### **Representation of phosphorus cycle in Joint UK Land** 1 **Environment Simulator (vn5.5 JULES-CNP)** 2

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#### 20 Abstract

- Most Land Surface Models (LSMs), the land components of Earth system models (ESMs), include
- representation of nitrogen (N) limitation on ecosystem productivity. However only few of these models have
- incorporated phosphorus (P) cycling. In topical ecosystems, this is likely to be important as N tends to be
- 21 22 23 24 25 26 27 28 29 abundant but the availability of rock-derived elements, such as P, can be very low. Thus, without a
- representation of P cycling, tropical forest response in areas such as Amazonia to rising atmospheric CO<sub>2</sub>
- conditions remains highly uncertain. In this study, we introduced P dynamics and its interactions with the N and
- carbon (C) cycles into the Joint UK Land Environment Simulator (JULES). The new model (JULES-CNP)
- includes the representation of P stocks in vegetation and soil pools, as well as key processes controlling fluxes
- 30 between these pools. We evaluate JULES-CNP using in situ data collected at a low fertility site in the Central
- Amazon, with a soil P content representative of 60% of soils across the Amazon basin, to parameterise, calibrate
- and evaluate JULES-CNP. Novel soil and plant P pool observations are used for parameterisation and calibration and the model is evaluated against C fluxes and stocks, and for those soil P pools not used for
- 31 32 33 34 35 36 37 38 39 parameterisation/calibration. We then apply the model under elevated CO<sub>2</sub> (600 ppm) at our study site to
- quantify the impact of P limitation on CO<sub>2</sub> fertilization. We compare our results against current state of the art
- CNP models using the same methodology that was used in the AmazonFACE model intercomparison study. The
- model is able to reproduce the observed plant and soil P pools and fluxes used for evaluation under ambient
- CO<sub>2</sub>. We estimate P to limit net primary productivity (NPP) by 24% under current CO<sub>2</sub> and by 46% under
- elevated CO<sub>2</sub>. Under elevated CO<sub>2</sub>, biomass in simulations accounting for CNP increase by 10% relative to at
- 40 contemporary CO<sub>2</sub>, although it is 5% lower compared with CN and C-only simulations. Our results highlight the
- 41 potential for high P limitation and therefore lower CO<sub>2</sub> fertilization capacity in the Amazon forest with low
- 42 fertility soils.
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#### 44 1. Introduction

45

46 Land ecosystems currently take up about 30% of anthropogenic CO<sub>2</sub> emissions (Friedlingstein et al., 2020), thus 47 buffering the anthropogenic increase in atmospheric CO<sub>2</sub>. Tropical forests play a major role in the land C cycle, 48 account for about half of global net primary production (NPP)(Schimel et al., 2015), and store the highest above 49 ground carbon among all biomes (Pan et al., 2011; Mitchard, 2018).

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51 The C sink capacity of tropical forests may be constrained by nutrient availability for plant photosynthesis and 52 growth (Vitousek and Howarth, 1991; Elser et al., 2007; LeBauer and Treseder, 2008) via either P (Nordin, 53 Högberg and Näsholm, 2001; Shen et al., 2011) and N related processes (DeLuca, Keeney and McCarty, 1992; 54 Perakis and Hedin, 2002). Global process-based models of vegetation dynamics and function suggest a 55 continued land C sink in the tropical forests, largely attributed to the CO<sub>2</sub> fertilization effect (Sitch *et al.*, 2008; 56 Schimel, Stephens and Fisher, 2015; Koch, Hubau and Lewis, 2021). However, many of these models typically 57 do not consider P constraints on plant growth (Fleischer et al., 2019), which is likely to be an important limiting 58 nutrient in tropical ecosystems, characterised by old and heavily weathered soils. The importance of nutrient 59 cycling representation in Earth System Models (ESMs), and the lack thereof, was highlighted by Hungate et al. 60 (2003) and Zaehle and Dalmonech (2011), showing the significance of nitrogen inclusion in ESMs for 61 generating more realistic estimations of the future evolution of the terrestrial C sink. However, in the Coupled 62 Climate C Cycle Model Inter-comparison Project (C4MIP), none of the participating ESMs included N 63 dynamics (Friedlingstein et al., 2006). Seven years later, for the update in CMIP5 (Anav et al., 2013), three 64 models out of eighteen with N dynamics were included (Bentsen et al., 2013; Long et al., 2013; Ji et al., 2014). Although much progress has been made in the inclusion of an N cycle in ESMs so far, none of the CMIP5

- 65 66 models included P cycling and in the most recent CMIP6, only one model includes P (ACCESSESM1.5 model)
- 67 (Arora et al., 2020).
- 68

69 The long history of soil development in tropical regions which involves the loss of rock-derived nutrients 70 through weathering and leaching on geologic timescales (Vitousek et al., 1997, 2010) results in highly 71 72 73 74 weathered soils. Soil P is hypothesized to be among the key limiting nutrients to plant growth in tropical forests (Vitousek et al., 1997, 2010; Hou et al., 2020), unlike temperate forest where N is hypothesised to be the main constraint(Aerts and Chapin, 1999; Luo et al., 2004). Low P availability in tropical soils is related to the limited un-weathered parent material or organic compounds as source of P (Walker and Syers, 1976), active sorption 75 (Sanchez, 1977) and high occlusion (Yang and Post, 2011) which further reduce plant available P. Although N 76 77 78 79 limitation can impact the terrestrial C sink response to increasing atmospheric CO<sub>2</sub> by changing plant C fixation capacity (Luo et al., 2004), this can be partially ameliorated over time by input of N into the biosphere via the continuous inputs of N into ecosystems from atmospheric deposition and biological N fixation (Vitousek et al., 2010). P-limitation is pervasive in natural ecosystems (Hou et al., 2020) and the lack of large P inputs into 80 ecosystems, especially those growing on highly weathered soil, may make P limitation a stronger constraint on 81 82 83 ecosystem response to elevated CO2 (eCO2) than N (Gentile et al., 2012; Sardans, Rivas-Ubach and Peñuelas, 2012). This causes considerable uncertainty in predicting the future of the Amazon forest C sink (Yang et al., 2014).

84

85 There is evidence to suggest P limitation on plant productivity in the Amazon forest (Malhi, 2012) where it has 86 been shown that the younger, more fertile west and south-west Amazon soils have higher tree turnover (Phillips 87 et al., 2004; Stephenson and Van Mantgem, 2005) and stem growth rates (Malhi et al., 2004) and lower above 88 ground biomass (Baker et al., 2004; Malhi et al., 2006) compared to their central and eastern counterparts. Total 89 soil P has been found as the best predictor of stem growth (Ouesada et al., 2010) and of total NPP (Aragão et 90 al., 2009) across this fertility gradient, and foliar P is positively related to plant photosynthetic capacity ( $V_{\rm cmax}$ 91 and  $J_{cmax}$ ) in these forests (Mercado *et al.*, 2011).

