

1           **A machine learning approach targeting parameter estimation for plant**  
2           **functional type coexistence modeling using ELM-FATES (v2.0)**

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17 **Highlight**

- 18       • Machine learning based surrogate models were developed and used to optimize the  
19       selection of the trait parameters in ELM-FATES demographic vegetation model.
- 20       • Trait parameters selected by the surrogate models significantly improve the modeling of  
21       plant functional type coexistence and reduce model errors.
- 22       • This approach represents a repeatable method for identifying parameter values that  
23       satisfy fidelity against observations and coexistence between plant functional types in  
24       vegetation demography models across different ecosystems.

25

26 **Abstract**

27 Tropical forest dynamics play a crucial role in the global carbon, water, and energy cycles.  
28 However, realistically simulating the dynamics of competition and coexistence between different  
29 plant functional types (PFTs) in tropical forests remains a significant challenge. This study aims  
30 to improve the modeling of PFT coexistence in the Functionally Assembled Terrestrial Ecosystem  
31 Simulator (FATES), a vegetation demography model implemented in the Energy Exascale Earth  
32 System Model (E3SM) land model (ELM), ELM-FATES. Specifically, we explore (1) whether  
33 plant trait relationships established from field measurements can constrain ELM-FATES  
34 simulations; and (2) whether machine learning (ML) based surrogate models can emulate the  
35 complex ELM-FATES model and optimize parameter selections to improve PFT coexistence  
36 modeling. We conducted three ensembles of ELM-FATES experiments at a tropical forest site  
37 near Manaus, Brazil. By comparing the ensemble experiments without (Exp-CTR) and with (Exp-  
38 OBS) consideration of observed trait relationships, we found that accounting for these  
39 relationships slightly improves the simulations of water, energy, and carbon variables when  
40 compared to observations, but degrades the simulation of PFT coexistence. Using ML based  
41 surrogate models trained on Exp-CTR, we optimized the trait parameters in ELM-FATES, and  
42 conducted another ensemble of experiments (Exp-ML) with these optimized parameters. The  
43 proportion of PFT coexistence experiments significantly increased from 21% in Exp-CTR to 73%  
44 in Exp-ML. After filtering the experiments that allow for PFT coexistence to agree with  
45 observations (within 15% tolerance), 33% of the Exp-ML experiments were retained, which is a  
46 significant improvement compared to the 1.4% in Exp-CTR. Exp-ML also well reproduces the  
47 annual means and seasonal variations of water, energy and carbon fluxes, and the field inventory  
48 of above ground biomass. This study represents a reproducible method, which utilizes machine  
49 learning to identify parameter values that improve model fidelity against observations and PFT  
50 coexistence in vegetation demography models for diverse ecosystems. Our study also suggests the  
51 need for new mechanisms to enhance the robust simulation of coexisting plants in ELM-FATES,  
52 and has significant implications for modeling the response and feedbacks of ecosystem dynamics  
53 to climate change.

54

55 **Plain Language Summary**

56 Modeling tropical forest dynamics is crucial for understanding global carbon, water, and energy  
57 cycles under climate change. However, vegetation demographic models commonly struggle to  
58 realistically simulate the competition and coexistence of different plant functional types (PFTs).  
59 Our study explores whether plant trait measurements and machine learning surrogate models can  
60 improve plant coexistence modeling. We tested our approach using ELM-FATES at a tropical  
61 forest site. We found that using observed trait relationships did not improve PFT coexistence  
62 modeling. However, optimizing the trait parameters using machine learning-based surrogate  
63 models significantly improved PFT coexistence modeling and reduced model errors. We  
64 demonstrate the benefits of using machine learning to enhance PFT coexistence modeling, with  
65 important implications for modeling the response and feedbacks of ecosystem dynamics to climate  
66 change. Our study also highlights that new mechanisms are needed to robustly simulate coexisting  
67 plants in ELM-FATES.

68

69 **1. Introduction**

70 Tropical ecosystems feature the highest biodiversity on Earth, maintaining more than 75% of all  
71 known species (Mora et al., 2011; Mitchard, 2018). The dynamics of tropical forests are closely  
72 related to the regional and global carbon, energy and water cycles (Bonan, 2008; Piao et al., 2020).  
73 Vegetation is expected to face more water stress from vapor pressure deficit increase and soil  
74 moisture reduction with global warming (McDowell et al., 2020). Tree mortality rates are  
75 accelerating in some tropical regions due to rising atmospheric water stress (Bauman et al., 2022;  
76 Hubau et al., 2020; Zuleta et al., 2017). Tropical forests currently make an approximately neutral  
77 contribution to the global carbon cycle as a result of a large land-use source balanced by sinks in  
78 recovering and undisturbed forests, but they may become a carbon source in the future under the  
79 threat of climate change and human-induced disturbance (Mitchard, 2018; Gatti et al., 2021).  
80 Therefore, understanding and modeling tropical forest dynamics and related feedbacks have  
81 crucial implications for projecting future changes in the global climate system.

82

83 Dynamic global vegetation models (DGVMs) are the primary tools to simulate terrestrial  
84 ecosystem dynamics of plant functional type distribution, ecosystem composition and functioning,  
85 and ecosystem response to and recovery from disturbance (e.g., fire and wind damage) (Longo et  
86 al., 2019; Fisher et al., 2018; Foley et al., 1996; Sitch et al., 2003; Cao and Woodward, 1998;  
87 Berzaghi et al., 2019; McMahon et al., 2011). Conventional DGVMs represent plant communities  
88 using an area-averaged representation of plant functional types (PFTs) in each grid cell. Their  
89 relatively simple structures have the advantage of high computational efficiency for use in Earth  
90 system models (Fisher et al., 2018; Snell et al., 2014). However, these models do not capture many  
91 demographic processes. For example, plants of each represented PFT typically have identical

92 properties (e.g., tree size), which limits the capability of modeling ecosystem dynamics and  
93 functioning of canopy gap formation, PFT competition, and disturbance reactions (Feeley et al.,  
94 2007; Stark et al., 2012; Hurtt et al., 1998; Moorcroft, 2003; Brister et al., 2020). To address these  
95 limitations, researchers have developed new generation DGVMs called vegetation demographic  
96 models (VDMs), commonly including individual-based models and cohort-based models (Fisher  
97 et al., 2018). The individual-based models, also known as forest gap models, explicitly represent  
98 vegetation as individual plants and simulate their birth, growth, and death (Fyllas et al., 2014;  
99 Christoffersen et al., 2016; Sato et al., 2007; Jonard et al., 2020; Maréchaux and Chave, 2017).  
100 These models incorporate the stochasticity and heterogeneity of the plant light environment  
101 mechanistically and thereby can typically represent PFT competitive exclusion, succession, and  
102 coexistence. However, explicit simulations of individual plants with stochastic processes suffer a  
103 substantial computational penalty and limit applicability over large or global scales (Fisher et al.,  
104 2018). To capture sufficient ecosystem dynamics and maintain relatively high computational  
105 efficiency, "cohort-based" models have been proposed (Haverd et al., 2013; Medvigy et al., 2009;  
106 Ma et al., 2021; Moorcroft et al., 2001; Weng et al., 2015; Longo et al., 2019; Belda et al., 2022).  
107 In cohort-based approaches, individual plants are grouped together as "cohorts" based on their  
108 similar properties, including size, age, and PFT (Fisher et al., 2018). Many cohort-based models  
109 have been developed and widely used across regional to global scales. Examples of cohort-based  
110 models include the Ecosystem Demography model (ED) (Moorcroft et al., 2001), the Functionally  
111 Assembled Terrestrial Ecosystem Simulator (FATES) (Fisher et al., 2018, 2015), and the  
112 Geophysical Fluid Dynamics Laboratory (GFDL) Land Model 3 with the Perfect Plasticity  
113 Approximation (LM3-PPA) (Weng et al., 2015). In this study, we employ the FATES model, a  
114 widely used tool for modeling ecosystem dynamics in multiple ecosystems, including tropical

115 (Holm et al., 2020; Koven et al., 2020; Chitra-Tarak et al., 2021; Cheng et al., 2021), boreal  
116 (Lambert et al. 2022) and mixed-conifer forests (Buotte et al., 2021), and forest disturbance  
117 (Huang et al., 2020).

118

119 Despite ongoing applications, robust simulations of competition and coexistence in cohort-based  
120 VDMs remain a major challenge. In niche-based coexistence theory, coexisting species require  
121 both convergence in strategy to adapt to the surrounding environment ("environmental filtering")  
122 and divergence in strategy to ensure differentiation in resource requirements ("niche partitioning")  
123 (Kraft et al., 2008; Adler et al., 2013). These same constraints apply to coexisting PFTs as modeled  
124 by VDMs. Thus, on the one hand, VDMs need to include mechanisms that capture critical niche  
125 dimensions (e.g., spatial and temporal variation in light, water, and nutrients). For example, the  
126 multi-layer canopy structure in FATES provides vertical light resource differentiation. Another  
127 essential aspect is to assign reasonable plant functional traits (i.e., the parameters that define a  
128 given plant functional type) to satisfy environmental filtering, ensure niche partitioning, and  
129 consequently preserve PFT coexistence. Considering the relatively high computational cost of  
130 VDMs and the host land surface models, it is not feasible to directly apply global optimization  
131 methods such as Shuffled Complex Evolution (Duan et al., 1992) to calibrate trait-related  
132 parameters, because this could be time-consuming and computationally intensive (Rouholahnejad  
133 et al., 2012). Therefore, most previous studies use the filtered ensemble approach to select trait-  
134 related parameters involving several steps: 1) generate a parameter ensemble based on reference  
135 trait ranges or correlations, 2) conduct ensemble model simulations, and 3) filter the parameter  
136 ensemble by coexistence and other criteria (e.g., observation constraints). For example, Huang et  
137 al. (2020) applied FATES implemented in the Community Land Model (CLM; herein CLM-

138 FATES) with two tropical PFTs to study forest dynamics at tropical sites. They performed 70 one-  
139 at-a-time experiments before obtaining one reasonable parameter set. Buottte et al. (2021) used  
140 CLM-FATES to simulate forest dynamics of pine and incense cedar over the Sierra Nevada of  
141 California, and their two stages of experiments (360 plus 72 runs) only yielded four sets of  
142 parameters that met the given criteria. The filtered ensemble approach has low efficiency, which  
143 hinders VDMs' application to modeling ecosystem dynamics under the changing climate. In  
144 addition, trait relationships derived from field measurements are often used to infer parameter  
145 selections when simulating coexistence. For example, Longo et al. (2020) used multiple trait  
146 relationships derived from various datasets to guide parameter selection for different PFTs in the  
147 ED-2.2 model simulations. However, whether the observed trait relationships can efficiently  
148 improve PFT coexistence simulation in current VDMs is still unclear. Earlier studies using FATES  
149 have also highlighted the importance of reproductive feedbacks in maintaining or prohibiting  
150 coexistence (Fisher et al. 2010; Maréchaux and Chave 2017). But fundamentally, if PFTs have  
151 highly contrasting reproductive output, the model tends towards competitive exclusion, so  
152 discerning areas with at least approximately equal fitness is necessary. While representing a large  
153 number of plant functional types may improve the likelihood of coexistence (Koven et al., 2020),  
154 this comes at a considerable computational expense.

