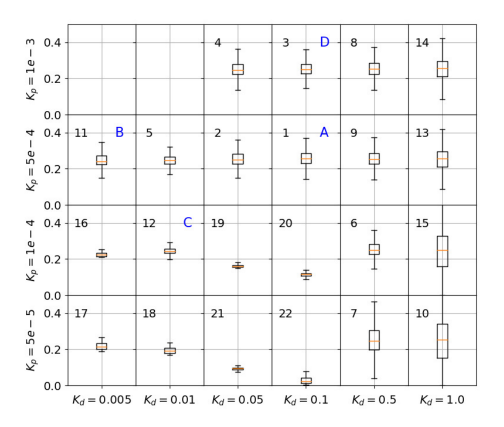


Figure F2. Mean and variance of the mean growth increment, over a range of PID proportion and derivative controller settings. Settings with letter designations are used in a follow-up simulation designed to identify a more optimal parameter couplet. PID stands for proportion integral derivative and is the fineroot biomass optimization controller.

1026 1027

F2 Experiment IV: Sensitivity to non-PID parameter constants that indirectly control fine-root growth

Since λ is driven directly by the differential nutrient storage ratios $f_{NC(s)}$, we focus on newly-introduced parameters that directly govern this differential, which includes the relative capacity the nutrient storage pools $\mu_{(s,pft)}$ and the allowable overflow of storage for all species μ_{ov} . While it would be interesting to investigate all correlations that exist between parameters, as well as how exiting parameters such as leaf photosynthetic traits govern this allocation, we leave that to future study.

In Experiment IV, we compare three simulations: the base simulation, one where 1034 the target N storage $\mu_{(N,pft)}$ is halved, and one where storage overflow μ_{ov} is halved. Time-1035 series of model indicators are compared, see Figure F3. In summary, major perturba-1036 tions to nutrient storage and overflow do not qualitatively change model results, and more-1037 over, the changes are explainable. Changes in NPP, Basal Area, Carbon Use Efficiency, 1038 1039 total nutrient uptake, organic nitrogen in the soil and aqueous nitrogen were also relatively unaffected. The largest changes were seen in those variables immediately con-1040 nected to the parameters, such as differential storage. 1041

With a smaller overflow capacity μ_{ov} , plants that acquire more nitrogen or carbon than required to fulfill construction costs, will be forced get rid of both at smaller tolerances. This was true for both species (see carbon use and nitrogen use inefficiency). However, halving overflow still generated a fairly high N allocation efficiency (i.e. plants were dumping less than 0.2% of what they acquired due to over-asking) and had small impacts on CUE.

A smaller nitrogen holding capacity $\mu_{(N,pft)}$ made plants slightly more net productive with higher carbon use efficiency, driven mostly by smaller allocations (and therefore respiration costs) to fine roots. With smaller holding capacity, a plant is more likely to fill up stores with excess nitrogen, which would signal to decrease root growth.

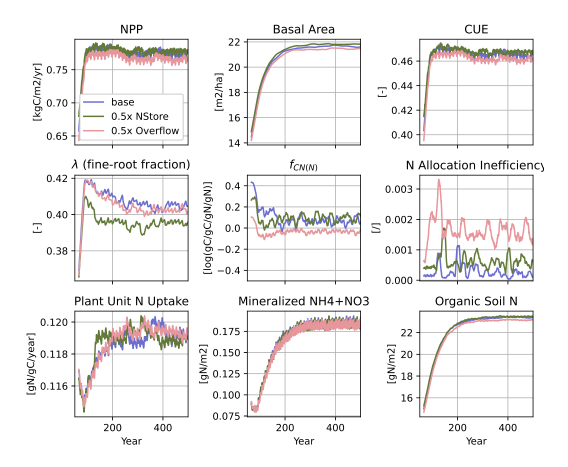


Figure F3. Sensitivity of model output to 0.5x reductions in plant Nitrogen Storage $\mu_{(N)}$ (0.5x Nstore) and overflow storage μ_{ov} (0.5x Overflow) parameters, as compared to a base simulation (1x for each).

1052 1053

F3 Experiment V: Sensitivity to alternative hypotheses for handling excess fine root and excess carbon gain

In experiment V, we evaluate how two subtle modifications to the core model hypotheses affect differences in the simulation compared to the base hypothesis. In the base hypothesis, when the fine-root biomass optimization algorithm signals a decrease in fineroot, the model will remove carbon and nutrients from the fine-root pool such that it matches the new (decreased) target. This removed mass is exported directly to the soil litter pool.

In an alternative configuration, called "efficient trim" (eff-trim), the carbon and nutri-1059 ent removed from the fine-root is sent to the plant's storage. The storage accepts this 1060 mass until full (defined by the target and overflow parameters), beyond which the mass 1061 is either burned as respiration (for carbon) or exuded to the labile soil pool (for nitro-1062 gen and phosphorus). The other alternative hypothesis, "c-exude" assumes that excess 1063 carbon acquired by the plant that can not fit in storage (defined by the target and over-1064 flow parameters) will be released into the labile litter pool, in a manner similar to the 1065 nutrients (instead of released as respiration). 1066

1067 In summary, there is very little difference in the model predictions between the three model hypotheses. Part of this may be due to evaluating these differences through a pa-1068 rameter set that has reasonable parameters and some modest optimization (in the case 1069 of the PID parameters). The plants in the base parameterization do not waste that much 1070 carbon and nutrients (see the efficiency metrics in Figure F3), and these alternative hy-1071 potheses are oriented around how these waste terms are treated. If future experiments 1072 wish to activate these alternative hypotheses, they are maintained in the FATES code 1073 behind software switches that other modelers can activate if there is interest. 1074