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92 93 94 95 However, modelling studies are unable to reproduce observed spatial patterns of NPP and biomass in the Amazon due to missing information on nutrient availability and soil fertility impact on productivity (Wang, Law

- and Pak, 2010; Vicca et al., 2012; Yang et al., 2014) and due to the lack of inclusion of soil P constraints on
- 96 plant productivity and function. Nevertheless, some modelling works have focused on improving process and
- 97 parameter representation using the observational data of spatial variation in woody biomass residence time
- 98 (Johnson et al., 2016), soil texture and soil P to parameterise the maximum RuBiCo carboxylation capacity
- 99  $(V_{cmax})$  (Castanho *et al.*, 2013). Results from these studies successfully represent observed patterns of Amazon
- 100 forest biomass growth increases with increasing soil fertility. However, the full representation of these
- 101 interactions and the impact of the soil nutrient availability on biomass productivity is still missing in most of
- 102 ESMs. 103

104 So far, several dynamic global vegetation models have been developed to represent P cycling within the soil 105 (Yang et al., 2013; Haverd et al., 2018) and between plant and soils for tropical forests particularly (Yang et al.,

106 2014; Zhu et al., 2016; Goll et al., 2017). Furthermore, a comprehensive study included several models with C-

107 N-P cycling and their feedbacks on the atmospheric C fixation and biomass growth in Amazon forests under

108 ambient and eCO<sub>2</sub> conditions (Fleischer et al., 2019). Despite these developments, data to underpin them and

109 their projections, particularly for the tropics, is sparse and remains challenging particularly for the Amazon

- 110 forest (Reed et al., 2015; Jiang et al., 2019). Moreover, due to the lack of detailed measurements, the P-related 111 processes such as ad/desorption and uptake represented in these models are under-constrained and likely
- 112 oversimplified, thus the future predictions of Amazon forest responses to eCO2 and climate change are
- 113 uncertain. To fill this gap, in this study, we will use data collected as part of the Amazon Fertilization
- 114 Experiment (AFEX), the first project that focuses on experimental soil nutrient manipulation in the Amazon,
- 115 with a comprehensive data collection program covering plant ecophysiology, C stocks and fluxes, soil processes

116 including P stocks. Thus, our model parameterization compared to prior P modelling studies includes detailed P

117 processes representation using the site measurements.

# 118

119 Here, we describe the development and implementation of the terrestrial P cycle in the Joint UK Land 120 Environment Simulator (JULES) (Clark et al., 2011), the land component of the UK Earth System Model 121

(UKESM), following the structure of the prior N cycle development (Wiltshire et al., 2021). The model

122 (JULES-CNP) is parameterized and calibrated using novel in situ P soil and plant data from a well-studied forest

- 123 site in Central Amazon near to Manaus, Brazil with soil P content representative of 60% of soils across the 124 Amazon basin. We then evaluate the model against carbon stocks and fluxes from data sets from our study site 125 and the nearby K34 field site. To test the model, we followed the protocol of Fleischer et al., (2019), to predict 126 nutrient limitations on land biogeochemistry under ambient and eCO<sub>2</sub>. Predictions of the CO<sub>2</sub> fertilization effect 127 in JULES-CNP are compared to those in current versions of the model with coupled C and N cycles (JULES-CN) and with C cycle only (JULES-C).
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# 2. Material and methods

#### 133 2.1 JULES

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135 JULES is a process-based model that integrates water, energy, C cycling (JULES-C) (Clark et al., 2011) and N 136 cycling (JULES-CN) (Wiltshire *et al.*, 2021) between the atmosphere, vegetation and soil (Best *et al.*, 2011; 137 Clark et al., 2011). Vegetation dynamics are represented in JULES using the TRIFFID model, using nine 138 distinct plant functional types (PFTs) (tropical and temperate broadleaf evergreen trees, broadleaf deciduous 139 trees, needle-leaf evergreen and deciduous trees, C3 and C4 grasses, and evergreen and deciduous shrubs), as 140 well as height competition (Harper et al., 2016). JULES simulates Gross Primary Productivity (GPP) based on a 141 coupled photosynthesis and water balance scheme, from which autotrophic respiration for each living tissue 142 (leaf, wood, root) is subtracted to estimate NPP. In JULES we assume a process-based leaf-level photosynthesis 143 scaled up to the canopy. Therefore, in JULES CNP in order to keep consistency with JULES C-CN, we also 144 assume a multi-level canopy, and leaf N and P in exponentially decreases through the canopy (CanRadMod 6) 145 (Clark et al., 2011). NPP is then allocated to increase tissue C stocks and to spread, i.e., expand the fractional 146 coverage of the PFT. The resultant PFT fractional coverages depend in addition on competition across PFTs for 147 resources, e.g., light. Tissue turnover and vegetation mortality add C into the litter pools. Representation of soil 148 organic C (SOC) follows the RothC equations (Jenkinson et al., 1990; Jenkinson and Coleman, 2008) defining 149 four C pools: decomposable plant material (DPM) and resistant plant material (RPM), which receive direct input 150 from litterfall, and microbial biomass (BIO) and humified material (HUM) which receive a fraction of 151 decomposed C from DPM and RPM which is not released to the atmosphere. The limitation of N on SOC is 152 applied to the vegetation and soil components using a dynamic C:N ratio to modify the mineralization and 153 immobilization processes as described in Wiltshire et al., (2021). Note that the soil component of JULES-CN 154 can be run either as a single box model or vertically resolved over soil depth (JULES-CN layered), and in this 155 paper we build upon the vertically resolved version described in Wiltshire et al. (2021). 156

#### 157 2.2 JULES-CNP

# 158

159 JULES-CNP includes the representation of the P cycle in JULES version (vn5.5). It includes P fluxes within the

- 160 vegetation and soil components, and the specification of P pools and processes related to P cycling within the 161 soil column (Figure 1). A parent material pool is introduced to consider the input of weathered P. The adsorbed,
- 162 desorbed and occluded fractions of P for both organic and inorganic P are also represented. However, except for
- parent material and occluded P pools, all other pools are estimated at each soil layer. The description of changes 163

164 in pools and associated relative fluxes are explained in detail in the next sections. However, despite JULES-CN

- 165 that includes N leaching and deposition, P leaching and deposition are omitted in the current version of JULES-CNP.
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- 167
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### Figure.1 – JULES CNP model scheme 171

#### 172 P pools 2.2.1 173

174 JULES represents eight P pools comprising organic and inorganic P: in plant P ( $P_p$ ) and soil pools (in each soil 175 layer (n)): litter P (Pol), soil organic P (Pos), soil inorganic P (Pin), organic sorbed (Porg-sorp), inorganic sorbed 176 (Pinorg-sorp), parent material (Ppm) and occluded (Pocc) P comprised of both organic and inorganic P. All pools 177 178 are in units of kg P m<sup>-2</sup> (Fig 1, Tables 1 and 2).