155

156 Machine learning (ML) has facilitated Earth science studies (Shen, 2018; Nearing et al., 2021; Zhu  
157 et al., 2022; Pal et al., 2019; Jung et al., 2019), possibly providing a promising approach to improve  
158 PFT coexistence modeling in VDMs. ML algorithms have been broadly and successfully  
159 employed in recent decades. They can be used as standalone models to predict variables of interest  
160 or integrated with process-based models to improve simulations (Xu and Liang, 2021; He et al.,



161 2022; Peatier et al., 2022). Among these applications, ML has shown advantages as a surrogate  
162 model for parameter optimization and sensitivity quantification, including its effectiveness and  
163 easy application, its ability to implicitly deal with complex nonlinear correlations and high  
164 dimensional data, and handle interactions between variables (Sit et al., 2020; Antoniadis et al.,  
165 2020; Tsai et al., 2021). One promising approach is to construct ML-based surrogate models using  
166 data from initial model simulations to emulate the relationship between inputs (i.e., model  
167 parameters) and model outputs (Wang et al., 2014). Then the computationally inexpensive  
168 surrogate model can be efficiently used for parameter optimization and sensitivity analysis. For  
169 example, Dagon et al. (2020) implemented artificial neural networks to emulate the satellite leaf  
170 area constrained version of CLM5 (Lawrence et al., 2019) and estimated optimal parameters to  
171 improve the global simulation of gross primary production and latent heat flux. Sawada (2020)  
172 developed an ML surrogate model to optimize the land surface model parameters and improve soil  
173 moisture and vegetation dynamics simulations. Watson-Parris et al. (2021) built a general tool to  
174 efficiently emulate Earth system models for uncertainty quantification and model calibration.  
175 Although employing ML based surrogate models to optimize the trait parameters and hence  
176 improve the vegetation dynamics modeling in VDMs is promising, this area of research remains  
177 under-explored.

178

179 This study aims to improve PFT coexistence modeling in VDMs. The cohort-based FATES  
180 implemented in the Energy Exascale Earth System Model (E3SM) land model (ELM; Golaz et al.,  
181 2019), i.e., ELM-FATES, is taken as our testbed. The ELM land model simulates surface energy  
182 fluxes, soil and canopy biophysics, hydrology, and soil biogeochemistry, whereas FATES  
183 simulates live vegetation processes, litter dynamics, and fire. We first examine whether trait

184 relationships constructed from field measurements can help improve ELM-FATES simulations.  
185 Second, we explore whether ML based surrogate models can help optimize key trait parameters in  
186 ELM-FATES to improve the simulation of PFTs coexistence. Our model experiments are  
187 conducted for a tropical rainforest site located in Manaus, Brazil. This paper is organized as  
188 follows. Section 2 describes ELM-FATES, summarizes the key functional trait-related parameters,  
189 introduces the machine learning algorithms, and explains the overall experimental design. Results  
190 are presented in Section 3, followed by Discussions and Conclusions in Section 4 and Section 5,  
191 respectively.

## 192 **2. Methodology**

### 193 **2.1 Study site and data**

194 Our study site is located at kilometer 34 (K34) of the ZF2 road, Manaus, Brazil (latitude: -2.6091  
195 S; longitude: -60.2093 W). The K34 site is an old-growth primary forest with minimal human  
196 disturbances (Holm et al., 2020). The annual precipitation is about 2252 mm, and the mean  
197 temperature is about 26.68 °C (<https://ameriflux.lbl.gov/sites/siteinfo/BR-Ma2>). The wet season is  
198 from November to May, and the dry season is from June to October (Fang et al., 2017). Hourly  
199 meteorological forcing (i.e., precipitation, air temperature, relative humidity, wind speed, surface  
200 pressure) at the K34 eddy covariance flux tower from 2002–2005 was obtained from the LBA-  
201 ECO CD-32 Flux Tower Network Data Compilation (Restrepo-Coupe et al., 2021). Observational  
202 reference datasets obtained from Holm et al. (2020) include gross primary production (GPP),  
203 evapotranspiration (ET), sensible heat flux (SH), Bowen ratio (BW, the ratio between sensible heat  
204 and latent heat), and inventory data-based aboveground biomass (AGB). The GPP, ET, SH, and  
205 BW observations are monthly climatological averages from 2000 to 2008 (Table S1). The AGB at  
206 this site is about  $303 \pm 2.3$  Mg/ha. These observational data were used to evaluate the ELM-  
207 FATES simulations and constrain the ML surrogate models.

208

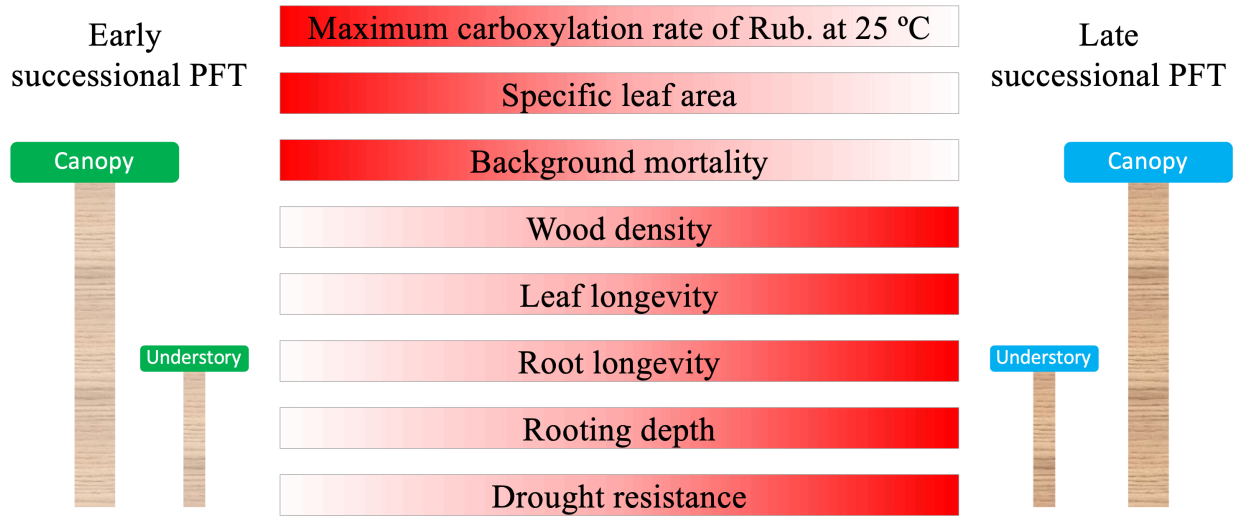
### 209 **2.2 ELM-FATES and parameters**

210 ELM-FATES is used as the model testbed. ELM is the land model of E3SM, which is the host  
211 land model of FATES (Golaz et al., 2019; Leung et al., 2020; Holm et al., 2020). FATES is a size-  
212 and age-structured vegetation model developed from the Community Land Model with ecosystem  
213 demography (CLM-ED) (Fisher et al., 2015; Koven et al., 2020). FATES includes two key  
214 structural components: ecosystem demography (ED; Moorcroft et al., 2001) and a modified

215 version of perfect plasticity approximation (PPA, Purves et al., 2008). FATES discretizes the  
216 simulated landscape into spatially implicit "*patches*" representing different disturbance histories  
217 of the ecosystem since the last disturbance. Within each patch, the hypothetical population of  
218 plants is grouped into "*cohorts*": a cohort consists of a population density of trees with similar size  
219 and the same plant functional type. Cohorts are organized, via the PPA concept, into canopy layers,  
220 and compete for light based on their canopy vertical positions (e.g., canopy layer vs. understory  
221 layer). The understory layer is formed when the canopy area becomes greater than the total ground  
222 area, and some fraction of each cohort is 'demoted' to the understory as a function of its height.  
223 The number of patches and cohorts varies depending on processes, including recruitment, growth,  
224 mortality, competition, and disturbance. The modified PPA probabilistically splits cohorts into  
225 discrete canopy and understory layers based on a function of their height (Strigul et al., 2008;  
226 Fisher et al., 2010). A detailed description of the FATES model can be found in its technical note  
227 (Zenodo, <https://doi.org/10.5281/zenodo.3517272>).

228

229 In this study, we configured two PFTs in ELM-FATES, i.e., early successional and late  
230 successional broadleaf evergreen tropical trees, which can represent a primary axis of variability  
231 in tropical forests (Huang et al., 2020; Reich, 2014; Díaz et al., 2016). There are tradeoffs between  
232 the plant traits of these two PFTs. Compared with the late successional PFT, the early successional  
233 PFT is more light-demanding and fast-growing, but with lower woody density, shorter leaf and  
234 root lifespans, and higher background mortality. To represent the drought impacts on forest  
235 dynamics, the early successional PFT is further assumed to be less drought resistant with shallower  
236 rooting depth and hence more easily affected by drought conditions (Oliveira et al., 2021). The  
237 corresponding tradeoffs and parameters between these two PFTs are shown in Figure 1 and Table



239

240 Figure 1. Schematic representation of tradeoffs between early and late successional PFTs. Dark  
 241 red denotes a higher parameter value. The tradeoffs of the top five traits are used to constrain the  
 242 parameter sampling.

243

Table 1 Summary of ELM-FATES trait parameters for two PFTs

Parameter type	Parameter name	Symbol	Unit	Early PFT	Late PFT	Range
Optimized parameter	Maximum carboxylation rate of Rub. at 25 °C, canopy top	$V_{cmax}$	$\mu\text{mol CO}_2/\text{m}^2/\text{s}$	$V_{cmax,early} > V_{cmax,late}$		40–105
	Specific leaf area, canopy top	$SLA$	$\text{m}^2/\text{gC}$	$SLA_{early} > SLA_{late}$		0.005–0.04
	Background mortality rate	$M_{bk}$	1/yr	$M_{bk,early} > M_{bk,late}$		0.005–0.05
	Wood density	$WD$	$\text{g}/\text{cm}^3$	$WD_{early} < WD_{late}$		0.2–1.0
	Leaf longevity	$L_{leaf}$	year	$L_{leaf,early} < L_{leaf,late}$		0.2–3.0
Fixed parameter	Maximum size of storage C pool, relative to the maximum size of leaf C pool	$CR_{s2l}$	—		same	0.8–1.5
	Root longevity	$L_{root}$	year	0.9	2.6	—
	Fine rooting distribution profile parameter a	$R_a$	—	7	7	—
	Fine rooting distribution profile parameter b	$R_b$	—	2	0.4	—
	BTRAN threshold below which drought mortality begins.	$M_{btran}$	—	0.4	1.0E-06	—
	Soil water potential at full stomatal closure	$\psi_{closure}$	mm	-113000	-242000	—

244

245

246

247

248

\*Parameter references (Huang et al., 2020; Koven et al., 2020; Longo et al., 2020; Holm et al., 2020; Cheng et al., 2021; Domingues et al., 2005; Chitra-Tarak et al., 2021; Buotte et al., 2021)

\* $R_a$  and  $R_b$  are parameters that determine the rooting depth and vertical distribution of fine roots.

\*BTRAN is the plant water stress factor. BTRAN  $\in [0,1]$ , 0 representing full water stress, 1 representing no water stress.