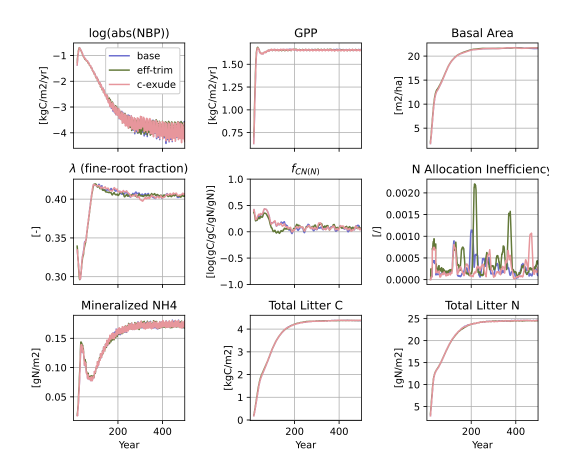


Figure F4. Comparison of model spin-ups for three different functional hypotheses. The "efftrim" simulation re-captures the carbon and nutrients from fine-roots when the PID controller decreases λ and moves it into storage (instead of releasing it to the litter pool, like turnover). The "c-exude" simulation exports carbon that is in excess of overflow storage to the labile pool in the soil, instead of burning it as respiration. The "base" simulation (the default used for all other simulations) assumes that fine-root tissues are lost as turnover when the PID controller decreases λ , and that carbon gained in excess of overflow storage is burned as respiration.

F4 Experiment VI: Sensitivity to total ecosystem fixation hypothesis: evapotranspiration versus net primary production based

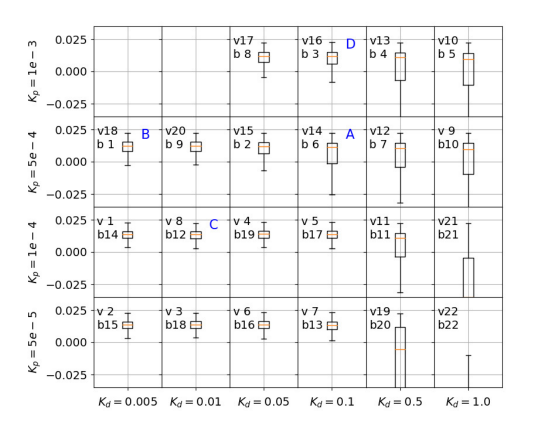
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1076

The ELM-FATES model can represent two sources of nitrogen fixation: 1) fixation 1077 that generates nitrogen in the soil and 2) obligate symbiotic nitrogen fixation that is avail-1078 able only to the plant individuals that spend the resources to fix. The nitrogen produced 1079 in the first method becomes available for competition between the nitrogen consumers 1080 in the soil (plants, nitrifiers and decomposers). In experiment IV, symbiotic nitrogen is 1081 turned off, and thus the fixation via method 1 is representative of total ecosystem fix-1082 ation, acting as a catch all for free-living microbial fixers in the soil, symbiotic fixers in 1083 root nodules as well and potentially other sources. 1084

Model response to the different total ecosystem nitrogen fixation (TEF) methods, proportionality with evapotranspiration (et-fix) and proportionality with net primary productivity (npp-fix) (Cleveland et al., 1999; P. Thornton et al., 2007), is evaluated over industrial-era atmospheric CO2 concentrations, see Figure F5. Both simulations were initialized with a pre-industrial 500 year spin-up.

Batterman et al. (2013) estimates that fixation at the BCI site is dominated by symbiotic fixation and almost negligible free-living fixation (with an exception for some old forest stands). Study at the nearby San Lorenzo (Stanton et al., 2019) site suggested that canopy microbial fixation can contribute significant sources of nitrogen.



 f_{CN}

Figure F5. Time series model response to the two different total ecosystem nitrogen fixation hypotheses in ELM, fixation proportional to NPP (npp-fix) and fixation proportional to evapotranspiration (et-fix).

The two methods generate qualitatively different results, particularly in reference 1094 to the total nitrogen fixation flux into the system. Both methods could be calibrated by 1095 adjusting scaling parameters, but we chose to use the default scaling coefficients in the 1096 ELM model, as the differences between the two options provide good end-points to study 1097 the system. Note that while the NPP method introduces more nitrogen as a whole, it 1098 also continues to increase indefinitely with CO2 increases, while the evapotranspiration 1099 method saturates and even decreases late in the 21st century. However, availability of 1100 nitrogen limits the models (as indicated by the plant unit N uptake, the nitrogen uptake 1101 per gram of fine-root, and the depleting mineralized (aqueous) N pool) in both scenar-1102 ios. The differences are mostly manifested in increased growth and biomass in the veg-1103 etation canopy with access to increased nitrogen, particularly late in the 21st century. 1104 The carbon usage efficiency is higher with more nitrogen availability, presumably because 1105 the vegetation can use the carbon it acquires to build tissues, instead of respiring it. 1106

Soil nitrogen responses are also provided in the supplement, see Figure F6. Note that in each fixation hypothesis, there are similar pattern responses to the fraction of mineralized nitrogen that can be immobilized for decomposition, as well as leaching, nitrification and denitrification response.

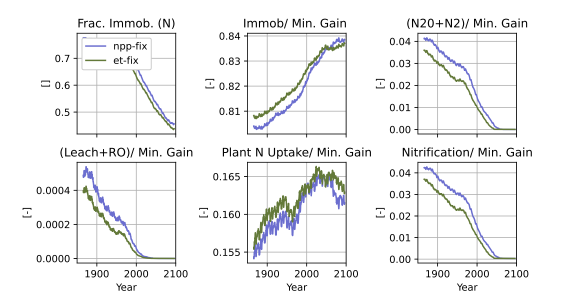


Figure F6. Differential responses of soil nitrogen flux to the two free-living fixation hypotheses (NPP based and evapotranspiration based).

Appendix G Open Research

The data and software code used to reproduce the model simulations and analysis in this manuscript have been made publicly available.