179 Plant P pool is composed of leaf  $(P_{leaf})$ , fine root  $(P_{root})$  and stem together with coarse root  $(P_{stem})$ , which are 180 related to their associated C pools ( $C_{leaf}$ ,  $C_{root}$ ,  $C_{stem}$ ) in (kg C m<sup>-2</sup>) and <u>fixed</u> C to P ratios 181  $(C: P_{leaf}, C: P_{root}C: P_{stem})$  as follows:

$$\begin{array}{l}
182\\
183\\
P_{leaf} = \frac{c_{leaf}}{c:P_{leaf}}\\
184
\end{array}$$
(eq.1)

185 
$$P_{root} = \frac{C_{root}}{C:P_{root}}$$
(eq.2)

$$\begin{array}{l}
186\\
187\\
P_{stem} = \frac{c_{stem}}{c:P_{stem}}\\
188
\end{array}$$
(eq.3)

189 Therefore, the plant P pool  $(P_p)$  is the sum of all vegetation P pools as follows: 190

$$\begin{array}{ccc}
191 \\
192 \\
192 \\
\end{array} P_p = P_{leaf} + P_{root} + P_{stem} \\
(eq.4)
\end{array}$$

193 Description of the plant P pool (P<sub>p</sub>) follows Zhu *et al.*, (2016) and is estimated as the difference between the 194 input, plant uptake  $F_P{}^{Up}(eq.\underline{26})$  and output of this pool, plant litter flux  $F_P{}^{lit}(eq.\underline{28})$ , with both fluxes 195 expressed in kg P m<sup>-2</sup> yr<sup>-1</sup> as follows: 196

$$\frac{197}{4t} = F_p^{\ up} - F_p^{\ lit}$$
(eq.5)  
198

199 The litter P pool  $(P_{O_l})$  is estimated as a sum of P<sub>DPM</sub> and P<sub>RPM</sub> pools. Each pool is formed by the fluxes of plant 200 litter input  $(F_P^{lit})$  and the outgoing decomposed P  $(dec_P^{lit})$  both expressed in kg P m<sup>-2</sup> yr<sup>-1</sup> (eq.<u>28-29</u>).

Furthermore, the plant litter input is modified based on the plant type material ratio  $\alpha$  (in order to distribute the litter input based on the DPM/RPM fraction) as follows:

$$\frac{dP_{DPM}}{dt} = F_{P_n}^{lit} \times \alpha - dec_{P_{DPM,n}}$$
(eq.6)
205

$$206 \quad \frac{dP_{RPM}}{dt} = F_{Pn}^{\ lit} \times (1 - \alpha) - dec_{PRM,n} \tag{eq.7}$$

$$\begin{array}{ccc}
207 \\
208 \\
209
\end{array} P_{o_l} = \sum_{n=1}^{N} P_{DPM_n} + \sum_{n=1}^{N} P_{RPM_n} \\
(eq.8)
\end{array}$$

The soil organic pool  $(P_{0_S})$  is represented as the sum of P<sub>BIO</sub> and P<sub>HUM</sub>. These pools are estimated from the difference between P inputs from total immobilized  $(F_{immob_P})$  distributed between BIO and HUM based on fixed fraction (0.46 for BIO, 0.54 for HUM) (Jenkinson *et al.*, 1990; Jenkinson and Coleman, 2008) and desorbed P  $F_{P_{0_S}}^{desorp}$  and P outputs from mineralized  $(F_{minl_P})$ , and adsorbed P fluxes  $(F_{P_{0_S}}^{sorp})$  (adsorption: eq. 40 and desorption: eq.41) with all fluxes expressed in kg P m<sup>-2</sup> yr<sup>-1</sup> as follows:

215  
216 
$$\frac{dP_{BIO}}{dt} = 0.46 \times F_{immob_{P_n}} + F_{P_{O_S}} \frac{desorp}{BIO,n} - F_{minl_{P_{BIO,n}}} - F_{P_{O_S}} \frac{sorp}{BIO,n}$$
 (eq.9)  
217

$$\frac{dP_{HUM}}{dt} = 0.54 \times F_{immobP_n} + F_{P_{O_{S}HUM,n}} - F_{minlP_{BIO,n}} - F_{P_{O_{S}HUM,n}}$$
(eq.10)  
219

$$P_{O_s} = \sum_{n=1}^{N} P_{BIO_n} + \sum_{n=1}^{N} P_{HUM_n}$$
(eq.11)

220

221 222

228 229

223 Description of the inorganic sorbed P pool ( $P_{inorg-sorp}$ ) follows Wang *et al.*, (2007) and is represented as the 224 difference between the input flux of inorganic sorption ( $F_{P_{in}}^{sorp}$ ) (eq. 37) and output fluxes of inorganic 225 desorption ( $F_{P_{in}}^{desorp}$ ) (eq. 38) and occluded P( $F_{P}^{occ}$ ) (eq. 39), with all fluxes expressed in kg P m<sup>-2</sup> yr<sup>-1</sup> as 226 follows:

$$\frac{dP_{inorg-sorp}}{dt} = \sum_{n=1}^{N} F_{P_{in}} \sum_{n=1}^{N} F_{P_{$$

 $\begin{array}{c} 230 \\ 231 \\ 231 \\ 232 \\ 233 \end{array} \xrightarrow{\text{Descripting of the occluded } (P_{occ}) P \text{ pool follows Wang et al., (2007) and Hou et al., (2019) and is represented} \\ as the sum of input fluxes of occluded P from both organic <math>(F_P^{\ or-occ})$  (eq. 42) and inorganic P pools  $(F_P^{\ occ})$  expressed in kg P m<sup>-2</sup> yr<sup>-1</sup>, as follows:

234 
$$\frac{dP_{occ}}{dt} = \sum_{n=1}^{N} F_{P_n}^{occ} + \sum_{n=1}^{N} F_{P_n}^{or-occ}$$
(eq.13)

236 Descripting of the organic sorbed P pool ( $P_{org-sorp}$ ) follows Wang *et al.*, (2007) and is represented as the 237 difference between the input flux of organic sorption ( $F_{P_{o_{s_n}}}^{sorp}$ ) and output fluxes of organic desorption 238 ( $F_{P_{o_{s_n}}}^{desorp}$ ) and occluded P( $F_{P_n}^{occ}$ ), with all fluxes expressed in kg P m<sup>-2</sup> yr<sup>-1</sup> as follows: 239

$$\begin{vmatrix} 240 & \frac{dP_{org-sorp}}{dt} = \sum_{n=1}^{N} F_{P_{O_{S_n}}} - \sum_{n=1}^{N} F_{P_{O_{S_n}}} - \sum_{n=1}^{N} F_{P_{O_{S_n}}} - \sum_{n=1}^{N} F_{P_{n}} \\ eq.14 \\ 241 \\ 242 \\ 243 \end{vmatrix}$$

244 Descripting of P from parent material  $(P_{pm})$  pool follows Wang *et al.*, (2007) and depends on the weathering 245 flux  $(F_{p}^{w})$  (eq. 43) in kg P m<sup>-2</sup> yr<sup>-1</sup> as follows:

246  
247 
$$\frac{dP_{pm}}{dt} = -\sum_{n=1}^{N} F_{P_n}^{w}$$
 (eq.15)