### 249 **2.3 Machine learning algorithm**

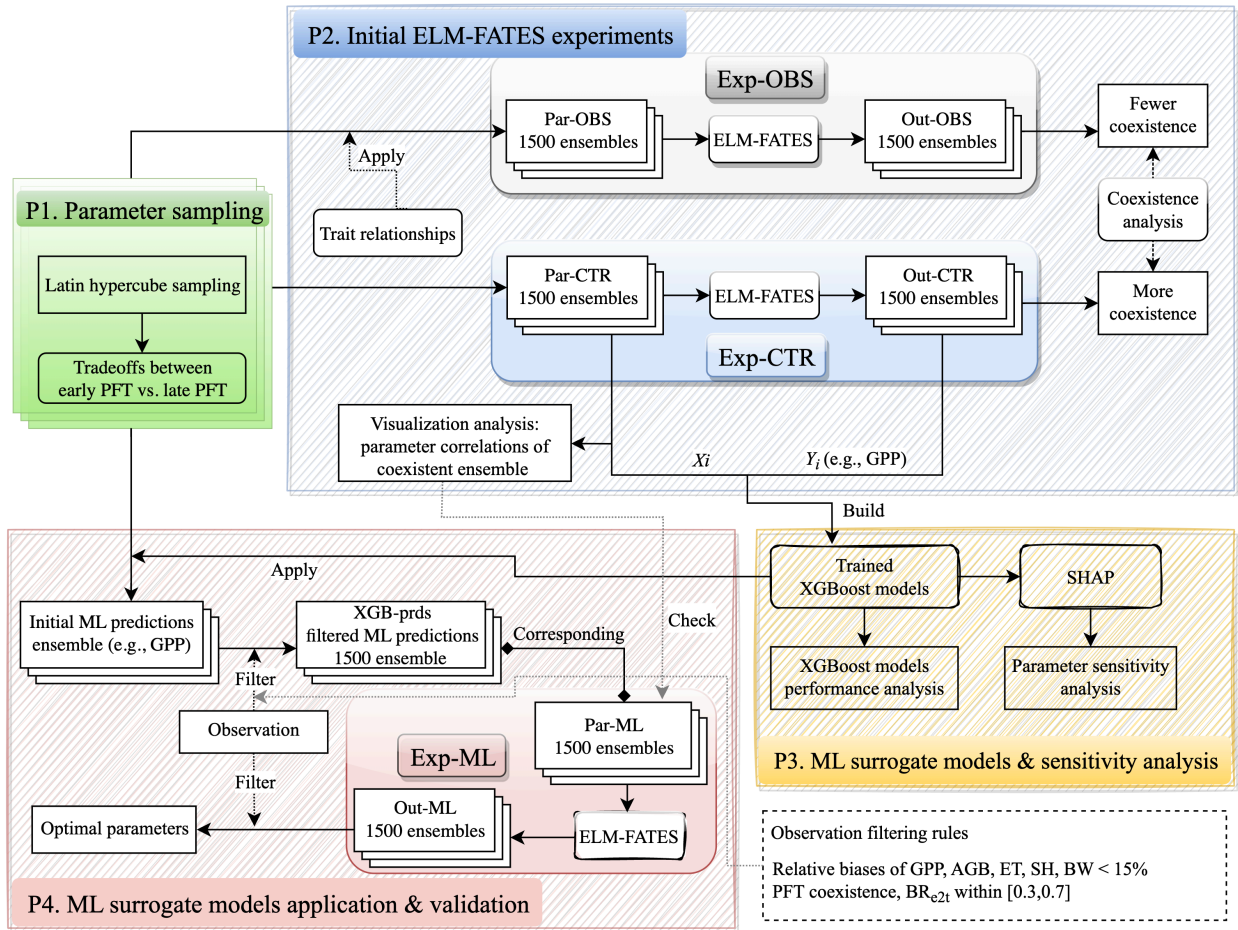
250 We built ML-based surrogate models to emulate ELM-FATES simulations. To represent the  
251 relationships between ELM-FATES parameters and simulations (e.g., AGB), we used eXtreme  
252 Gradient Boosting (XGBoost; Chen and Guestrin, 2016), a decision-tree-based ensemble machine  
253 learning algorithm. Ensemble learning techniques combine the predictions of multiple independent  
254 base models (e.g., decision trees) to produce more accurate predictions, with popular algorithms  
255 such as Random Forest (Breiman, 2001) and XGBoost. While Random Forest builds an ensemble  
256 of parallel trees using bagging and produces the final prediction by averaging the outputs of all  
257 individual trees, XGBoost sequentially trains a set of decision trees using boosting (Friedman,  
258 2001), where each successive tree corrects the mistakes of its predecessors, and the final prediction  
259 is obtained by combining the predictions of all trees using a weighted sum. XGBoost not only  
260 handles complex nonlinear interactions and collinearity between different features, but also  
261 provides a parallel implementation that effectively solves a range of data science problems. It has  
262 been successfully applied in a variety of fields within Earth and Environmental Sciences, such as  
263 urban temperature emulation (Zheng et al., 2021b), wildfire burned area (Wang et al., 2021), and  
264 emissions prediction (Wang et al., 2022), flash flood risk assessment (Ma et al., 2021), and aerosol  
265 property estimation (Zheng et al., 2021a, c).

266

### 267 **2.4 Experimental design**

268 The experimental design flowchart is shown in Figure 2. Overall, we generated three ensembles  
269 of parameter values, i.e., Par-CTR, Par-OBS, and Par-ML, and conducted three ensembles of  
270 corresponding ELM-FATES experiments, i.e., Exp-CTR, Exp-OBS, and Exp-ML. Exp-CTR is the  
271 control experiment without being constrained by the observed trait relationships. Exp-OBS

272 considered the constraint of the observed trait relationships. The Par-ML was generated by  
 273 machine learning surrogate models, which were trained based on Exp-CTR, and then used to  
 274 conduct Exp-ML. The detailed experiment procedures are described below.



275

276

277

Figure 2. Overall flowchart of experimental design and associated analysis.

278 **2.4.1 Procedure 1: Parameter sampling**

279 The procedure "P1" in Fig. 2 is used to generate an ensemble of parameter values for each  
280 experiment ensemble, i.e., Exp-CTR, Exp-OBS, and Exp-ML. First, a number of initial parameter  
281 sets (e.g., 5000 sets) were generated using Latin Hypercube Sampling (LHS; Mckay et al., 2000).  
282 Second, the initial parameter sets were filtered by the trait tradeoffs between early and late  
283 successional PFTs (Figure 1). We repeatedly increased the number of initial parameter sets in the  
284 first step until 1500 parameter sets were obtained in the second step. Each ELM-FATES  
285 experiment starts from bare ground and runs for 350 years to reach an equilibrium state, by cycling  
286 the meteorological forcing during 2002–2005, and the last four years of the simulations were  
287 analyzed.

288

289 **2.4.2 Procedure 2: Initial ELM-FATES experiments of Exp-CTR and Exp-OBS**

290 To test whether plant trait relationships established from field measurements can improve the  
291 ELM-FATES simulations, we derived three trait relationships based on the tropical studies of  
292 Koven et al. (2020) and Longo et al. (2020). Using the digitized data from Figure 3 in Koven et al.  
293 (2020), background mortality  $M_{bk}$  (see table 1 for parameter definitions) can be empirically  
294 computed from  $V_{cmax}$ ,

295 
$$M_{bk} = 0.0082 \times e^{(0.0153 \times V_{cmax})} \quad (1)$$

296 Based on the equations in Figure S18 of Longo et al. (2020),  $L_{leaf}$  and  $WD$  can be calculated via  
297  $SLA$ -(see table 1 for parameter definitions),

298 
$$L_{leaf} = 0.0001 \times SLA^{(-2.32)} \quad (2)$$

299 
$$WD = -0.583 \times \ln(SLA) - 1.6754 \quad (3)$$

300 These trait relationships were used to generate parameters for Par-OBS.



301 Two initial sets of experiment ensembles, i.e., Exp-CTR and Exp-OBS (procedure "P2" in Figure  
302 2), were conducted based on Par-CTR and Par-OBS, respectively. For Par-CTR, 1500 parameter  
303 sets were generated from the procedure "P1" based on the entire eleven parameters' space (i.e.,  
304  $V_{cmax,early}$ ,  $V_{cmax,late}$ ,  $SLA_{early}$ ,  $SLA_{late}$ ,  $M_{bk,early}$ ,  $M_{bk,late}$ ,  $WD_{early}$ ,  $WD_{late}$ ,  $L_{leaf,early}$ ,  
305  $L_{leaf,late}$ ,  $CR_{s2l}$ ). For Par-OBS, 1500 parameter sets were generated from the procedure "P1" but  
306 only based on five parameters' space (i.e.,  $V_{cmax,early}$ ,  $V_{cmax,late}$ ,  $SLA_{early}$ ,  $SLA_{late}$ ,  $CR_{s2l}$ ). The  
307 other six parameters ( $M_{bk,early}$ ,  $M_{bk,late}$ ,  $WD_{early}$ ,  $WD_{late}$ ,  $L_{leaf,early}$ ,  $L_{leaf,late}$ ) in Par-OBS  
308 were calculated based on the traits relationships defined by Equations (1) ~ (3). Therefore,  
309 compared to Par-CTR, the parameters in Par-OBS are constrained by the observed trait  
310 relationships. The distributions of these two parameter sets are shown in Figure S1.  $V_{cmax}$ ,  $SLA$ ,  
311 and  $CR_{s2l}$  have similar distributions between Par-CTR and Par-OBS. Compared with Par-CTR,  
312 Par-OBS has a narrower distribution of  $M_{bk}$  but broader distributions of  $WD$  and  $L_{leaf}$ .

313

314 Exp-CTR and Exp-OBS each include 1500 350-year ELM-FATES simulations. We averaged the  
315 last four years of these simulations for analysis, i.e., simulation outputs: Out-CTR and Out-OBS,  
316 respectively. To quantify the PFT coexistence, we computed the biomass ratio between early  
317 successional PFT and the total biomass, denoted as  $BR_{e2t}$ . For brevity, we denote the ELM-  
318 FATES experiments with  $BR_{e2t} \in [0.1, 0.9]$  as "coexistence",  $BR_{e2t} \in [0.0, 0.1)$  as "late",  
319  $BR_{e2t} \in (0.9, 1.0]$  as "early". We calculated  $BR_{e2t}$  based on Out-CTR and Out-OBS, and then  
320 computed the fraction of coexistence experiments in each ensemble. As we will show in section  
321 3.1, considering the observed trait relationships, Exp-OBS has a lower fraction of coexistence  
322 experiments. Therefore, only Exp-CTR was used for further ML-related analysis. We also

323 performed some analysis of Exp-CTR to explore whether the parameters of the coexistence  
324 experiments have correlations with each other (Section 3.2).

### 325 **2.4.3 Procedure 3: ML surrogate models & sensitivity analysis**

326 Based on Exp-CTR, we trained XGBoost models to emulate the ELM-FATES model behavior and  
327 analyzed the parameter sensitivity using SHAP (procedure "P3" in Figure 2). Sixteen variables  
328 were used as XGBoost model features, including 11 parameters in Par-CTR and 5 parameter  
329 differences between early and late successional PFTs. The corresponding ELM-FATES annual  
330 average outputs were used as XGBoost model targets. Specifically, six models were built, i.e.,  
331 XGB\_ET, XGB\_SH, XGB\_BW, XGB\_GPP, XGB\_AGB, XGB\_BR for predicting ET, SH, BW,  
332 GPP, AGB, and  $BR_{e2t}$ , respectively. The ML models were trained and tested as described in  
333 Section 2.5, and subsequently utilized to perform the parameter sensitivity analysis (Section 2.5).

### 334 **2.4.4 Procedure 4: ML surrogate models application & validation**

335 The trained XGBoost models were then used to help select ELM-FATES parameters (procedure  
336 "P4" in Figure 2). First, initial parameter sets were generated from procedure "P1" based on the  
337 entire eleven parameters' space (Table 1, identical to the parameters' space used for the generation  
338 of Par-CTR). Second, these parameter sets and parameter differences were sent to six XGBoost  
339 surrogate models to predict ET, SH, BW, GPP, AGB, and  $BR_{e2t}$ . Third, the predictions were  
340 further filtered by two criteria: (1) compared to observations, the relative biases of the predicted  
341 ET, SH, BW, GPP, and AGB should be less than 15%; (2) the XGBoost model predicted  $BR_{e2t}$   
342 should be within [0.3, 0.7], which corresponds to the range where the XGB-BR model exhibited  
343 relatively better performance (Figure 5). We repeated these three steps until we obtained 1500 sets  
344 of XGBoost model predictions that matched the criteria. Finally, we obtained 1500 sets of  
345 XGBoost model predictions and their corresponding 1500 sets of parameters (Par-ML). We also

346 checked whether the selected Par-ML could match the empirical relationships derived from the  
347 empirical analysis in procedure "P2" (see Sections 3.2 and 3.5 for details). Then, the 1500 sets of  
348 parameters in Par-ML were sent to ELM-FATES to conduct 350-year runs (i.e., Exp-ML). The  
349 last four years of the simulations were averaged (i.e., Out-ML) for further analysis. We then  
350 filtered Out-ML based on a relative bias of 15% or less compared to observations and PFT  
351 coexistence to identify the optimal experiments and corresponding parameters.