1114 G1 Data Availability

The "Next Generation Ecosystem Experiment - Tropics" project provides model driver data at the Barro Colorado Island Site, including soils and meteorological data (Knox et al., 2019). The data can be found here: https://ngt-data.lbl.gov/dois/ NGT0086/

1119 G2 Software Availability

Both the FATES and E3SM models use Git (https://git-scm.com/) version con-1120 trol to manage their software, and Github (https://github.com/) to host their soft-1121 ware. The model software of both projects and their dependencies are publicly available. 1122 Readers who wish to either reproduce or do similar work in this manuscript are encour-1123 aged to install git and use it to clone the E3SM model, also Zenodo DOIs are provided. 1124 FATES will be imported as a submodule of E3SM. To initialize submodules following 1125 a clone, and assuming the user has "checked out" the correct tag or branch, they should 1126 run the command "git submodule update -init -recursive". 1127

The specific E3SM tag used in this research is DOI 10.5281/zenodo.7684977, https:// doi.org/10.5281/zenodo.7684977; or the github tag can be found here: https://github .com/rgknox/E3SM/releases/tag/elm-fates-cnp-ms (?, ?).

The specific FATES tag used in this research is DOI 10.5281/zenodo.7685350, https:// doi.org/10.5281/zenodo.7685350; or the github tag can be found here: https://github .com/rgknox/fates/releases/tag/fates-cnp-ms-anlsys (?, ?).

- The python analysis scripts (contained in (?, ?)) used to generate the figures in this manuscript are provided in the directory: "./ms-analysis/".
- This file (contained in (?, ?)) will patch the default FATES parameter file to generate parameterizations specifically for one tropical evergreen plant functional type at Barro Colorado Island Panama: https://github.com/rgknox/fates/blob/fates-cnp -ms/parameter_files/patch_default_bciopt224.xml

A nix-type "shell" script (contained in (?, ?)) is provided, that was used to build 1140 and setup the simulations. This script should be executed from the directory: "./cime/scripts". 1141 This file also assumes that the driver data package listed above is unpacked in the same 1142 directory as well. The user will need to modify many of the paths in the script to ac-1143 commodate their file structure. This script should facilitate other users running simu-1144 lations at BCI, but this file is provided as-is, and absolutely no support will be provided 1145 for making this script work. https://github.com/rgknox/fates/blob/fates-cnp-ms/ 1146 parameter_files/create_bci_fatescnp_mscopy.sh 1147

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Figure 1.