248 249

## 250 2.2.2. <u>C and P fluxes</u>

251 252 NPP in JULES is calculated as the difference between GPP and autotrophic respiration. In JULES-CNP, 253 potential NPP represent the amount of C, available for tissue growth (C density increase) on a unit area, and 254 spreading (vegetation cover increase as a result of reproduction and recruitment), ie to increase the area covered 255 by the vegetation type, assuming no nutrient limitation. The reported NPP in the literature often includes other C 256 fluxes related to the exudates, volatiles production and non-structural carbohydrates (Malhi et al., 2009; Chapin 257 et al., 2011; Walker et al., 2021) which are challenging to measure (Malhi, Doughty and Galbraith, 2011). 258 Therefore, actual NPP is for our purposes equal to Biomass Production (BP), and is calculated as potential NPP 259 minus excess C (lost to the plant through autotrophic respiration), with the latter the C that cannot be used to 260 growth new plant tissue due to insufficient plant nutrient supply. Hence, if the system is limited by the 261 availability of N and/or P, NPP will be adjusted to match the growth that can be supported with the limited N or 262 P supply, with any excess carbohydrate lost through excess C. 263 The total excess C term ( $\psi_{\star}$ ) (kg C m<sup>-2</sup> yr<sup>-1</sup>) is calculated as: 264 265  $\Psi_t = \Psi_g + \Psi_s$ (eq.16) 266 where  $\psi_g$  and  $\psi_s$  are the excess C fluxes due to growth (g) and spread (s) and are assumed to be rapidly respired 267 268 by plants. 269 270 Therefore, BP is calculated as the difference between potential NPP ( $\Pi_c$ ) and total excess C: 271 272  $BP = \Pi_c - \Psi_t$ (eq.17)273 The litter production in JULES before limitation is estimated based on the as follows: 274 275 276  $F_{C_n}^{lit} = \gamma_{leaf} C_{leaf} + \gamma_{root} C_{root} + \gamma_{wood} C_{wood}$ (eq.18) 277 278 where  $\lambda$  is the leaf, root and stem re-translocation (at daily timestep) coefficient (Clark *et. al.*, 2011) and  $\gamma$  is a 279 temperature dependent turnover rate representing the phenological state (Clark et al., 2011). P limitation is 280 applied on the C litter production similar to the N scheme of JULES (JULES-CN) (Wiltshire et al., 2021). In 281 JULES-CN the N limitation effect on the litter production is captured by estimating the available C for litter 282 production as a difference between the NPP and excess C (Wiltshire et al., 2021). 283 284 Similar to other P-enabled models (Yang et al., 2014; Goll et al., 2017), JULES-CNP follows the same structure 285 as its N model component. Description of the plant P and N demand follow Wang et al., (2007) and are represented by the sum of demand  $(\phi_t)$  to sustain growth (P-related:  $(\phi_{g_P})$ , N-related:  $(\phi_{g_N})$ ) and to sustain 286 vegetation spreading (to increment PFT fractional coverage) (P-related:  $(\phi_{S_P})$ , N-related:  $(\phi_{S_N})$ ) and is 287 288 expressed in (P-related in kg P m<sup>-2</sup> yr<sup>-1</sup>; N-related in kg N m<sup>-2</sup> yr<sup>-1</sup>). The total demand for growth ( $\phi_a$ ) and 289 spreading  $(\phi_s)$  is controlled by the dominant demand between P  $(\phi_{a_P})$  and N  $(\phi_{a_N})$  as follows: 290

$$323 F_p {}^{up}_n = \begin{cases} \emptyset_t & \emptyset_t \le u^{max} \\ u^{max} & \emptyset_t > u^{max} \end{cases}$$
(eq.26)

Description of the plant P uptake  $(F_p^{up})$  varies spatially depending on the root uptake capacity  $(u^{max})$  followed by Goll *et al.*, (2017). Therefore, in regions with limited P supply, the plant P uptake is limited to the  $u^{max}$  and consequently impacts the excess C and BP. The root uptake capacity depends on the maximum root uptake capacity  $(v_{max})$  (kg P kg<sup>-1</sup> C yr<sup>-1</sup>), root depth  $(d_{root})$ , the concentration of inorganic p at different soil depths  $(P_{in})$ , and a half saturation term at which half of

The root uptake capacity depends on the maximum root uptake capacity  $(v_{max})$  (kg P kg<sup>-1</sup> C yr<sup>-1</sup>), root depth ( $d_{root}$ ), the concentration of inorganic p at different soil depths  $(P_{in})$ , and a half saturation term at which half of the maximum uptake capacity is reached using inorganic p at different soil depths  $(P_{in})$ , a scaling uptake ratio ( $K_p$ ) (µmol P l<sup>-1</sup>), unit conversion ( $C_f$ ) (1 kg P<sup>-1</sup>), and soil moisture ( $\theta$ ) (1 m<sup>-2</sup>), as follows:

$$333 \quad u^{max} = v_{max} \times d_{root} \times \sum_{n=1}^{N} P_{in_n} \times \left(\frac{1}{\sum_{n=1}^{N} P_{in_n} + c_f \times K_p \times \theta_n}\right)$$
(eq.27)

334

335 Description of the litter production of P  $(F_{P_n}^{lit})$  (arrow b in Fig 1) follows JULES-CN as in Wiltshire *et al.*, 336 (2021) and is calculated based on the litter flux of C (kg C m<sup>-2</sup> yr<sup>-1</sup>) using leaf, root and wood turnovers (yr<sup>-1</sup>), 337 and through the vegetation dynamics due to large-scale disturbance and litter production density, as follows: 338

$$\begin{array}{l} 339 \quad F_{P_n}^{lit} = \left(1 - k_{leaf}\right) \gamma_{leaf} C_{leaf} \times C : P_{leaf} + (1 - k_{root}) \gamma_{root} C_{root} \times C : P_{root} + \gamma_{wood} C_{wood} \times C : P_{stem} \\ 340 \\ 341 \end{array}$$
 (eq.28)

342 where  $\lambda$  is the leaf, root and stem re-translocation (at daily timestep) coefficient (Zaehle and Friend, 2010; Clark 343 *et al.*, 2011) and the related *C*: *P* ratios for P fraction and  $\gamma$  is a temperature dependent turnover rate representing 344 the phenological state (Clark *et al.*, 2011).