352

### 353 **2.5 ML model development and SHAP analysis**

354 The process of building each of the six ML surrogate models is described. Taking  $BR_{e2t}$  as an  
355 example, the 1500 pairs of sixteen features and the corresponding simulated  $BR_{e2t}$  were randomly  
356 split into two groups, 90% used for training and the remaining 10% used for testing. Given that  
357 the coexistence experiments only account for 20.6% in the simulations of Exp-CTR (Section 3.1),  
358 we used 90% of the data for training to ensure sufficient coexisting samples were included in the  
359 training process. Optimizing the hyperparameters of the XGBoost model is crucial for its  
360 performance. To achieve this, we employed the Bayesian optimization method during the training  
361 process (Snoek et al., 2012). In addition, to avoid overfitting during hyperparameter optimization,  
362 we utilized a five-fold cross-validation method (Feigl et al., 2021). The mean squared error was  
363 used as the objective function to achieve the optimal hyperparameters. The root mean squared  
364 error (RMSE) and R-squared ( $R^2$ ) are used to quantify the overall model performance for the  
365 training and testing data prediction.

366 Based on the trained XGBoost models, we subsequently employed a game theoretic approach  
367 called SHapley Additive exPlanations (SHAP; Lundberg and Lee, 2017; Lundberg et al., 2018,  
368 2020) to gain insights into the parameter sensitivity of ELM-FATES. SHAP assumes that features

369 (predictive variables) interact and collaborate in a prediction game, with each feature receiving a  
370 payout for its contributions. This approach provides a unified measure of feature importance to  
371 explain both individual samples and the entire dataset, which is distinct from intrinsic feature  
372 importance methods such as the feature importance in XGBoost (Lundberg and Lee, 2017). This  
373 approach has been widely used in various fields, including interpreting a digital soil mapping  
374 model (Padarian et al., 2020) and identifying the critical drivers of wildfires (Wang et al., 2021).  
375 In this study, we performed SHAP analysis for each XGBoost model and used the SHAP values  
376 as a proxy to quantify the relative importance of ELM-FATES parameters.

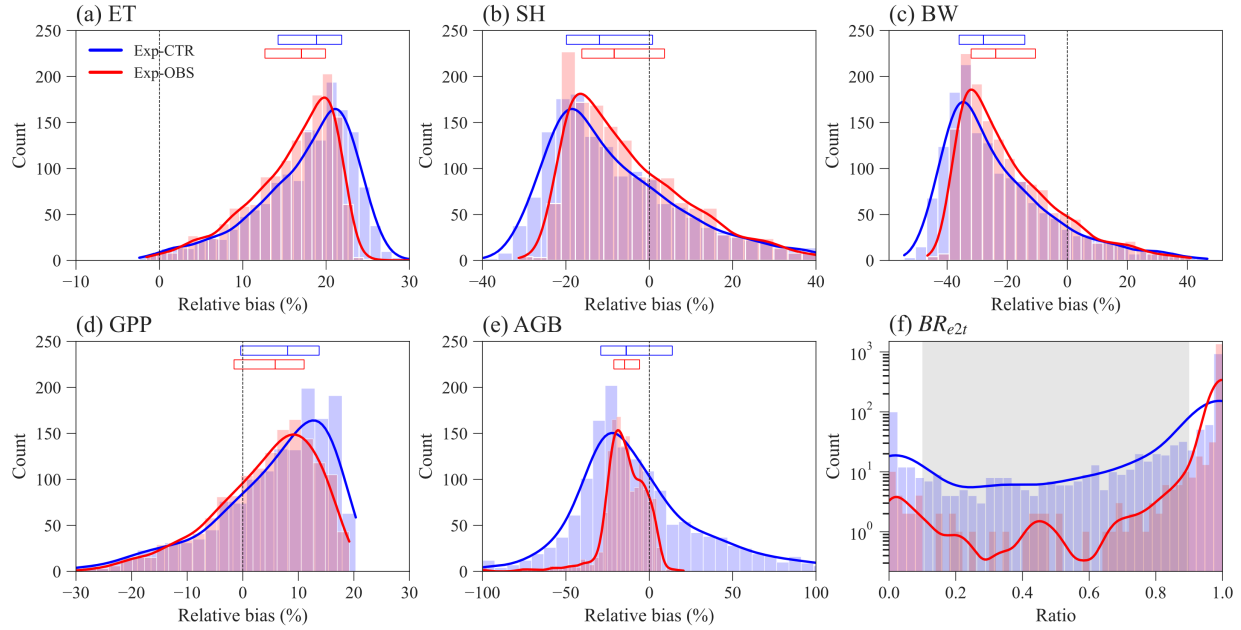
377

### 378 3. Results

#### 379 3.1 Comparison between Exp-CTR and Exp-OBS

380 Constraining the input traits using the observed trait relationships yields slightly better ELM-  
381 FATES simulations of water, energy, and carbon variables (Figures 3a~3e). The distributions of  
382 the relative biases of ET, SH, BW, and GPP have similar ranges between the two sets of  
383 experiments (Figures 3a~3d). Compared with Exp-CTR, the 50<sup>th</sup> percentiles of relative biases of  
384 ET, SH, BW, and GPP for Exp-OBS (with constrained traits) are closer to zero, indicating Exp-  
385 OBS is slightly better than Exp-CTR. The distribution of simulated AGB for Exp-OBS is much  
386 narrower than Exp-CTR (Figure 3e), which could be due to the narrower distribution of  $M_{bk}$   
387 (Figure S1).

388 Exp-CTR has a much higher fraction of PFT coexisting simulations than Exp-OBS (Figure 3f and  
389 Table S2). Overall, 70.6 % of experiments in Exp-CTR, and 94.5% of experiments in Exp-OBS  
390 have high simulated  $BR_{e2t}$  that is greater than 0.9. This indicates that both Par-CTR and especially  
391 Par-OBS favor the early successional PFT. As for the coexisting experiments with  $BR_{e2t} \in$   
392  $[0.1, 0.9]$ , Exp-CTR has about five times more coexisting experiments (20.6%) than Exp-OBS  
393 (4.1%). Further filtering the coexisting cases by observations (Table S1), only 21 experiments  
394 remain in Exp-CTR, and 6 experiments in Exp-OBS (Table S2). Even though Exp-OBS considered  
395 the observed trait relationships, it has fewer coexisting cases within the reasonable observation  
396 ranges than Exp-CTR. Therefore, Exp-OBS is not used in our remaining analysis.



397

398 Figure 3. Distribution of ELM-FATES simulations for Exp-CTR and Exp-OBS. The y-axis in (f)

399 is logarithmic.  $Relative\ bias = \frac{simulation - observation}{observation} \times 100\%$ . In (a)~(e), the top horizontal

400 bars with three vertical lines denote the relative bias at the 25<sup>th</sup>, 50<sup>th</sup>, and 75<sup>th</sup> percentiles,

401 respectively. The grey shaded area in (f) represents the coexistence biomass ratio between 0.1

402 and 0.9.

### 403 3.2 Parameter analysis of Exp-CTR

404 We also tested whether simple parameter correlations can be constructed to guide the simulation

405 of PFTs coexistence. No simple parameter correlations can be built to distinguish the coexisting

406 cases from the early and late cases in Exp-CTR (Figures 4, S2, and S3). Most parameter (or

407 parameter difference) spaces show large overlaps between early, late, and coexisting cases

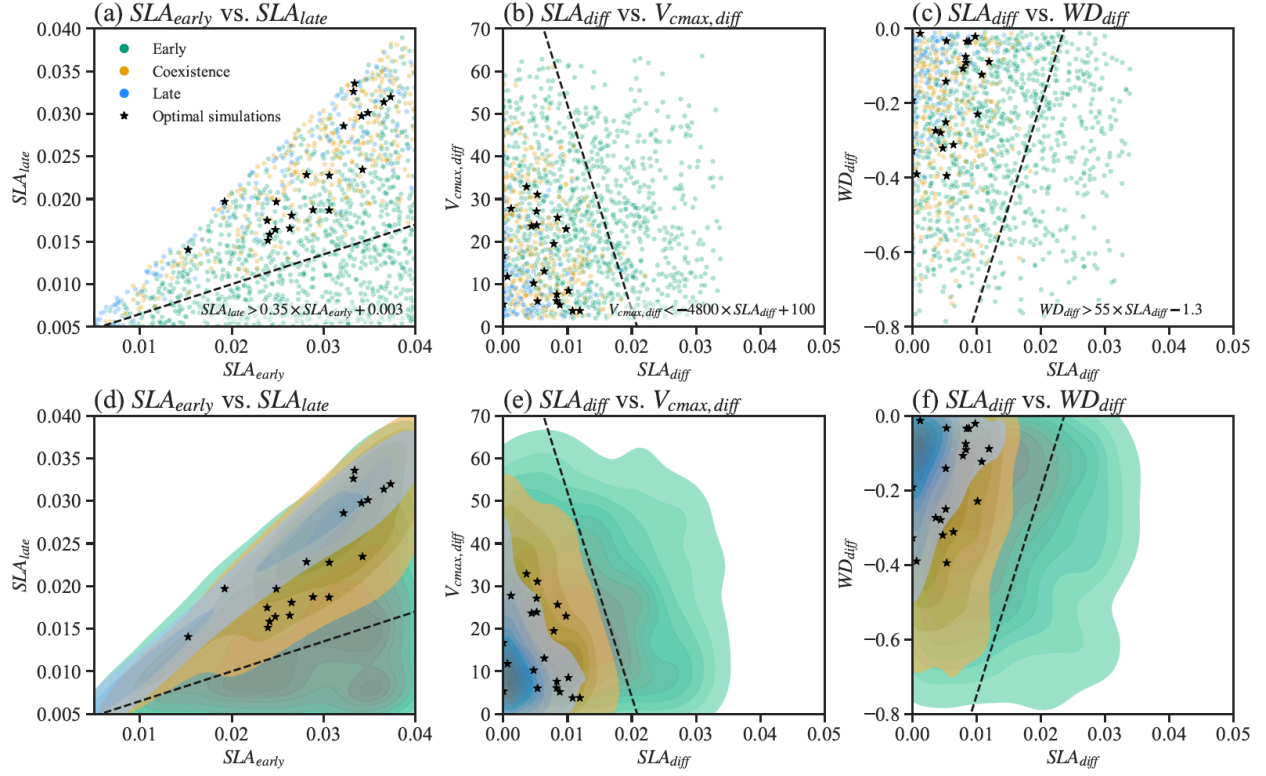
408 (Figures S2 and S3). Notably, we empirically built three linear equations based on the boundaries

409 in the parameter spaces for the coexisting cases (Figure 4). Coexisting cases are primarily located

410 in spaces with  $SLA_{late} > 0.35 \times SLA_{early} + 0.003$  (Figures 4a and 4d),  $V_{cmax,diff} < -4800 \times$

411  $SLA_{diff} + 100$  (Figures 4b and 4e), and  $WD_{diff} > 55 \times SLA_{diff} - 1.3$  (Figures 4c and 4f),

412 where  $V_{cmax,diff} = V_{cmax,early} - V_{cmax,late}$ , and  $SLA_{diff}$  and  $WD_{diff}$  are defined likewise.  
413 Within these constrained parameter spaces, the percentage of coexisting cases increases from the  
414 original 20.6% (i.e., 309 out of 1500) to 32.6% (i.e., 304 out of 932). Therefore, these empirical  
415 correlations could help guide ELM-FATES parameter selection for coexisting PFTs. On the other  
416 hand, a dominant proportion (i.e., 67.4% (1-32.6%)) of experiments are still either early or late  
417 cases within the constrained parameter spaces and cannot robustly predict PFT coexistence.  
418 Moreover, despite further considering the observational constraints (black scatters in Figure 4;  
419 Table S2), the 21 experiments (2.3%, 21 out of 932) are still sparsely distributed in the parameters'  
420 space of the coexisting cases, so no simple correlations can be developed based on these  
421 simulations. Therefore, simple empirically built relationships between plant traits provide limited  
422 benefit to guiding ELM-FATES parameter selection for modeling PFTs coexistence while  
423 matching the observations. This finding provides additional motivation for the ML-based  
424 approaches.