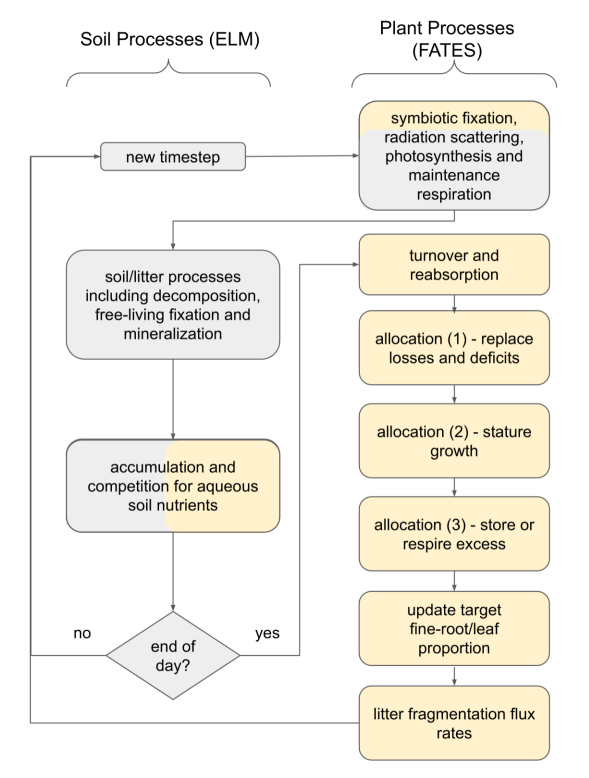


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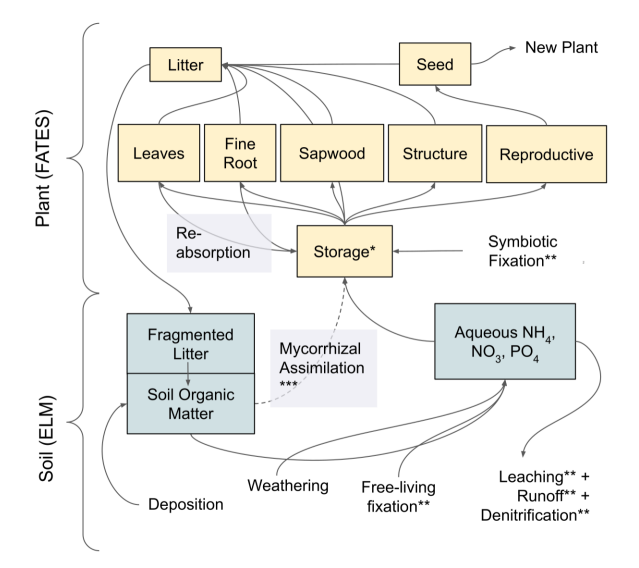
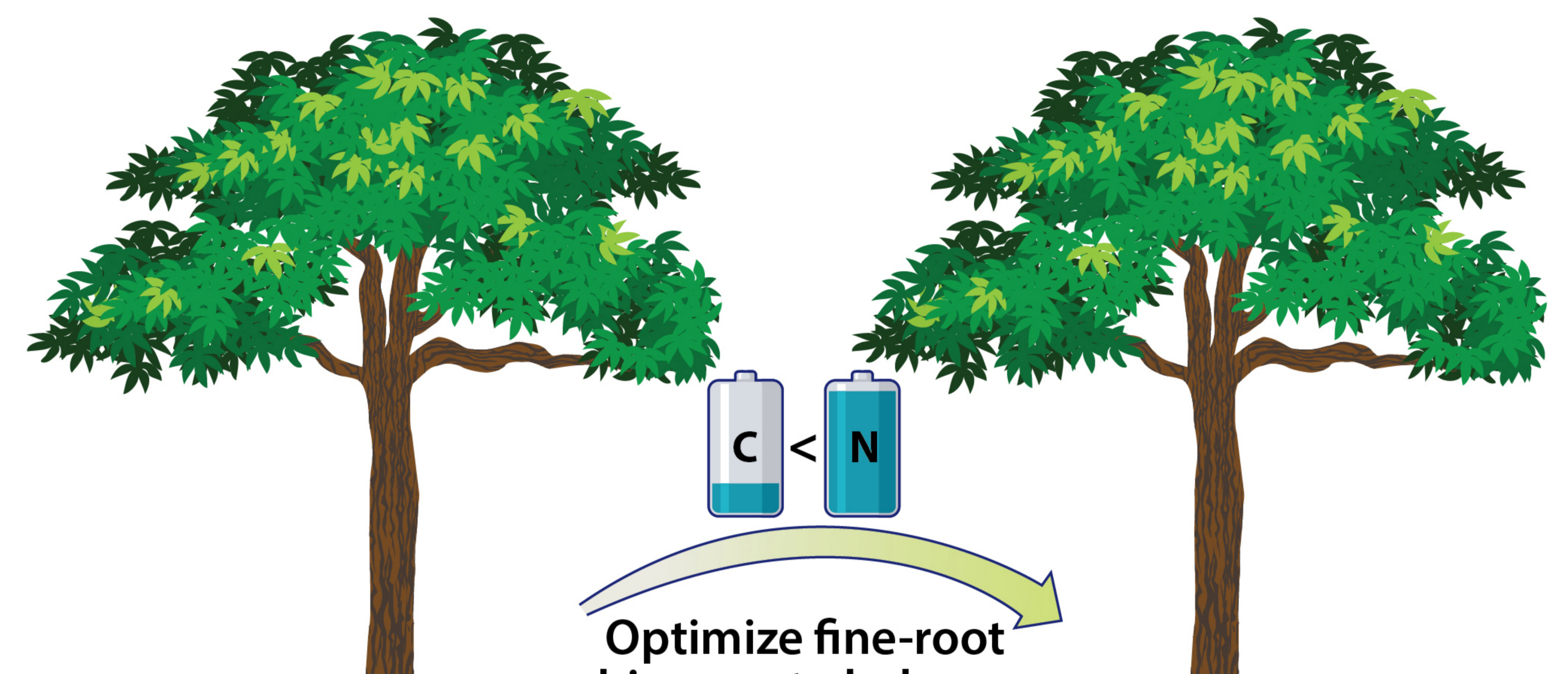
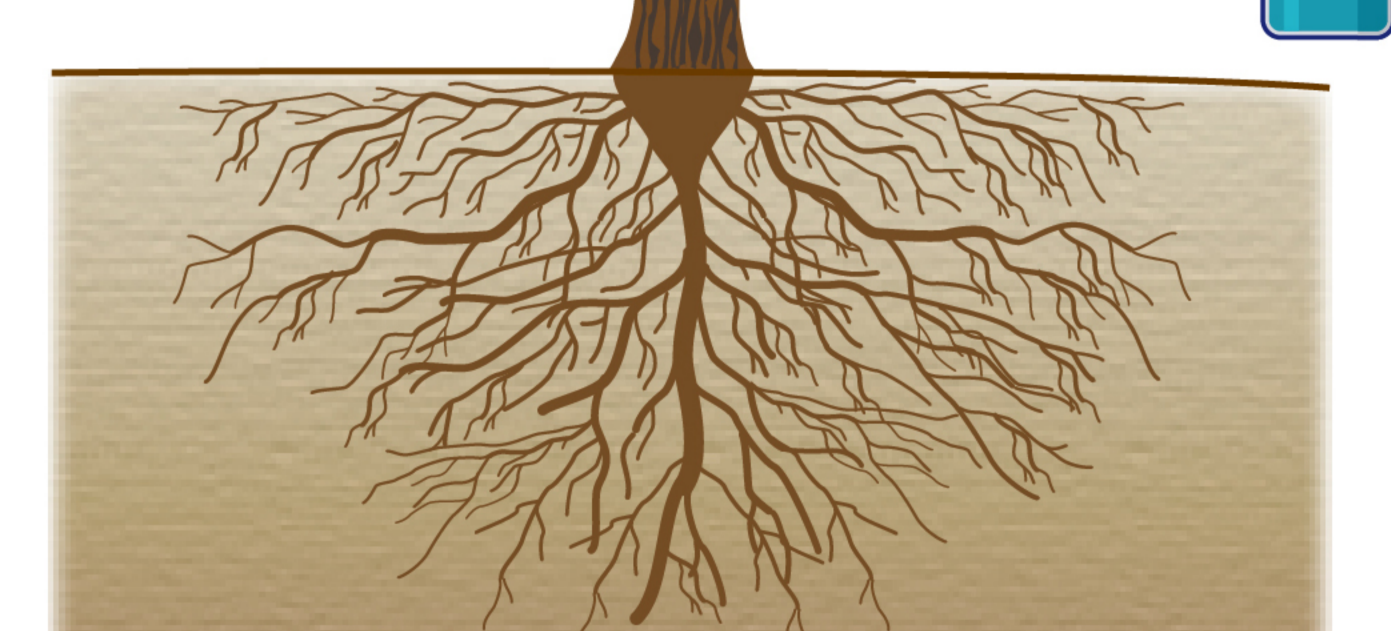