The decomposition of litter  $(dec^{lit})$  (arrow c in Fig 1) depends on soil respiration (R) (kg C m<sup>-2</sup> yr<sup>-1</sup>), the litter 348 C:P ratio ( $C: P_{lit}$ ) at each soil layer (n) as follows: 349  $dec_P^{lit} = \frac{\sum_{n=1}^{N} R_n}{CP_{lit}}$ 350 (eq.2<u>9</u>) 351 352 where the C: P<sub>lit</sub> is calculated based on litter C pool (DPM and RPM) (lit<sup>C</sup>) (kg C m<sup>-2</sup> yr<sup>-1</sup>) and litter P pool 353  $(P_{O_1})$  as follows: 354  $C: P_{lit} = \frac{\sum_{n=1}^{N} lit_n^C}{P_{O_{ln}}}$ 355 (eq.<u>30</u>) 356 357 The mineralized  $(F_{minl_P})$  (arrow d in Fig 1) and immobilized  $(F_{immob_P})$  (arrow e in Fig 1) P fluxes are 358 calculated based on C mineralization and immobilization, C:P ratios of plant (i) (DPM/RPM) (C: Pplant) and 359 soil (HUM/BIO)(C:  $P_{soil}$ ), soil pool potential respiration ( $R_{POT_i}$ ) (kg C m<sup>-2</sup> yr<sup>-1</sup>) and the respiration partitioning 360 fraction (resp\_frac) as follows: 361  $F_{minl_{P_n}} = \frac{\sum_{n=1}^{N} R_{POT_{i,n}}}{\varepsilon_{cn}}$ 362 (eq.31)363  $F_{immob_{P_n}} = \frac{\sum_{n=1}^{N} R_{i,n} \times resp_{frac}}{C:P_{soil}}$ 364 (eq.32)365 366 The soil respiration from each soil layer  $(R_{i,n})$  is estimated from potential soil respiration  $(R_{POT_{i,n}})$  for the 367 <u>DPM, RPM pools and the litter decomposition rate modifier  $(F_{P_n})$  as follows:</u> 368 369  $R_{i,n} = R_{POT_{i,n}} \times F_{P_n}$ (eq.33) 370 where the description of  $F_{P_n}$  for P pools  $(F_{P_{P_n}})$  follows Wang *et al.*,(2007) and is estimated based on the soil 371 372 <u>pool (BIO/HUM) mineralization (minl<sub>P-BIOn</sub>, minl<sub>P-HUMn</sub>) and immobilization (immob<sub>P-BIOn</sub></u>. 373  $immob_{P-HUM_n}$  (in kg P m<sup>-2</sup> yr<sup>-1</sup>), soil inorganic P ( $P_{inorg_n}$ ) (in kg P m<sup>-2</sup>), and litter pools (DPM/RPM) demand 374 (in kg P m<sup>-2</sup> yr<sup>-1</sup>) as follows: 375  $F_{P_{P_n}} = \frac{(minl_{P-BIO_n} + minl_{P-HUM_n} - immob_{P-BIO_n} - immob_{P-HUM_n}) + P_{inorg_n}}{DEM_{DPM_n} + DEM_{RPM_n}}$ 376 (eq.34)377 The net demand associated with decomposition of litter pools  $(DEM_{k,n})$  represents the P required by microbes 378 379 which convert DPM and RPM into BIO and HUM. The limitation due to insufficient P availability is estimated 380 based on the potential mineralization  $(minl_{p-pot})$  and immobilization  $(immob_{p-pot})$   $(in \text{ kg P m}^{-2} \text{ yr}^{-1})$  of pools 381 (k) as follows: 382 383  $DEM_{k,n} = immob_{p-pot,k} - minl_{p-pot,k}$ (eq.35) 384 385 <u>The  $F_{P_n}$  estimated for N pools ( $F_{P_{N_n}}$ ) follows the same formulation as P (see Wiltshire *et al.*, 2021 for detail)</u> 386 and the  $F_{P_n}$  is estimated based on a higher rate modifier between N and P as follows: 387  $F_{P_n} = \begin{cases} F_{P_{P_n}} F_{P_{P_n}} > F_{P_{N_n}} \\ F_{P_{N_n}} F_{P_{N_n}} > F_{P_{P_n}} \end{cases}$ 388 (eq.36)389 <u>Description of the fluxes of adsorption</u>  $(F_{P_{in_n}}^{sorp})$  (arrow e in Fig 1) and desorption  $(F_{P_{in_n}}^{desorp})$  (arrow f in Fig 390 391 <u>1)</u> of inorganic P in kg P m<sup>-2</sup> yr<sup>1</sup> follow Wang *et al.*, (2010) and are calculated based on soil inorganic ( $P_{in_n}$ ) and sorbed inorganic  $(P_{inorg-sorbed_n})$  P pools and inorganic adsorption  $(K_{sorp-in})$ , desorption  $(K_{desorp-in})$ 392 393 coefficients (kg P m<sup>-2</sup> yr<sup>-1</sup>) and maximum sorbed inorganic ( $P_{in-max}$ ) (kg P m<sup>-2</sup>) as follows: 394  $F_{P_{in_n}}^{sorp} = P_{in_n} \times K_{sorp-in} \times \frac{\left(P_{in-max_n} - P_{inorg-sorbed_n}\right)}{P_{in-max_n}}$ 395 (eq.37)

Variable	Unit	Definition
ψ	kg C m <sup>-2</sup> yr <sup>-1</sup>	Excess C <u>flux</u>
Ø	kg P m <sup>-2</sup> yr <sup>-1</sup>	Plant demand for uptake
П <sub>с</sub>	kg C m <sup>-2</sup> yr <sup>-1</sup>	Potential NPP
u <sup>max</sup>	kg P kg <sup>-1</sup> C yr <sup>-1</sup>	Root uptake capacity
DEM	kg P m <sup>-2</sup> yr <sup>-1</sup>	Plant pool P associated decomposition demand
dec <sub>P</sub> <sup>lit</sup>	kg P m <sup>-2</sup> yr <sup>-1</sup>	Litter decomposition
$F_D$	kg P m <sup>-2</sup> yr <sup>-1</sup>	Plant diffusion flux
$\overline{F_P}$	-	Plant litter decomposition rate modifier
$F_p^{lit}$	kg P m <sup>-2</sup> yr <sup>-1</sup>	Plant litter flux
$F_p^{up}$	kg P m <sup>-2</sup> yr <sup>-1</sup>	Plant uptake
$F_{PO_{S}}^{sorp}$	kg P m <sup>-2</sup> yr <sup>-1</sup>	Sorbed organic P flux
$F_{P_{in}}^{sorp}$	kg P m <sup>-2</sup> yr <sup>-1</sup>	Sorbed inorganic P flux
$F_{Pos}^{desorp}$	<u>kg P m<sup>-2</sup> yr<sup>-1</sup></u>	Desorbed organic P flux
$F_{\rm P}$ . desorp	kg P m <sup>-2</sup> yr <sup>-1</sup>	Desorbed inorganic P flux
$F_p^{occ}$	kg P m <sup>-2</sup> yr <sup>-1</sup>	Occluded inorganic P flux
$F_p^{or-occ}$	kg P m <sup>-2</sup> yr <sup>-1</sup>	Occluded organic P flux
$F_{p}^{W}$	kg P m <sup>-2</sup> yr <sup>-1</sup>	Weathered P flux
Fimmoha	kg P m <sup>-2</sup> yr <sup>-1</sup>	Immobilized P flux
lit <sub>c</sub>	kg C m <sup>-2</sup> yr <sup>-1</sup>	C litter flux
lit <sub>frac</sub>	-	Litter fraction
lit <sub>leaf</sub>	kg C m <sup>-2</sup> yr <sup>-1</sup>	Leaf litter flux
litroot	kg C m <sup>-2</sup> yr <sup>-1</sup>	Root litter flux
litwood	kg C m <sup>-2</sup> yr <sup>-1</sup>	Woody litter flux
F <sub>minlp</sub>	kg P m <sup>-2</sup> yr <sup>-1</sup>	Mineralized P flux
$P_n$	kg P m <sup>-2</sup>	Plant P pool
$P_{O_1}$	kg P m <sup>-2</sup>	Litter organic pool
$P_{O_s}$	kg P m <sup>-2</sup>	Soil organic pool
$P_{in}$	kg P m <sup>-2</sup>	Soil inorganic pool
P <sub>inora-sorn</sub>	kg P m <sup>-2</sup>	Soil inorganic sorbed pool
P <sub>org-sorp</sub>	kg P m <sup>-2</sup>	Soil organic sorbed pool
P <sub>occ</sub>	kg P m <sup>-2</sup>	Soil occluded pool
$P_{pm}$	kg P m <sup>-2</sup>	Parent material pool
R	kg C m <sup>-2</sup> yr <sup>-1</sup>	Total respiration
R <u>pot</u>	kg C m <sup>-2</sup> yr <sup>-1</sup>	Total potential respiration
R <sup>s</sup>	kg C m <sup>-2</sup> yr <sup>-1</sup>	Soil respiration
$R_d$	kg C m <sup>-2</sup> yr <sup>-1</sup>	Leaf dark respiration
T <sub>ref</sub>	Κ	Soil reference temperature
$T_s$	Κ	Soil temperature
Veg <sub>c</sub>	kg C m <sup>-2</sup>	Sum of biomass
Z	m	Soil depth