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Figure 4. Relationships between selected parameters of Par-CTR. These parameters are presented in three groups, i.e., green color for the late cases with  $BR_{e2t} \in [0.0,0.1)$ , orange color for the coexisting cases with  $BR_{e2t} \in [0.1,0.9]$ , and blue color for the early cases with  $BR_{e2t} \in (0.9,1.0]$ . Black star represents coexistence cases further filtered by observational constraints. (d)~(f) are the corresponding kernel density estimate plots of the scatter plots (a)~(c).  $V_{cmax,diff} = V_{cmax,early} - V_{cmax,late}$ .  $SLA_{diff}$  and  $WD_{diff}$  are defined likewise.

### 3.3 XGBoost model performance

434

435

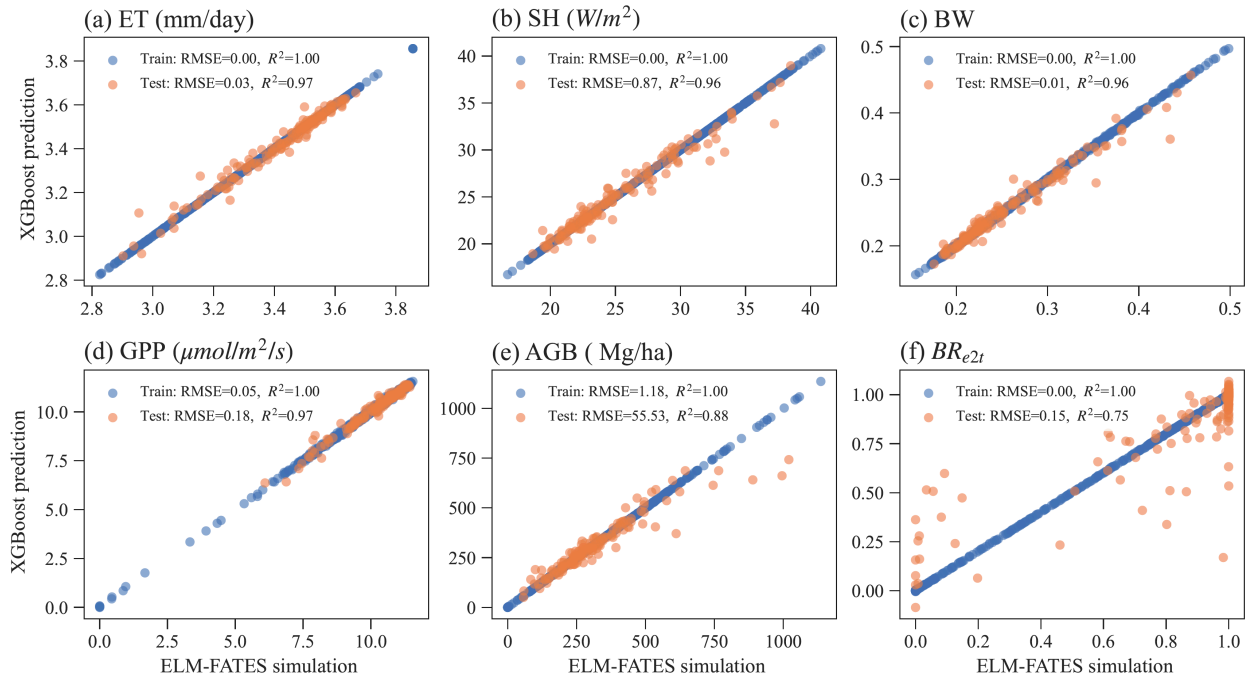
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437

Overall, the XGBoost surrogate models show good performance in predicting ELM-FATES simulations (Figure 5). Based on Exp-CTR (i.e., Par-CTR and Out-CTR), six XGBoost models were trained. In training, the RMSEs for the six models are zero or nearly zero, and  $R^2$ s are close to one. In the testing, four XGBoost models (i.e., XGB\_ET, XGB\_SH, XGB\_BW, XGB\_GPP)



438 still show good performance with small RMSE and large  $R^2$  ( $>0.95$ ). XGB\_AGB shows a little  
 439 degradation with  $R^2$  of 0.88. The performance of XGB\_BR also shows degradation with  $R^2$   
 440 decreasing from 1.0 in training to 0.75 in testing. XGB\_BR cannot well predict the ELM-FATES  
 441 simulated  $BR_{e2t}$  of 0 or 1 when only one PFT survives. This indicates that PFT competition  
 442 processes in ELM-FATES, which determine  $BR_{e2t}$  and AGB, are highly nonlinear and difficult to  
 443 emulate even using a state-of-the-art machine learning algorithm.



444  
 445 Figure 5. The performance of XGBoost surrogate models in the training and testing for  
 446 predicting (a) ET, (b) SH, (c) BW, (d) GPP, (e) AGB, and (f)  $BR_{e2t}$ .

447

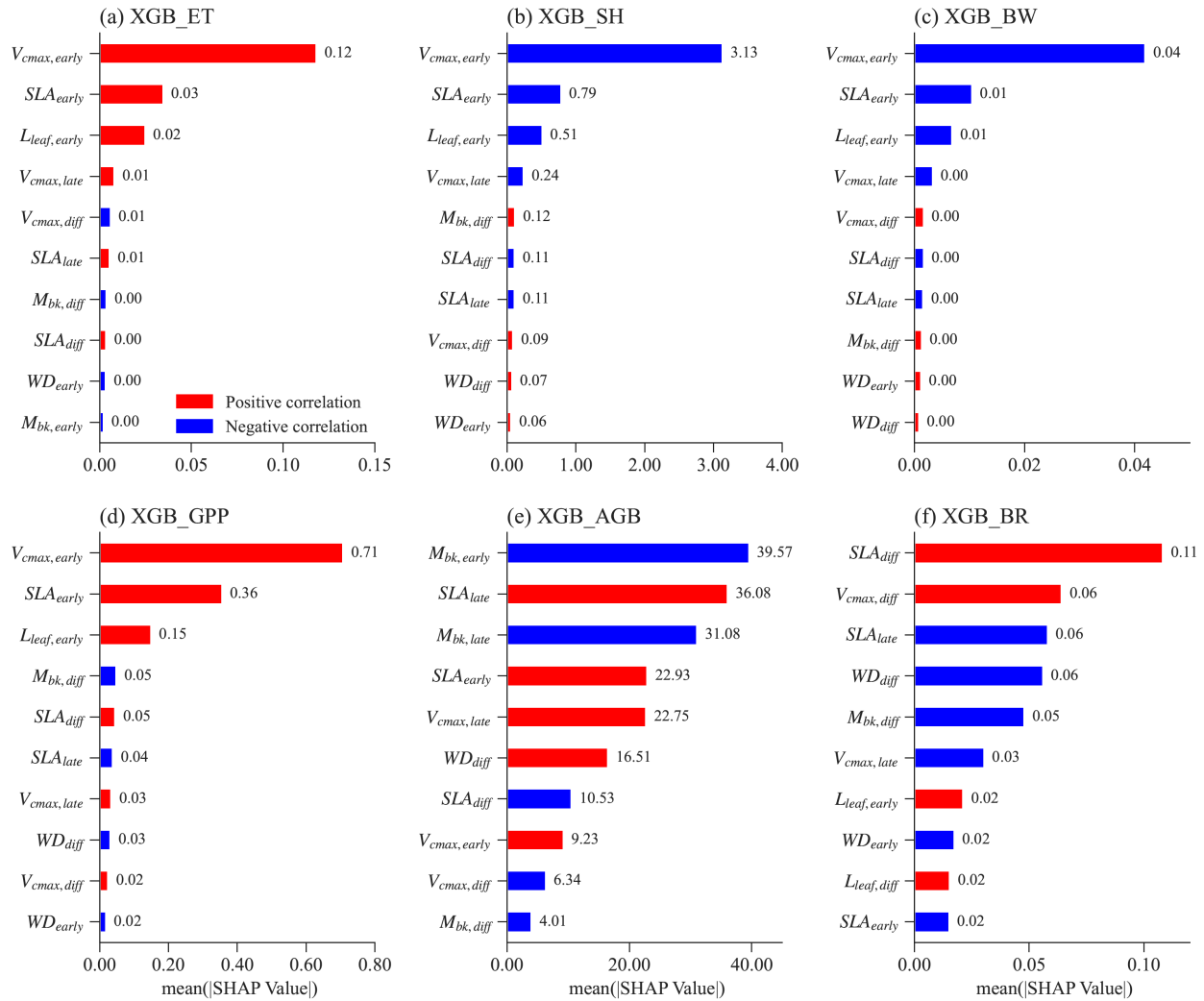
### 448 3.4 SHAP parameter importance analysis

449 Figure 6 shows the feature importance, including parameters and parameter differences, for  
 450 different XGBoost models. Features (on the y-axis) with a higher mean absolute SHAP value (on  
 451 the x-axis) denote a larger contribution to the XGBoost model prediction. The number of most

452 important features is different for predicting ET, SH, BW, and GPP compared with predicting  
453 AGB and  $BR_{e2t}$ .

454 For the XGBoost models that predict ET, SH, BW, and GPP, the top three features have the largest  
455 SHAP values compared to the rest (Figures 6a~5d). Notably, these top three features are the same  
456 and correspond to the early successional PFT, i.e.,  $V_{cmax,early}$ ,  $SLA_{early}$ , and  $L_{leaf,early}$ . Most  
457 ELM-FATES experiments in Exp-CTR used as the training samples for the XGBoost models are  
458 early cases. Therefore, the parameters of early successional PFT have dominant contributions in  
459 the XGBoost model predictions of overall grid-level fluxes. These three parameters are positively  
460 correlated with ET and GPP and negatively correlated with SH and BW (red vs. blue bars in  
461 Figures 6a~d; Figure S4 for more details), reflecting the fundamental carbon metabolism of the  
462 typically dominant early successional plant.

463 For the XGBoost surrogate models of AGB and  $BR_{e2t}$ , more than eight features have large SHAP  
464 values (Figures 6e and 6f). Both early and late successional PFT parameters contribute to  
465 predicting the two variables. Compared with the predictions of ET, SH, BW, and GPP with only  
466 three major features, predicting AGB and  $BR_{e2t}$  is relatively more complex. This is because AGB  
467 and particularly  $BR_{e2t}$  are closely related to the PFT competition process in which both the early  
468 and late PFT traits are crucial. Especially for  $BR_{e2t}$ , the most important features are the parameter  
469 difference between the early and late successional PFTs. For example,  $SLA_{diff}$  is positively  
470 correlated to  $BR_{e2t}$ . Therefore, to have coexisting PFTs with  $BR_{e2t} \in [0.1,0.9]$ , the SLA of two  
471 PFTs should neither be too large nor too small.