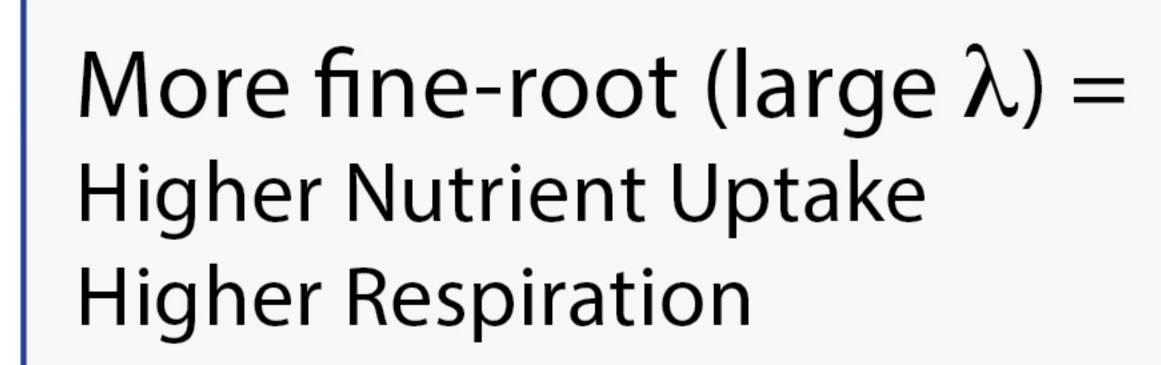
Figure 3.



biomass to balance Carbon and Nutrient storage

Ν





Less fine-root (small λ) = Lower Nutrient Uptake Lower Respiration



Figure 4.

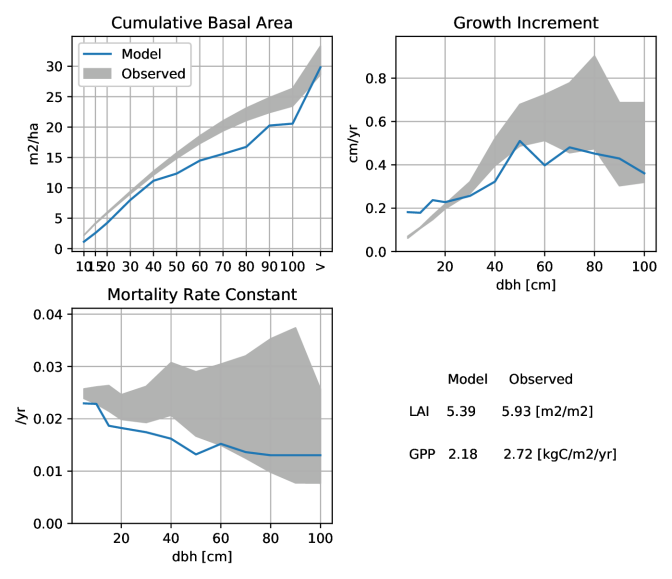
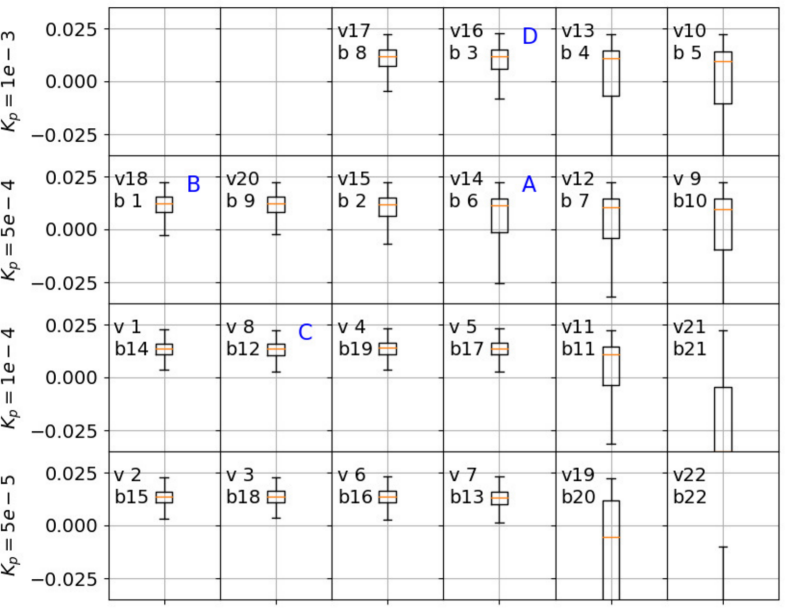


Figure 5.



 $K_d = 0.005$ $K_d = 0.01$ $K_d = 0.05$ $K_d = 0.1$ $K_d = 0.5$ $K_d = 1.0$

Figure 6.

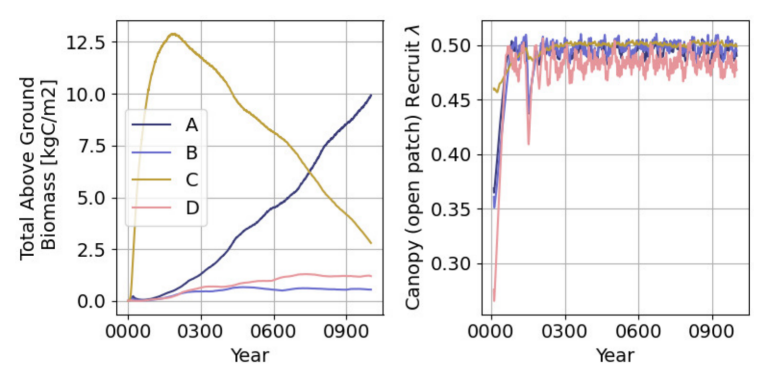
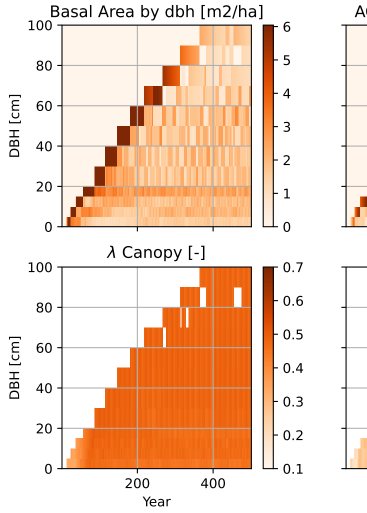