### Table 1. Model variables

<b>Parameter</b>	<b>Value</b>	<u>Unit</u>	Eq.	<b>Description</b>	<b>Source</b>
		<u>C</u> and	N rela	ated	
<u>α</u>	<u>0.25</u>	<b>_</b>	<u>6</u>	Plant type material ratio	(Clark et al., 2011)
$\overline{a}_{wl}$	<u>1.204</u>	<u>kg C m<sup>-2</sup></u>	<u>50</u>	Allometric coefficient	<u>calibrated</u>
$\sigma_l$	<u>0.0375</u>	kg C m <sup>-2</sup> per unit LAI	<u>48</u>	Specific leaf density	Clark et al., 2011
$b_{wl}$	<u>1.667</u>	<b>_</b>	<u>50</u>	Allometric exponent.	Clark et al., 2011
$f_{dr}$	0.005	<b>-</b>	<u>47</u>	Respiration scale factor	Calibrated
resp_frac	<u>0.25</u>	=	<u>32</u>	Respiration fraction	(Clark et al., 2011)
k <sub>leaf</sub>	<u>0.5</u>	=	<u>28</u>	Leaf N re-translocation coeffi-	(Zaehle and
				<u>cient</u>	Friend, 2010)
k <sub>root</sub>	<u>0.2</u>	<b>=</b>	<u>28</u>	Root N re-translocation coeffi-	(Zaehle and
				cient	Friend, 2010)
$d_{root}$	<u>3.0</u>	-	27	Root fraction in each soil layer	<u>(Clark <i>et al.</i>, 2011)</u>
$v_{int}$	<u>7.21</u>	$\mu mol CO_2 m^{-2} s^{-1}$	<u>45</u>	Intercept in the linear regres-	Calibrated
	10.00		4.5	sion between $V_{\rm cmax}$ and $N_{\rm area}$	(Clark <i>et al.</i> , 2011)
$v_{sl}$	<u>19.22</u>	$\mu$ mol CO <sub>2</sub> gN <sup>-1</sup> s-1	<u>45</u>	Slope in the linear regression	<u>Calibrated</u>
	121 571052	a m 0	15	Observed Losf Mass row Area	(Clark <i>et al.</i> , 2011)
LMA	1 70007506	<u>g m-2</u>	<u>45</u>	Observed Lear Mass per Area	Study site
Leaj N	1./900/390	<u>g g-1</u>	$\frac{43}{46}$	basis	<u>Study site</u>
			<u>40</u>	04515	
		Р	related	l	
$C: P_{soil}$	1299.6	-	32	Soil C:P ratio	(Fleischer et al.,
5011		-			2019)
$v_{max}$	0.0007	kg P kg <sup>-1</sup> C yr <sup>-1</sup>	<u>27</u>	Maximum root uptake capacity	Calibrated (Goll et
					<u>al., 2017)</u>
Р	0.7083062	<u>g kg<sup>-1</sup></u>	<u>46</u>	Foliar P concentrations	Study site
$C_{f}$	<u>3.1×10<sup>-5</sup></u>	<u>1 kg P<sup>-1</sup></u>	<u>27</u>	Conversion factor	<u>(Goll et al., 2017)</u>
$D_z$	<u>0.001</u>	$m^2 s^{-1}$	<u>44</u>	Diffusion coefficient	(Burke et al, 2017)
K <sub>occ</sub>	$1.2 \times 10^{-5}$	<u>yr-1</u>	<u>39,</u>	<u>P occlusion rate</u>	<u>(Yang et al., 2014)</u>
			<u>42</u>		
K <sub>p</sub>	<u>3.0</u>	<u>kg P 1<sup>-1</sup></u>	<u>27</u>	Scaling uptake ratio	<u>Calibrated</u>
K <sub>sorp-in</sub>	<u>0.0054</u>	<u>kg P m<sup>-2</sup> yr<sup>-1</sup></u>	<u>37</u>	Inorganic P adsorption coeffi-	Calibrated (Hou et
				cient	<u>al., 2019)</u>
K <sub>sorp-or</sub>	0.00054	<u>kg P m<sup>-2</sup> yr<sup>-1</sup></u>	<u>40</u>	Organic P adsorption coeffi-	<u>Calibrated</u>
				cient	
K <sub>in-max</sub>	0.0075	$\frac{\text{kg P m}^{-2} \text{ yr}^{-1}}{1}$	<u>37</u>	Maximum sorbed inorganic P	Study site
K <sub>or-max</sub>	0.0042	$\frac{\text{kg P m}^{-2} \text{ yr}^{-1}}{1}$	<u>40</u>	Maximum sorbed organic P	Study site
K <sub>w</sub>	$3 \times 10^{-6}$	<u>kg P m<sup>-2</sup> yr<sup>-1</sup></u>	<u>43</u>	<u>P weathering rate</u>	<u>(Wang <i>et al.</i>, 2010)</u>

457

# 458 **2.3** Study sites

459

460 This study uses data from two nearby sites in Central Amazon in Manaus, Brazil. The main site from here on 461 termed study site (2°35''21.08'' S, 60°06''53.63'' W) (Lugli et al., 2020) is for model development and 462 evaluation. The second site is the Manaus K34 flux site (2°36''32.67'' S, 60°12''33.48'' W) which provides 463 meteorological station data for running the model but also provides data for model evaluation. Our study site is 464 the main lowland tropical forest site maintained by the National Institute for Amazon Research (INPA). 465 Research at this site focuses on pre-experimental, plot, and full-scale long-term projects, combining 466 experimental approaches (Keller et al., 2004; Malhi et al., 2009) with modelling (Lapola and Norby, 2014). 467 Moreover, a recent manipulation experiment at this site provides an opportunity for future model testing under P 468 fertilization. We use detailed novel soil and plant P pool data from the study site (Lugli et al., 2020, 2021) for 469 model parameterisation and calibration and carbon stock data for model validation. The study site has a very 470 similar forest, geomorphology, soil chemistry and species composition to the well-known and studied K34 eddy 471 covariance flux site (Araújo et al., 2002). The average reported annual precipitation is 2431 (mm yr<sup>-1</sup>), with a 472 monthly range of 95 to 304 (mm month<sup>-1</sup>), and averaged temperature is 26°C (Araújo et al., 2002). Moreover, 473 the soil class at this site is Geric Ferrosol with a high clay content and weathering activities (Malhi et al., 2004).