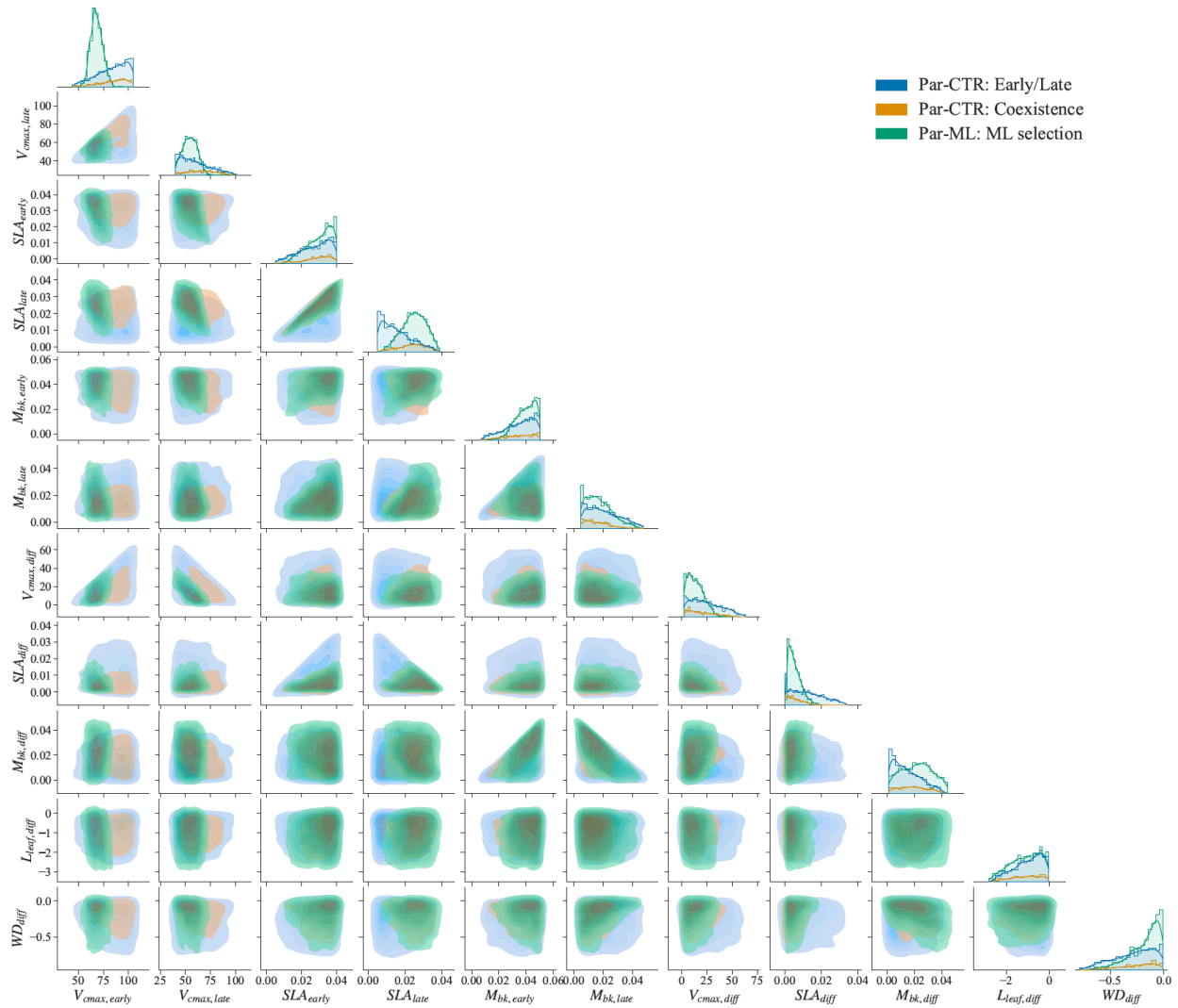
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473 Figure 6. Mean absolute SHAP values for different XGBoost surrogate models for the top ten most  
 474 important features. Absolute SHAP values are sorted in decreasing order from top to bottom. For  
 475 each feature (y-axis) in each XGBoost model, the Spearman correlation coefficient is calculated  
 476 between the feature values and the corresponding SHAP values (Figure S4). The red color means  
 477 that a given feature is positively correlated with the predicting variable, whereas blue denotes a  
 478 negative correlation.

### 479 3.5 XGBoost model parameter selection

480 Using the XGBoost surrogate models, the Par-ML was selected, including 1500 sets of parameters  
481 and the corresponding parameter differences between the early and late successional PFTs (Section  
482 2.4, procedure "P4" in Figure 2). We examined whether Par-ML matches the empirical  
483 relationships shown in Figure 4 (Section 3.2), i.e.,  $SLA_{late} > 0.35 \times SLA_{early} + 0.003$  ,  
484  $V_{cmax,diff} < -4800 \times SLA_{diff} + 100$  , and  $WD_{diff} > 55 \times SLA_{diff} - 1.3$  . In total, 99.1%  
485 (1486 out of 1500) of parameter sets are consistent with the empirical relationships, indicating the  
486 XGBoost models implicitly learned these simple relationships.

487 The parameter distributions of Par-ML show different patterns from the early/late parameters of  
488 Par-CTR (green vs. blue regions in Figure 7), but there are large overlaps between the coexistence  
489 parameters of Par-CTR and Par-ML (orange vs. green regions, e.g., the third column in Figure 7).  
490 This indicates that the XGBoost surrogate models learned to select parameters around the  
491 parameters' space of the coexisting cases. Par-ML also tends to have a smaller parameter difference  
492 between the early and late successional PFTs in terms of  $SLA_{diff}$  and  $V_{cmax,diff}$ . However, Par-  
493 ML also shows different patterns from the coexisting parameters of Par-CTR, probably because  
494 the XGBoost selected parameters were also constrained by multiple observations and implicitly  
495 considered parameter tradeoffs. For example, the  $V_{cmax,early}$  and  $V_{cmax,late}$  of Par-ML are located  
496 in narrower ranges than the coexisting parameters of Par-CTR (first two columns in Figure 6).



497

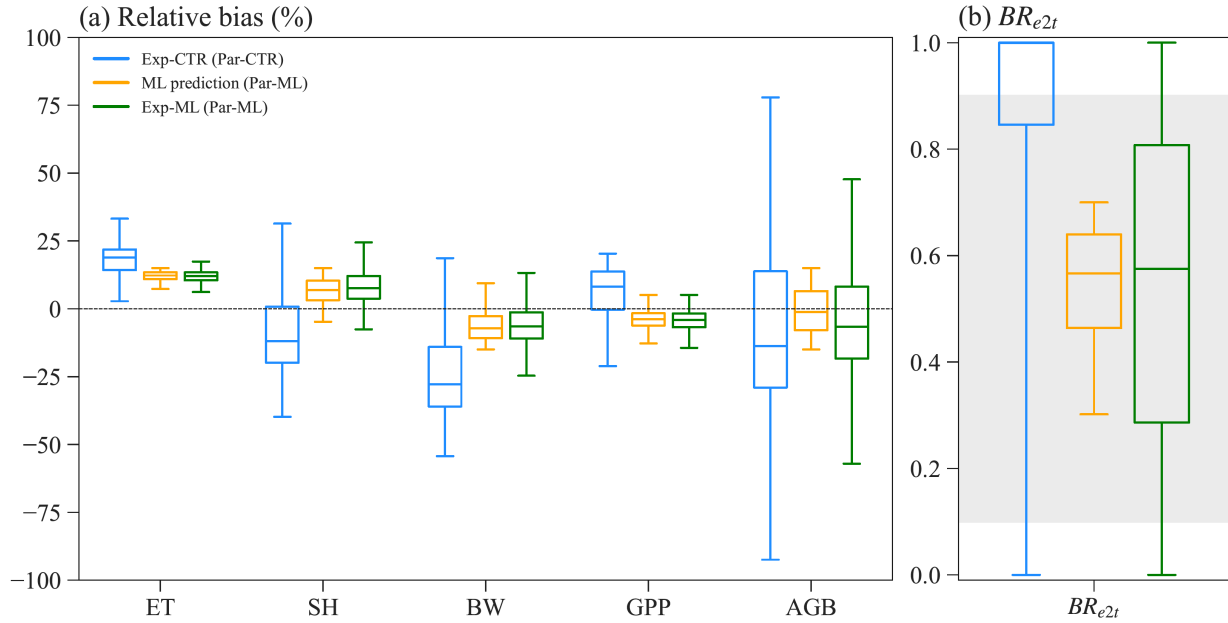
498 Figure 7. Comparison of parameter or parameter difference in Par-CTR vs. Par-ML for eleven  
 499 features. The diagonal plots represent each parameter's distribution, and the rest of the subplots are  
 500 kernel density estimate plots. There are three groups, i.e., blue for the early/late cases of Par-CTR,  
 501 orange for the coexisting cases of Par-CTR, and green for Par-ML selected by XGBoost models.

502

### 503 3.6 Validation of ML selected parameters

504 ELM-FATES simulations of Exp-ML based on the ensemble parameters of Par-ML selected by  
505 the XGBoost surrogate models can better capture the observations and have more coexisting cases  
506 than Exp-CTR (Figure 8). The median values of simulated variables for Exp-ML are closer to  
507 observations with relative biases closer to zero than Exp-CTR (Figure 8a, blue vs. green boxes).  
508 The Exp-ML simulated variables also have more concentrated distributions than Exp-CTR.  
509 Compared to the skewed distribution of  $BR_{e2t}$  in Exp-CTR with a large proportion of early cases,  
510 Exp-ML has a more normally distributed  $BR_{e2t}$  (Figure 8b). Specifically, Exp-ML has about 3.6  
511 times more coexisting cases than Exp-CTR, i.e., 73.1% (1097 out of 1500) in Exp-ML vs. 20.6%  
512 (309 out of 1500) in Exp-CTR (Table S3). After being further constrained by observation (Table  
513 S3), one-third of the experiments (i.e., 495 out of 1500) in Exp-ML remain, and this ratio is 23.6  
514 times more than 1.4% (21 out of 1500) in Exp-CTR.

515 The XGBoost surrogate model predicted variables also match well with those simulated using  
516 ELM-FATES in Exp-ML (Figure 8, orange vs. green boxes), indicating the overall reasonable  
517 accuracy for the XGBoost model predictions. Compared to the ELM-FATES results using Par-  
518 ML, the XGBoost models show better performance for ET, SH, BW, and GPP, but relatively  
519 degraded performance for AGB and  $BR_{e2t}$  (Figure S5). It is consistent with the performance of  
520 the XGBoost models' training and testing results (in Section 3.3).



521  
 522 Figure 8. Comparison between the ELM-FATES simulations for Exp-CTR and Exp-ML. (a)  
 523 Relative bias for simulated ET, SH, BW, GPP, and AGB. (b) Simulated  $BR_{e2t}$ . ML prediction  
 524 represents the selected XGBoost model predictions after filtering with observation and biomass  
 525 ratio (i.e., the XGB\_prds, procedure "P4" in Figure 2).

526

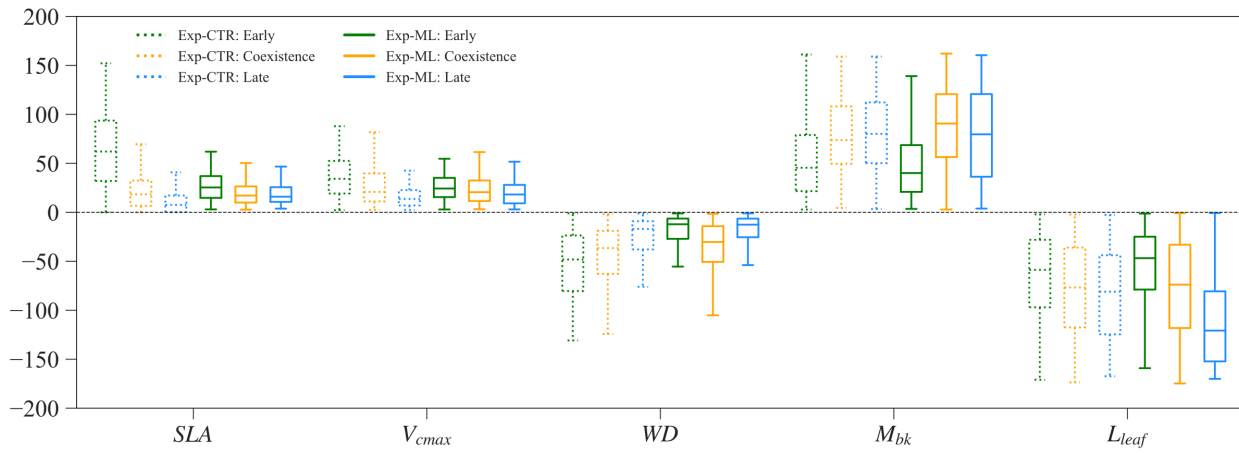
### 527 3.7 Parameter tradeoff for coexistence experiments

528 Parameters of the early and late successional PFTs show tradeoffs for the coexisting experiments.  
 529 Large relative differences in  $SLA$ ,  $V_{cmax}$ , and  $WD$  (more negative) favor the early successional  
 530 PFT, while large relative differences in  $M_{bk}$  and  $L_{leaf}$  favor the late successional PFT. Therefore,  
 531 in Exp-CTR, compared to the early and late cases, the coexisting cases have intermediate relative  
 532 differences in  $SLA$ ,  $V_{cmax}$ ,  $WD$ ,  $M_{bk}$ , and  $L_{leaf}$  (dashed boxes in Figure 9). The coexisting cases  
 533 in Exp-ML have similar patterns with intermediate relative differences in  $SLA$ ,  $V_{cmax}$  and  $L_{leaf}$   
 534 compared to the early and late cases (solid boxes in Figure 9). However,  $M_{bk}$  and especially  $WD$   
 535 show the largest relative difference for the coexisting cases compared to the early and late cases

536 in Exp-ML. These two parameters still show a tradeoff in determining coexisting PFTs, because  
 537 larger  $WD$  favors the early PFT while larger  $M_{bk}$  favors the late PFT.