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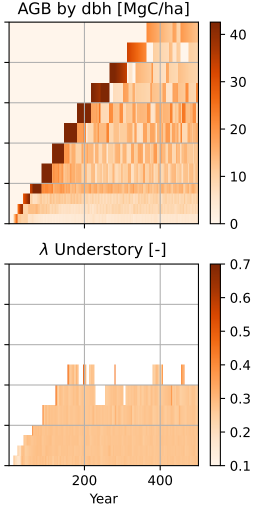


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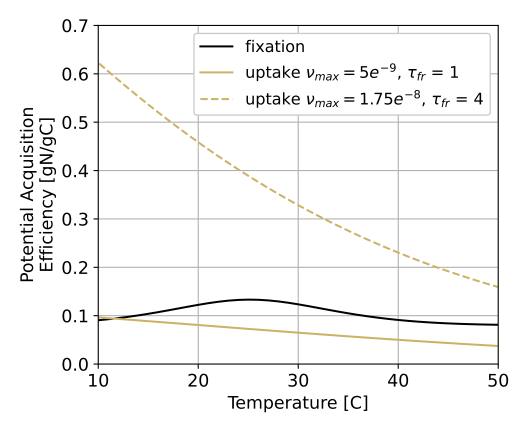


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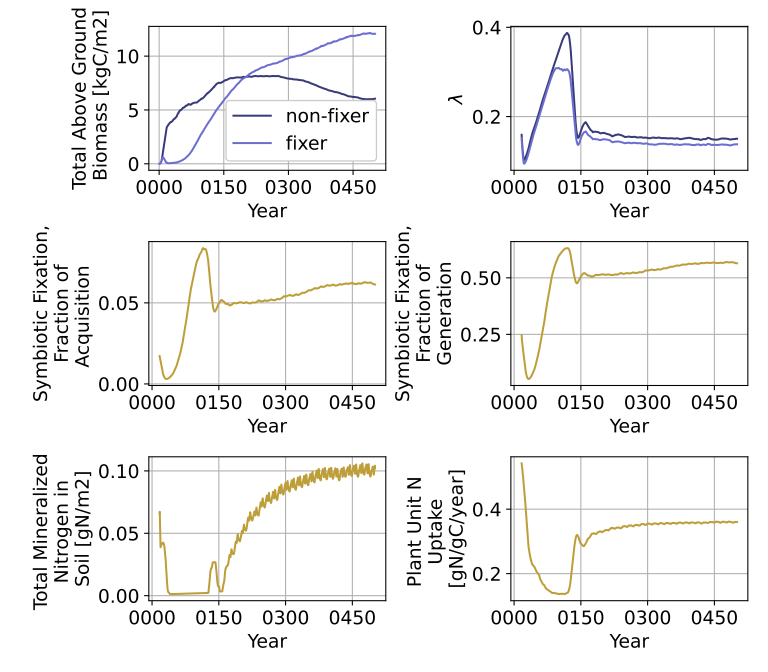
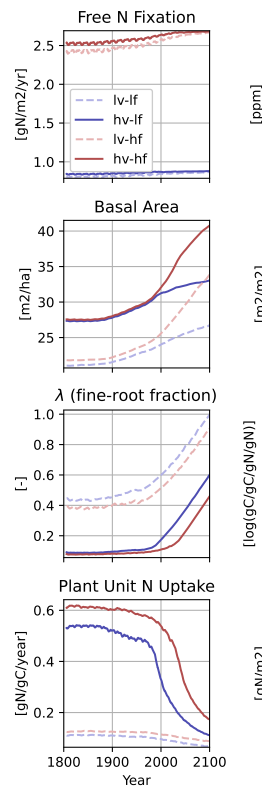
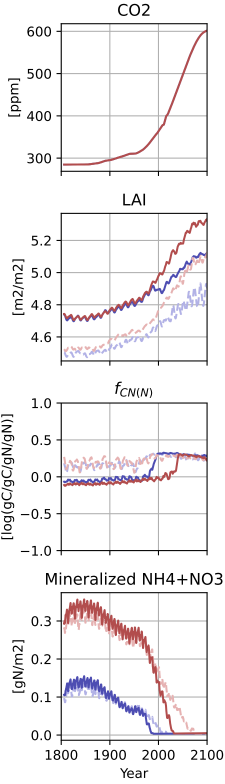
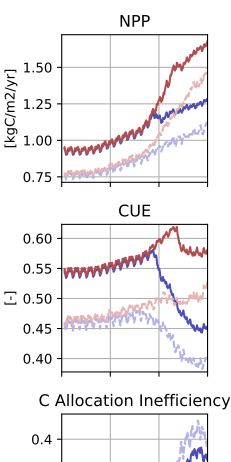
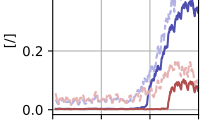


Figure 10.









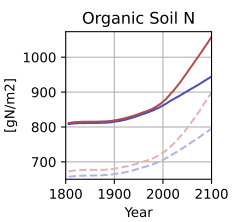


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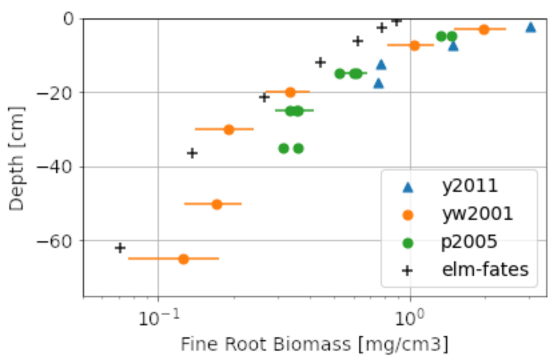


Figure E1.