### 475 2.4 Model parameterisation, calibration and evaluation 476

477 We use observations from the four control plots of the study site to parameterise, calibrate and evaluate different 478 processes in JULES (Table 3). The observations were collected at 4 soil depths and processed using the Hedley 479 sequential fractionation (Hedley, Stewart and Chauhan, 1982; Quesada et al., 2010). Observed Leaf Mass per 480 Area (LMA) leaf N and leaf P estimated from fresh leaves were used as input parameters to JULES to estimate 481 photosynthetic capacity and respiration parameters. JULES vn5.5 (JULES CN in this study) estimates  $V_{\rm cmax}$ 482 ( $\mu$ mol m<sup>-2</sup> s<sup>-2</sup>) based on Kattge et al. (2009) using foliar N concentrations in area basis (*nleaf*), as follows: 483

$$484 \quad V_{cmax} = v_{int} + v_{sl} * nleaf$$

(eq.45)

486 where  $v_{int}$  is the estimated intercept and  $v_{sl}$  is the slope of the linear regression derived for the  $V_{cmax}$  estimation. 487 We incorporated an additional P dependency on the estimation of V<sub>cmax</sub> following Walker et al. (2014) as 488 follows: 489

$$490 \quad \ln(V_{cmax}) = 3.946 + 0.921 \ln(N) + 0.121 \ln(P) + 0.282 \ln(N) \ln(P)$$
(eq.46)

491

485

492 Where N and P are foliar concentrations in area basis. 493

494 Implementation of eq. 46 resulted in higher  $V_{\rm cmax}$  than in the original version of JULES. A higher  $V_{\rm cmax}$  predicted 495 higher leaf and plant respiration (eq.47). Constrained by observations of NPP and plant respiration at the study 496 site, we modified one of the most uncertain parameters in the description of plant respiration  $(f_{dr})$  (eq. 47) which 497 is the scale factor  $(f_{dr})$  for leaf dark respiration  $(R_d)$  as follows: 498

$$\begin{array}{c} 499 \\ 500 \end{array} \quad R_d = f_{dr} \, V_{cmax} \tag{eq.47}$$

501 The default value for this scale factor is 0.01 (Clark et al., 2011), and for JULES-CNP simulations at our study 502 site it was modified to 0.005.

503 Observations of aboveground biomass were used to calibrate the non PFT dependent allometric relationships in 504 JULES (Clark et al 2011) (eq  $\frac{48-50}{10}$ ) for leaf, root and stem C. Specifically, the  $a_{wl}$  parameter (eq  $\frac{50}{10}$ ) was 505 modified from 0.65 to 1.204 to match better tropical forest allometry:

505	modified from 0.05 to 1.204 to match better tropical forest anom	cuy.
506	-	-
507	$C_{leaf} = \sigma_l L_b$	(eq.4 <u>8</u> )
508	$C_{root} = C_{leaf}$	_(eq.4 <u>9</u> )
509	$C_{stem} = a_{wl} L_b^{b_{wl}}$	_(eq. <u>50</u> )
510		

511 Where  $\sigma_l$  is specific leaf density (kg C m<sup>-2</sup> per unit LAI),  $L_b$  is balanced (or seasonal maximum) leaf area index 512  $(m^2 m^{-2})$ ,  $a_{wl}$  is allometric coefficient (kg C m<sup>-2</sup>) and  $b_{wl}$  is allometric exponent.

Note that JULES CNP uses C3 and C4 photosynthesis model from Collatz et al., 1991; Collatz, Ribas-Carbo and Berry, 1992, which does not include estimation of J<sub>max</sub>. 513 514

515 JULES-CNP has fixed stoichiometry and C:P ratios of leaf and root (measured), and wood (estimated from

516 fresh coarse wood (Lugli, 2013)) which were taken from the study site and prescribed in JULES to simulate P

517 dynamics in the plant. The following belowground data were used to represent various soil P pools: Resin and

518 bicarbonate inorganic P (inorganic P: $P_{in}$ ), organic bicarbonate P (organic P: $P_{0s}$ ), NaOH organic P (sorbed

519 organic P:  $P_{org-sorp}$ ), NaOH inorganic P (sorbed inorganic P:  $P_{inorg-sorp}$ ), residual P (occluded P:  $P_{occ}$ ) and

520 HCL P (parent material P: Ppm) (Table 3). The measurements were collected between 2017 and 2018 in control

- plots. All measurements were conducted at four soil layers (0-5, 5-10, 10-20, 20-30 cm). However, to be
- 521 522 523 524 consistent with the JULES model soil layer discretization scheme, we defined 4 soil layers (0-10 cm, 10-30 cm, 30-100 cm and 100-300 cm) and we used the average between 0 and 30 cm to compare against the measurement from the same depth for model evaluation.
- 525 Vegetation C stocks were derived based on tree diameter measurements at breast height, that are linked to
- 526 allometric equations and wood density databases to estimate the C stored in each individual tree, and then scaled 527 528 to the plot (Chave et al., 2014).

#### 529 The organic and inorganic soil P assumed to be always at equilibrium with the relative sorbed pools (Wang, 530 Law and Pak, 2010). Thus, in order to cap P sorption and uptake capacity, the maximum sorption capacities

- 531 (Pin-maxn, Por-maxn, eq.37 and 39) (adopted from (Wang, Houlton and Field, 2007)) were prescribed using
- 532 maximum observed sorbed inorganic and organic P. Hence, the maximum sorption capacity defines the

- 533 equilibrium state of sorbed and free-soil P. Moreover, as the magnitude of changes in the occluded and parent
- 534 material pools are insignificant over a short-term (20 years) simulation period (Vitousek et al., 1997), these two
- 535 pools were prescribed using observations. Remaining parameters used to describe soil P fluxes (eq.s <u>27-44</u>)
   536 were prescribed using values from the literature (Table 3).

We used a combination of data from *Study site* and the nearby site K34 for model evaluation of C fluxes (GPP,
 NPP) and C pools (soil and vegetation C, leaf, root and stem C) with no calibration on plant and soil organic and
 soil inorganic P polls included (Table 3).

541

Table 3. Observations from study site (taken during 2017-2018) and from Manaus site K34 used for model parameterisation and evaluation

Process	Variables	Purpose of use	Reference and site	
C associated	GPP	Evaluation	Fleischer et al., 2019, K34	
	NPP	Evaluation	Fleischer et al., 2019, K34	
	Soil C	Evaluation	Malhi et al., 2009, K34	
	CUE	<b>Evaluation</b>	Malhi et al., 2009, K34	
	Veg C	Evaluation	Study site	
	Leaf C	<b>Evaluation</b>	Study site	
	Stem C	<b>Evaluation</b>	Study site	
	Root C	<b>Evaluation</b>	Study site	
	LAI	Initialisation	Study site	
	LMA	Parameterisation	Study site	
Р	Resin	Evaluation	Study site	
associated	Pi Bic	Evaluation	Study site	
	Po Bic	Evaluation	Study site	
	Po NaOH	<b>Calibration</b>	Study site	
	Pi NaOH	<b>Calibration</b>	Study site	
	P residual	Parameterisation	Study site	
	P HCL	<b>Parameterisation</b>	Study site	
	Leaf N	<b>Parameterisation</b>	Study site	
	Leaf P	<b>Parameterisation</b>	Study site	
	Root P	Parameterisation	Study site	
	Plant C:P ratio	Parameterisation	Study site	