538

539 In Exp-ML, the parameter spaces of the coexisting cases show large overlaps with the early/late  
 540 cases (Figure S6). There are no simple correlations between these parameters to distinguish the  
 541 coexisting cases from the early and late cases (also see Section 3.2). Although  $WD_{diff}$  of the  
 542 coexisting cases still overlap with the early/late cases, when  $WD_{diff}$  is less than roughly  $-0.4$   
 543 ( $\text{g/cm}^3$ ), only coexisting cases exist (Figure S6). Nevertheless, this rule (i.e.,  $WD_{diff} < -0.4$ ) alone  
 544 cannot ensure PFT coexistence (see Figure 7).



545

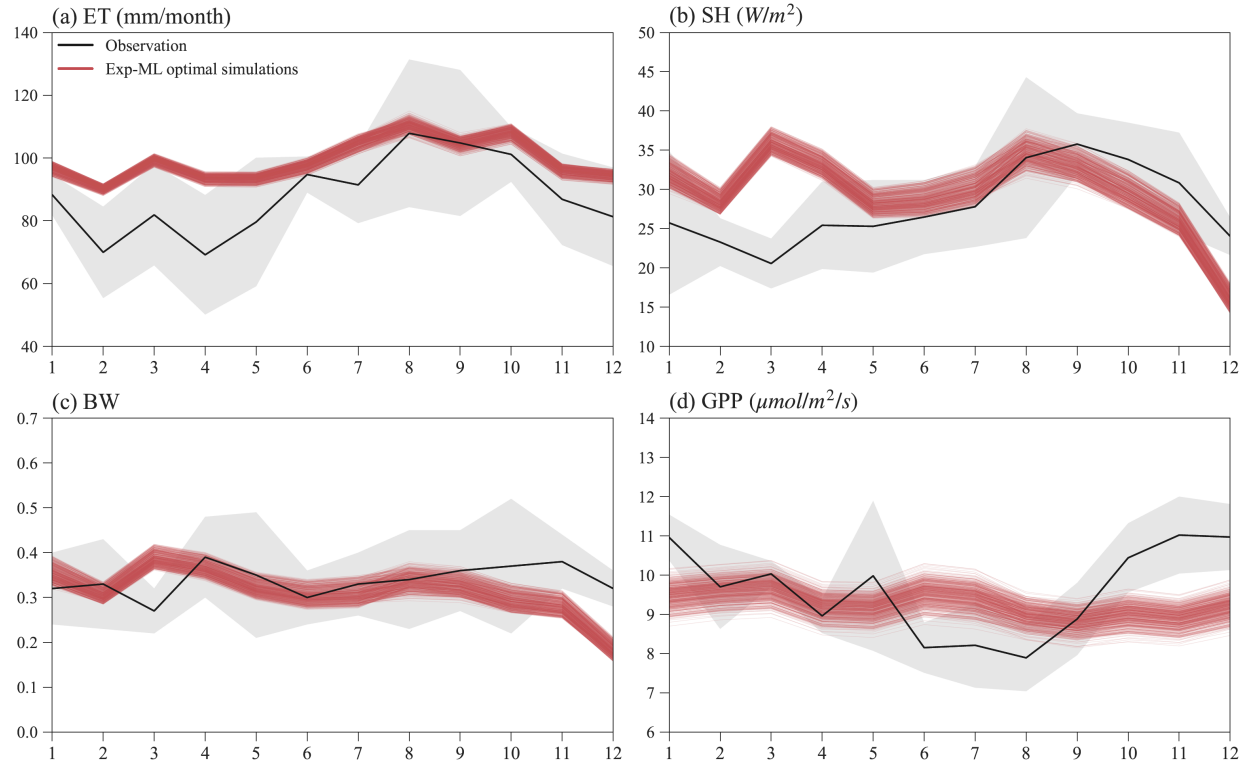
546 Figure 9. Parameter relative difference (%) between early successional PFT and late successional  
 547 PFT for Exp-CTR (box with dash line) and Exp-ML (box with solid line). Parameter relative

548 difference is calculated as, taking SLA as an example,  $\frac{SLA_{early} - SLA_{late}}{(SLA_{early} + SLA_{late})/2} \times 100$  (%).



### 549 3.8 Seasonal variation comparison

550 Figure 10 shows the seasonal variations of ET, SH, BW, and GPP for observations and simulations  
551 of the finally selected 495 experiments in Exp-ML with good model performance (Table S3).  
552 Overall, the simulated ET shows a similar seasonal variation to ET observation (Figure 10a), with  
553 relatively small ET in the wet season (November–May), high ET in the dry season (June–October),  
554 and ET peaks in August. However, compared to the observations, ELM-FATES overestimates ET,  
555 especially during the wet season. The simulated SH also shows a similar seasonal variation with  
556 the SH observation except in March. ELM-FATES overestimated SH from January to May but  
557 underestimated SH from September to December (Figure 10b). Due to the discrepancy between  
558 simulated ET and SH, the model underestimates BW from September to December (Figure 10c).  
559 The simulated GPP has minor seasonal variability compared to the observed GPP. ELM-FATES  
560 overestimates GPP from June–August in the dry season, but underestimates GPP over October–  
561 December. The lower GPP over June–August indicates that plants may be relatively water-stressed  
562 or energy limited during these months. However, the large ET observation over the same period  
563 implies that this site is unlikely water limited or strongly energy limited. The ELM-FATES  
564 simulations also display little water stress year-round (Figure S7). Therefore, there are likely  
565 elements of the seasonal cycle (e.g., phenological responses of photosynthetic capacity) that are  
566 not yet captured here. Additionally, tower estimates of GPP may also have large uncertainties.



567

568 Figure 10. Mean monthly observations and selected optimal ELM-FATES simulations in Exp-ML  
 569 for (a) ET, (b) SH, (c) BW, and (d) GPP. Each red line represents one experiment simulation (four-  
 570 year simulation average). The black curves are monthly climatologic averages from 2000 to 2008,  
 571 and the grey shaded area represents the interannual variabilities (i.e., *mean*  $\pm$   
 572 *standard deviation*).

## 573 4. Discussion

### 574 4.1 Limited guidance of observed trait relationships for PFT coexistence modeling in FATES

575 We found degraded PFT coexistence in ELM-FATES simulation when observed trait relationships  
576 are considered. More specifically, constrained by observed trait relationships, Exp-OBS has fewer  
577 coexisting cases than Exp-CTR which does not consider the observed trait relationships. The  
578 observed trait relationships were derived from site measurements in the species-rich tropical  
579 ecosystem where plant coexistence commonly happens (Kraft et al., 2008), which is expected to  
580 enhance the PFT coexistence simulations. This inconsistency could be due to several possible  
581 reasons. First, ELM-FATES is a typical "trait filtering" model (Fisher et al., 2018), and the realistic  
582 simulation of PFT dynamics largely depends on the fidelity with which trait tradeoff surfaces are  
583 prescribed in the model (Scheiter et al., 2012). Implicit representation of trait tradeoff in the current  
584 ELM-FATES model may not be well balanced, which may differ from the observed trait  
585 relationships that lead to coexistence in the real world (at least for the ecosystem at our study site).  
586 In particular, there may be correlated tradeoffs that are measured (e.g., with below ground  
587 processes, Chitra-Tarak et al. 2021) but not represented in the model. A second reason could be  
588 the mismatch between different spatial scales. The observed trait relationships are derived from  
589 field measurements across tropical forests over a large region with diverse species and climate,  
590 e.g., the relationship in equation (1) is for plant species in Panama. In contrast, ELM-FATES  
591 simulations were conducted at the K34 site scale with specific species composition. Therefore, the  
592 large-scale trait relationships may not reflect the small-scale trait relationships. Wright et al. (2005)  
593 showed that trait relationships fitted for individual sites varied considerably. Third, the observed  
594 trait relationships are based on simplified equations, which may not be able to comprehensively  
595 reflect PFT coexistence. For example, although equation (2) derived from Longo et al. (2020) can

596 reflect the negative relationship between SLA and  $L_{leaf}$ , the  $R^2$  of this equation is about 0.49,  
597 which may not be accurate enough to represent trait relationships. Additionally, these equations  
598 (1)~(3) do not consider the uncertainty of traits covariance. In Koven et al. (2020), the uncertainties  
599 between trait covariance were considered when sampling parameters for FATES experiments.  
600 Furthermore, machine learning models can also be employed to extract the relationships between  
601 plant traits, which can then be incorporated into ELM-FATES and evaluated in future studies.

602

#### 603 **4.2 Advantages of ML surrogate models on improving PFT coexistence modeling**

604 ELM-FATES simulations driven by parameters selected using the XGBoost models essentially  
605 improved PFT coexistence and better captured observations. Compared to the initial Exp-CTR,  
606 which was used to train the XGBoost models, the proportion of coexisting PFTs in Exp-ML  
607 reaches 73.1%, 3.6 times more than 20.6% in Exp-CTR. Further filtering the coexistence  
608 experiments by observations, Exp-ML still has 33.0% of experiments left with good model  
609 performance, 23.6 times that of 1.4% of experiments in Exp-CTR with good performance. Our  
610 ML-based approach also outperforms the empirical correlations built in Section 3.2, which only  
611 yields 32.5% of coexistence experiments and this reduces to 2.3% of experiments if further  
612 constrained by observation. The large proportion of optimal experiments selected by our ML  
613 approach also outperforms previous studies using direct filtering approaches. Buotte et al. (2021)  
614 conducted two stages of experiments to select optimal parameters for CLM-FATES modeling with  
615 two conifer species; only 0.3% (1 out of 360) of the cases met the given criteria in the first stage  
616 experiments, which increased to 5.5% in the second stage experiments. Huang et al. (2020)  
617 conducted CLM-FATES modeling with two tropical PFTs at the Tapajós National Forest sites;  
618 only one parameter set out of seventy (about 1.4%) was selected with reasonable fractions of two

619 PFTs and minor errors compared to observations. In addition, the parameter selection procedures  
620 of these two studies require some degree of subjective decision making and expert knowledge. On  
621 the other hand, our ML-based approach takes a more objective procedure, and little expert  
622 knowledge is required except for the initial determination of the parameter reference ranges.  
623 Importantly, we believe this approach can be repeatable as, e.g., model developments lead to  
624 changes between the parameter values and model predictions of forest structure and function, and  
625 can be used to define constrained ensemble values that will allow assessment of confidence in  
626 model predictions. Even though simulating the coexistence of different plants may not be a big  
627 concern for individual-based VDMs, e.g., LPJmL-FIT (Sakschewski et al., 2015, 2016) and  
628 TROLL (Maréchaux and Chave, 2017), our approach also could be applied to the selection of key  
629 parameters that regulate vegetation dynamics in these models.

630

631 Our study also reproduced the observations satisfactorily. Holm et al. (2020) conducted the ELM-  
632 FATES simulation with only one PFT considered at the same K34 site. Our study yields better or  
633 similar performance in the magnitude of AGB, and the magnitude and seasonal variation of GPP,  
634 ET, SH, and BW (Table 2 and Figure 3 in Holm et al. 2020 vs. Figures 8 and 10 in this study). It  
635 should also be noted that the overestimation of simulated energy fluxes (latent heat and SH) from  
636 January to May could be associated with the energy-related processes (e.g., energy partition,  
637 surface albedo) in ELM-FATES. Other potential reasons could be related to the uncertainties in  
638 atmospheric forcing and the common issue of incomplete energy budget closure at eddy covariance  
639 towers (Wilson et al., 2002; Foken, 2008; Rocha et al., 2009).