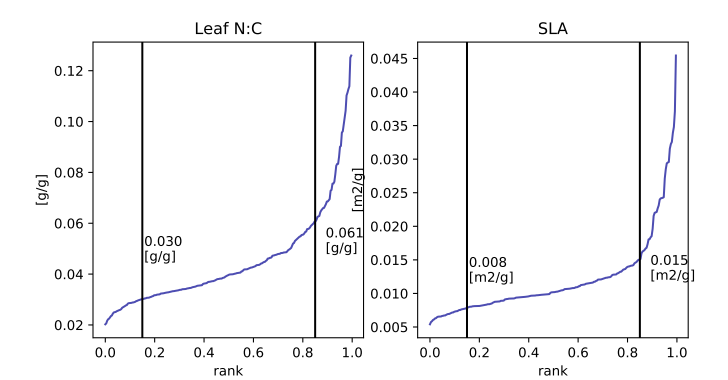


Figure E2.

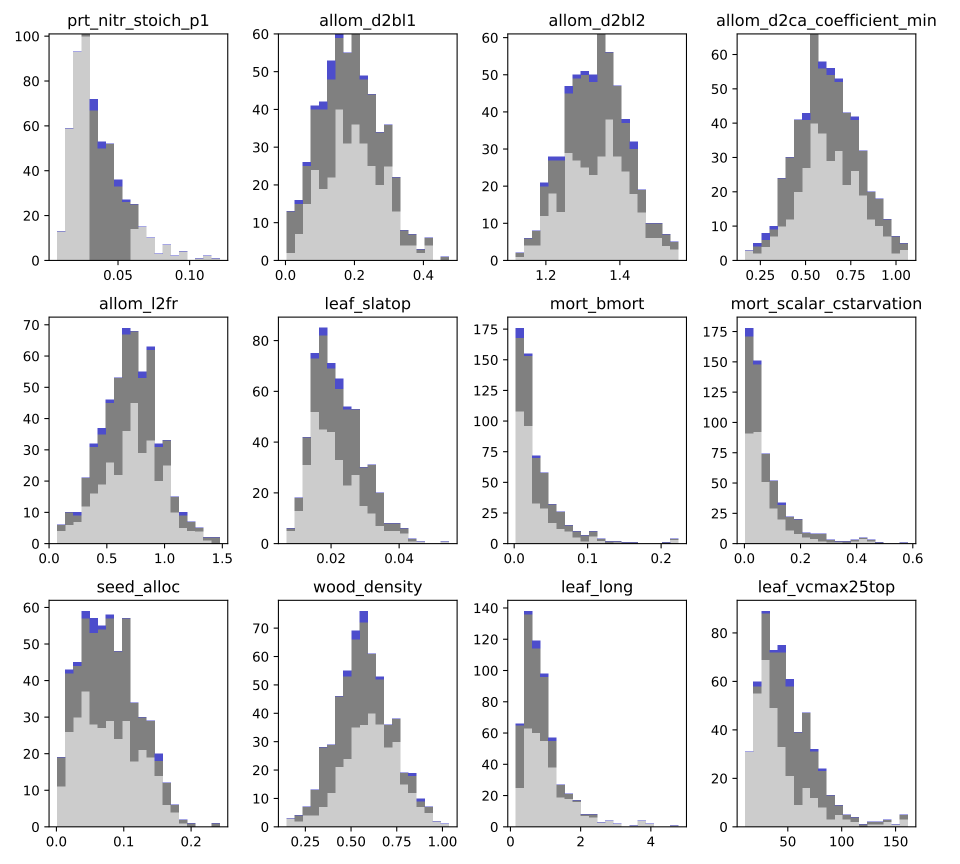
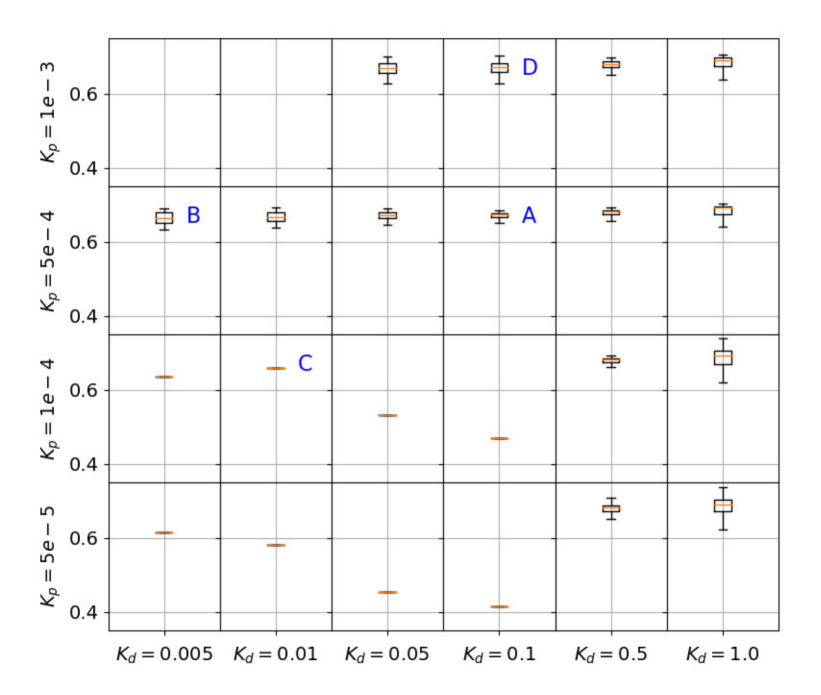


Figure F1.



λ

Figure F2.