640

641 Compared to the predictions of GPP, ET, SH, and BW simulated by ELM-FATES, the XGBoost  
642 surrogate models show slightly degraded performance in predicting the simulated  $BR_{e2t}$  and AGB  
643 (Figures 5 and S5). Three parameters ( $V_{cmax,early}$ ,  $SLA_{early}$ , and  $L_{leaf,early}$ ) mainly control the  
644 predictions of ET, SH, BW, and GPP, while eight features are crucial for predicting AGB and  
645  $BR_{e2t}$ . Even though the XGBoost algorithm has an excellent ability to capture complex nonlinear  
646 relationships, it does not predict well the PFT competition related variables of AGB and  $BR_{e2t}$   
647 because the physical model cannot robustly predict coexisting PFTs due to the higher  
648 dimensionality of predicting PFT composition as compared to other ecosystem variables. Another  
649 important point worth mentioning is the small sample size of coexistence cases in Exp-CTR, with  
650 only 309 cases having  $BR_{e2t}$  in the range of [0.1, 0.9], while the majority of cases are dominated  
651 by either early or late successional PFT. This limited sample size may not provide enough data to  
652 train the XGBoost surrogate model sufficiently for predicting  $BR_{e2t}$  within the range of [0.1, 0.9].  
653 Therefore, further studies are still needed to improve the emulation of PFT competition related  
654 variables. Other approaches that have been applied in VDMs but not specifically for PFT  
655 coexistence modeling, for example, the generalized likelihood uncertainty estimation (GLUE)  
656 approach (Zhang et al., 2022) and the Bayesian model emulation approach (Fer et al., 2018), could  
657 provide alternative ways. Furthermore, we suggest exploring other machine learning algorithms,  
658 such as Gaussian process and neural network algorithms, which may be better suited for capturing  
659 non-linear correlations and learning from sparse data.

660

661 Overall, our study presents a reproducible approach that utilizes machine learning to identify  
662 parameter values that improve model fidelity against observations and promote coexistence  
663 between plant functional types in vegetation demography models across diverse ecosystems. This

664 approach has the potential to enhance the modeling of PFT coexistence in other ecosystems, such  
665 as the mixed conifer forests in Sierra Nevada, California (Buotte et al., 2021), Amazon forests  
666 subject to selective logging (Huang et al., 2020) and tropical forests with heterogeneous soils and  
667 subject to droughts in Panama (Cheng et al., 2021).

668

### 669 **4.3 Trait tradeoffs between coexisting PFTs**

670 Trait-related parameters show tradeoffs between early and late successional PFTs for the ELM–  
671 FATES simulated coexisting experiments. The relative differences between the two PFTs in *SLA*,  
672 *V<sub>cmax</sub>*, and *WD* complementarily coordinate with the relative difference in *M<sub>bk</sub>* and *L<sub>leaf</sub>*, hence  
673 avoiding competitive exclusion (Figure 9). These ELM-FATES reflected tradeoffs are consistent  
674 with the niche-based species coexistence mechanisms of environmental filtering and niche  
675 partitioning (MICHALKO and PEKÁR, 2015; Adler et al., 2013). On the one hand, in the  
676 coexisting cases, the relative differences between the two PFTs' parameters should not be  
677 considerable. For example, a large difference in SLA more likely favors the early cases (green  
678 dash box in Figure 9). This is related to environmental filtering in which coexisting species require  
679 some degree of convergence in strategy to survive and persist under given environmental  
680 conditions (Cadotte and Tucker, 2017; Thakur and Wright, 2017). On the other hand, some degree  
681 of differences should exist between the two PFTs' parameters in the coexisting cases. This is  
682 related to niche partitioning to ensure either differences in resource requirements or differences in  
683 tolerance to surrounding conditions (Kraft et al., 2015; Fowler et al., 2013). Phenomenological  
684 evidence has shown that functional trait variation promotes coexistence or increases species  
685 richness (Uriarte et al., 2010; Angert et al., 2009; Adler et al., 2006; Mason et al., 2012; Ben-Hur  
686 et al., 2012).

687

688 In our ELM-FATES simulations, the primary axis of competition for resources is light. The  
689 tradeoffs between the two PFTs' parameters differentiate their vertical competition in light  
690 absorption, which has been shown to strongly control tropical forest community composition  
691 (Farrion et al., 2016; Poorter et al., 2003). Even though the early PFT has a shallower rooting depth  
692 than the late PFT, there is no critical dry condition during our simulation period (i.e., corresponding  
693 to values of the water stress factor (BTRAN) close to 1.0 in Figure S7). Therefore, competition for  
694 water resource access negligibly contributes to PFT coexistence in this study. Previous tropical  
695 studies also revealed these coexistence mechanisms. At a tropical forest site in eastern Ecuador,  
696 Kraft et al. (2008) found that cooccurring trees are often less ecologically similar, and both  
697 environmental filtering (different topographic habitats of ridgetops vs. valley) and niche  
698 differentiation simultaneously contribute to species coexistence. Swenson & Enquist (2009) also  
699 found that at small spatial scales in a tropical forest, most traits of coexisting species were under-  
700 dispersed, consistent with environmental filtering, while the seed mass and maximum height were  
701 over-dispersed, reflecting niche partitioning.

702

#### 703 **4.4 Limitations and further model development**

704 Some limitations exist in our experiments. Niche partitioning is a critical aspect of promoting  
705 species coexistence, which is closely related to spatial heterogeneity, temporal heterogeneity,  
706 disturbances (e.g., nature enemy, fire), and resource partitioning (Adler et al., 2013). In our current  
707 ELM-FATES simulations, some processes that have been or are being developed in the model are  
708 not considered. These processes include nutrient limitation (Holm et al., 2020), fire disturbance  
709 (Fisher et al., 2015), subsurface lateral flow (Fang et al., 2022), and plant hydraulics (Chitra-Tarak



710 et al., 2021; Li et al., 2021). Ignoring these processes could limit the potential of niche partitioning  
711 among PFT in our ELM-FATES simulations. Topography has been recognized as an essential  
712 spatial heterogeneity factor for tropical forests, but it is not considered in ELM-FATES (Kraft et  
713 al., 2008; Costa et al., 2022). For example, Fang et al. (2022) coupled a three-dimensional  
714 hydrology model (ParFlow) with ELM-FATES and found that lateral flow plays a prominent role  
715 in governing aboveground biomass, and Cheng et al. (2021) also found a critical role for subsurface  
716 hydrology on coexistence. As these processes are added to the model, the reproducibility aspects  
717 of the XGBoost method to identify PFT combinations that match a broad range of criteria will be  
718 particularly important.

719

720 Lacking other features or processes could also affect PFT coexistence in the current FATES. For  
721 example, plant trait plasticity, that plants can adjust their morphological and/or physiological traits  
722 to better adapt to the environment (Nicotra et al., 2010; Bloomfield et al., 2018; McDowell et al.,  
723 2022), is also not well considered in FATES. Leaf traits such as  $V_{cmax}$  and SLA do vary vertically  
724 through the canopy in FATES, via a prescribed relationship described by Lloyd et al., 2010. Liu  
725 and Ng (2019) found that the SLA of a desert shrub is significantly correlated with seasonal water  
726 availability. Additionally, FATES only considers the inter-PFT variance of functional traits (e.g.,  
727 different  $V_{cmax}$  for early and late PFT). However, studies revealed that trait variations commonly  
728 exist within and between species (Wright et al., 2005; Engemann et al., 2016; Meng et al., 2015;  
729 Dong et al., 2020; Siefert et al., 2015), which play a vital role in maintaining plant diversity (Violle  
730 et al., 2012; Lu et al., 2017). Reproductive features that enhance competitive exclusion tendencies  
731 have been illustrated to affect coexistence (Maréchaux and Chave, 2017; Fisher et al., 2018).  
732 Hanbury-Brown et al. (2022) discussed the importance of the representation of forest regeneration,

733 including improving parameters and algorithms for reproductive allocation, dispersal, seed  
734 survival and germination, environmental filtering in the seedling layer, and tree regeneration  
735 strategies adapted to wind, fire, and anthropogenic disturbance regimes. Besides, both growth-  
736 survival and stature-recruitment tradeoffs are critical to accurately predict successional patterns in  
737 tropical forest structure and competition (see details in Rüger et al., 2020), which should also be  
738 better considered in future model development. Furthermore, measured plant traits are increasingly  
739 available, e.g., the TRY datasets (Kattge et al., 2020) can be used to improve the model process  
740 and parameterizations. Future studies on properly and adequately using these datasets to guide  
741 VDM parameterizations are advocated.

742

#### 743 **4.5 Enhancing VDM prediction with machine learning**

744 We provide a brief overview of how machine learning can be applied to improve the modeling of  
745 plant dynamics, specifically in the context of vegetation demographic models. Firstly, ML can be  
746 used to derive trait parameter values. For instance, in this study, ML could be applied to replace  
747 the simple equations to derive the relationships between measured traits (Section 4.1). By  
748 integrating multiple datasets, including in situ measurements, atmospheric forcing, and remote  
749 sensing, ML could derive the spatial patterns and temporal variations of trait parameters for use in  
750 large-scale VDM modeling. Secondly, ML can be utilized to optimize parameters by developing  
751 surrogate models that emulate the relationships between the parameters and the VDM simulations,  
752 and using the surrogate models to identify optimal parameter values. This application has  
753 demonstrated success in this study and previous studies (e.g., Tsai et al., 2021; Dagon et al., 2020;  
754 Watson-Parris et al., 2021). Another benefit of using ML in VDMs is the ability to develop  
755 benchmark datasets. For example, studies have successfully employed ML to derive AGB datasets

756 for various ecosystems (Morais et al., 2021; Zhang et al., 2020; Li et al., 2020; Bispo et al., 2020;  
757 Pham et al., 2020). These datasets can serve as benchmarks to evaluate the accuracy of VDM  
758 simulations. Lastly, ML can be used to replace semiempirical sub-models with little theoretical  
759 bases in DGVMs (Reichstein et al., 2019). For example, accurately modeling wildfire using  
760 process-based wildfire models integrated in DGVMs remains challenging. However, ML-based  
761 wildfire models have shown advantages in accuracy and computational efficiency (Rodrigues and  
762 Riva, 2014; Jain et al., 2020; Sayad et al., 2019), and have the potential to be employed in Earth  
763 system models to improve wildfire simulations (e.g., Zhu et al., 2022).

764 **5. Conclusions**

765 In this study, we explored two possible solutions to improve PFT coexistence modeling in a cohort-  
766 based model (ELM-FATES): (1) using plant trait relationships established from field  
767 measurements and (2) using machine learning surrogate models to optimize trait parameter values.  
768 Three ensembles of ELM-FATES experiments were conducted over a tropical forest site at  
769 Manaus, Brazil. We found that considering the observed trait relationships (Exp-OBS) slightly  
770 improves the simulations of water (ET), energy (SH and BW), and carbon (GPP, AGB) when  
771 compared against observations, but degrades the simulation of PFT coexistence. Based on Exp-  
772 CTR, the ML surrogate models were built to optimize the ELM-FATES parameters by integrating  
773 the observations (i.e., ET, SH, BW, GPP, and AGB) and PFT coexistence criteria. Exp-ML, with  
774 parameters selected by the ML surrogate models, vastly improves the simulation of PFT  
775 coexistence, and also better reproduces the annual means and seasonal variations of ET, SH, BW,  
776 GPP, and the field inventory of AGB. This study demonstrates the benefits of using machine  
777 learning models to improve the modeling of PFT coexistence in ELM-FATES, with important  
778 implications for modeling the response and feedback of ecosystem dynamics to climate change.  
779 Our results also suggest that the incorporation of additional mechanisms into ELM-FATES is  
780 essential for robust modeling of coexisting PFTs.

781

782 *Code and Data Availability.* The ELM-FATES source code, surface and domain data, forcing data,  
783 and ML codes used in this study are archived on Zenodo (<https://doi.org/10.5281/zenodo.7730685>).  
784 The observational reference datasets of GPP, ET, SH, BW, and AGB are obtained from Holm et  
785 al. (2020). The forcing data are available from Oak Ridge National Laboratory Distributed Active  
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788

789 *Author contributions.* LL and YF designed and conducted the experiments and analysis, and  
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