DDBH [cm/year]

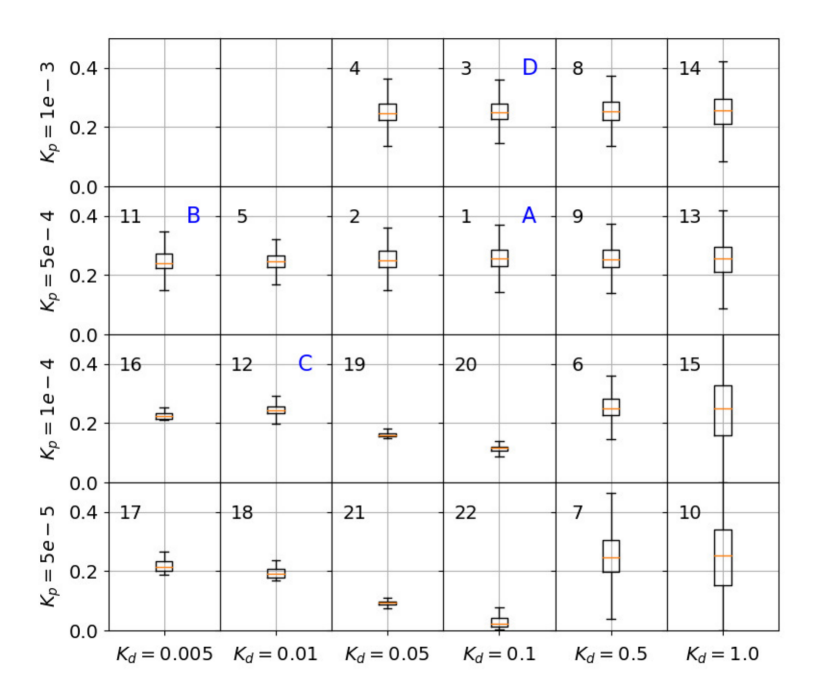


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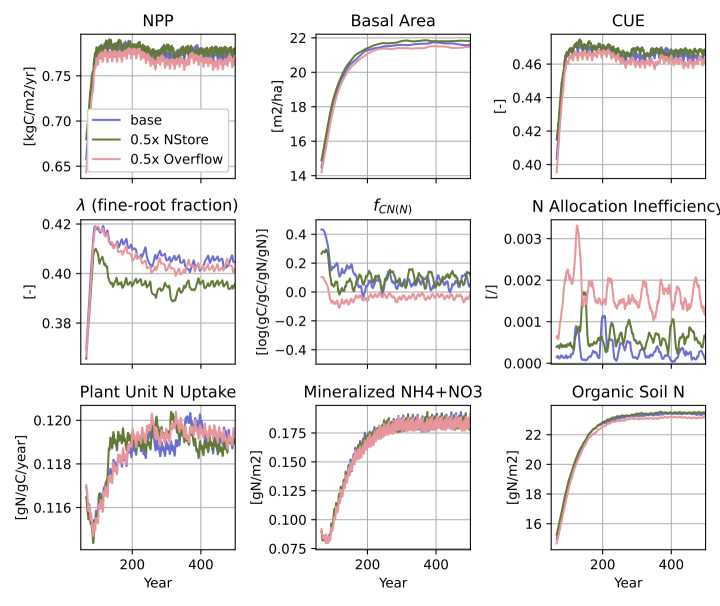


Figure F4.

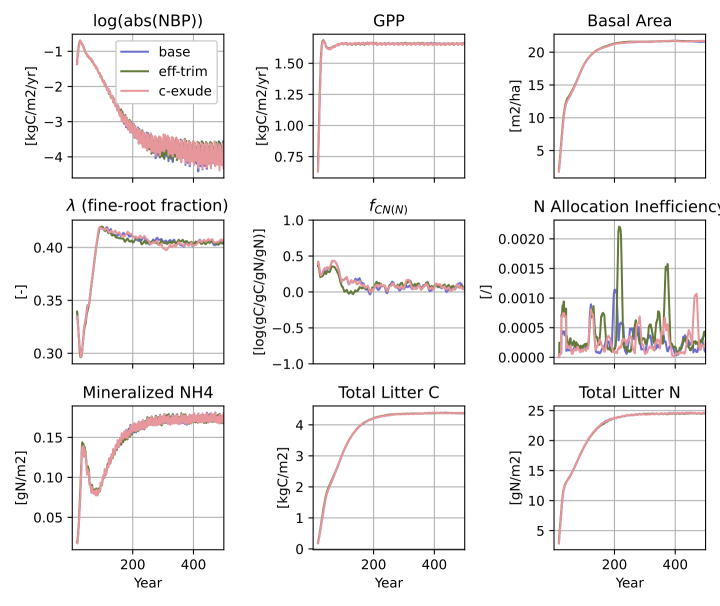


Figure F5.

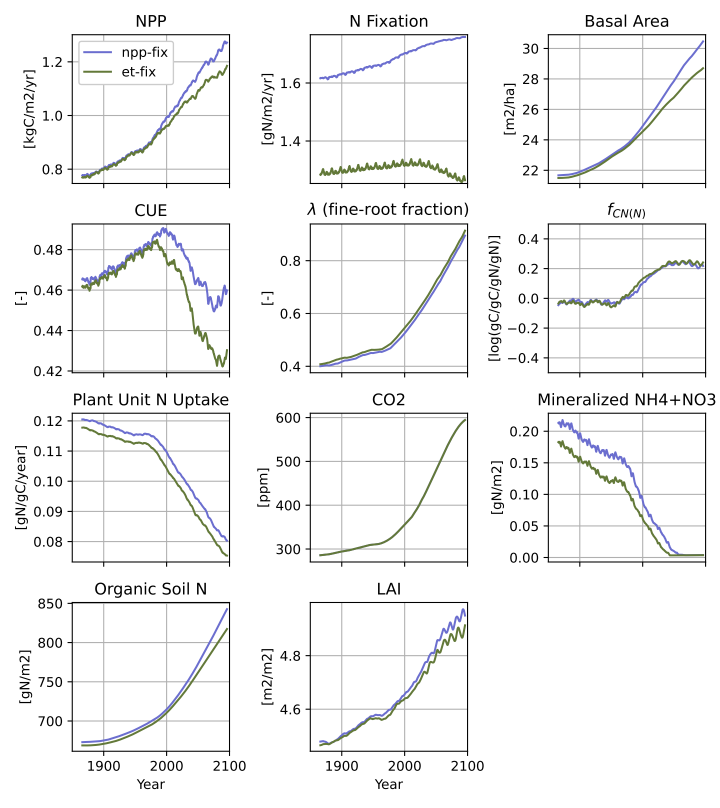


Figure F6.