# 544

# 545 2.5 JULES simulations

546

547 JULES was applied at the K34 flux tower site using observed meteorological forcing data from 1999-2019 548 (Fleisher et a 2019) at half hourly resolution. The following meteorological variables are needed to drive JULES 549 (model inputs) (Best et al., 2011): atmospheric specific humidity (kg kg<sup>-1</sup>), atmospheric temperature (K), air 550 pressure at the surface (Pa), short and longwave radiation at the surface (W  $m^{-2}$ ), wind speed (m  $s^{-1}$ ) and total 551 precipitation (kg m<sup>-2</sup> s<sup>-1</sup>). Furthermore, the averaged measured LAI from study site was used to initialise the 552 vegetation phenology module, but was allowed to vary in subsequent prognostic calculations. Soil organic and 553 inorganic sorbed P pools were initialised with study site observations. The JULES CNP simulations were 554 initialized following the same methodology as in Fleischer et al., (2019), by the spin-up from 1850 recycling 555 climatology to reach equilibrium state (Figure S1) and spin up was performed separately for three versions of 556 JULES (C/CN/CNP) following the same procedure. Furthermore, the transient run was performed for the period 557 1851-1998 using time-varying CO<sub>2</sub> and N deposition fields. Finally, for the extended simulation period (1999-558 2019) two runs were performed, the first with ambient the second elevated  $CO_2$  concentrations. 559 560 We evaluate the impact of including a P cycle in JULES using three model configurations (JULES C, CN and 561 CNP). We apply JULES in all three configurations using present day climate under both ambient CO<sub>2</sub> and 562 elevated CO<sub>2</sub> (eCO<sub>2</sub>). Ambient and eCO<sub>2</sub> were prescribed following Fleischer et al., (2019), with present-day 563  $CO_2$  based on global monitoring stations, and an abrupt (step) increase in atmospheric  $CO_2$  of +200 ppm on the 564 onset of the transient period (i.e., 1999). However, the comparison period is limited to 2017-18 for which the P 565 measurements are available. We compare simulated C fluxes (GPP, NPP, litterfall C), C stocks (total vegetation, fine root, leaf, wood, soil) 566 567 and the CO<sub>2</sub> fertilization effect across model configurations. The CO<sub>2</sub> fertilization effect  $(CO2_{fert-eff})$  (eq.51) 568 is calculated based on simulated vegetation C under ambient (VegC  $(aCO_2)$ ) and eCO<sub>2</sub> (VegC  $(eCO_2)$ ) as

570  
571  
572
$$CO2_{fart-eff} = \frac{(vegc (ec0_2)-vegc (ac0_2))\times 100}{vegc (ac0_2)}$$
(eq.51)573  
574  
575Furthermore, the net biomass increases due to CO; fertilization effect ( $\Delta C veg$ ) is estimated as follows:574  
575  
576  
577 $\Delta C_{veg} = \Delta BP - \Delta litter fail C$ (eq.52)576  
577  
577  
578We studied the Water Use Efficiency (WUE) (eq. 53) at half-hourly timestep, then aggregated per month as one  
of the main indicators of OPP changes (Xiao *et al.*, 2013), and soil moisture content (SMCL), as one of the  
main controllers of maximum uptake capacity (eq. 27), in order to better understanding the changes in GPP, P  
demand and uptake as well as excess C fluxes.588  
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Figure. 2- Modelled vs measured soil phosphorus pools under ambient  $CO_2$  (for the soil depth of 0-30cm). Black line represents standard deviation

Table 4. Observed and simulated phosphorus pools and fluxes. Occluded and weathered P pools were prescribed using the observed values (between period 2017-18).

	Pl	Phosphorus pools and fluxes				
	Measured	Modelled	Modelled			
		Ambient CO <sub>2</sub>	Elevated CO <sub>2</sub>			
<b>Organic P</b> (g P m <sup>-2</sup> )	1.09±0.53	1.6	1.57			
<b>Inorganic P</b> $(g P m^{-2})$	1.05±0.33	1.07	0.96			
<b>Sorbed organic P</b> (g P m <sup>-2</sup> )	$1.04{\pm}0.42$	1.04	1.03			
Sorbed inorganic $P(g P m^{-2})$	2.1±0.55	2.4	2.4			
<b>Occluded P</b> ( $g P m^{-2}$ )	7.98±2.38	prescribed	prescribed			
Weathered $P (g P m^{-2})$	0.59±12	prescribed	prescribed			
Total vegetation P (g P m <sup>-2</sup> )	4.15	4.66	5.11			
Soil $P - 30 \text{ cm} (\text{g P m}^{-2})$	13.85	14.7	14.56			
Total ecosystem P (g P m <sup>-2</sup> )	-	35.97	35.97			
<b>P litter flux</b> (g P m $^{-2}$ yr $^{-1}$ )	0.3	0.28	0.29			

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### 3.1.2 Model evaluation

JULES CNP-CNP could reproduce the plant and soil C (Figure.2 and Table 5) and N pools and fluxes (Figure
 S6 and Table 6) pools and fluxes under ambient CO<sub>2</sub>. Our results show that simulated GPP, is within the range
 of measurement (3.02 kg C m<sup>-2</sup> yr<sup>-1</sup> model vs 3-3.5 kg C m<sup>-2</sup> yr<sup>-1</sup> observed, respectively, Table 5).

636 637 Simulated NPP, is close to the measured values (NPP: 1.14 - 1.31 observed vs 1.26 modelled kg C m<sup>-2</sup> yr<sup>-1</sup>) with 638 autotropic respiration (RESP) also closely following the observations (1.98 observed vs 1.81 modelled kg C m<sup>-2</sup> 639 yr<sup>-1</sup>). Biomass production is estimated as a difference between NPP and the amount of C which is not fixed by 640 plants due to the insufficient P in the system (excess C) (eq. 27). The excess C flux is highly dependent on the 641 plant P and the overall P availability to satisfy demand (Table 5). Simulated flux of excess C is 0.3 kg C m<sup>-2</sup> yr<sup>-1</sup>

642 under ambient CO<sub>2</sub>. In JULES-CNP this flux is subtracted from NPP in order to give the BP (eq. 17) (Table 5).

643 Our simulated litterfall overestimates the observations by 32%, however simulated vegetation and its

644 components (fine root, leaf and wood) and soil C stocks match well the observations (Table 5).

<b>1 able 5.</b> Observed and simulated carbon pools and nuxes with JULES CMP (between period 2017-18)
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Carbon pools and fluxes								
	Measured	Modelled Ambient CO <sub>2</sub>	Modelled Elevated CO <sub>2</sub>					
<b>GPP</b> (kg C m <sup>-2</sup> yr <sup>-1</sup> )	3.0-3.5	3.06	3.9					
<b>NPP</b> <sub>pot</sub> (kg C $m^{-2}$ yr <sup>-1</sup> )		1.27	1.77					
<b>Plant respiration</b> (kg C m <sup>-2</sup> yr <sup>-1</sup> )	1.98	1.78	2.12					
Excess C flux (kg C m <sup>-2</sup> yr <sup>-1</sup> )	-	0.30	0.81					
<b>Biomass Production</b> (kg C m <sup>-2</sup> yr <sup>-1</sup> )	<u>1.14±0.12</u>	0.96	0.94					
Litter C flux (kg C m <sup>-2</sup> yr <sup>-1</sup> )	0.69 <u>±0.15</u>	0.91	0.83					
Leaf C (kg C m <sup>-2</sup> )	0.37 <u>±0.2</u>	0.38	0.40					
Wood C (kg C $m^{-2}$ )	22.01	22.4	24.71					
<b>Root</b> C (kg C $m^{-2}$ )	0.37 <u>±0.2</u>	0.38	0.40					
Vegetation C (kg C m <sup>-2</sup> )	22.75 <u>±0.3</u>	23.16	25.52					
Soil C stock (kg C m <sup>-2</sup> )	12.7	13.2	12.71					
LAI $(m^2 m^{-2})$	5.6 <u>±0.36</u>	5.77	6.12					

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## 3.1.3 Comparison of JULES C, CN and CNP under ambient CO<sub>2</sub>

649 We compare simulated C pools and fluxes from JULES-C, JULES-CN and JULES-CNP (Figure. 3). There is no 650 difference between C stocks and fluxes in simulations from JULES C and CN indicating that there is no N 651 limitation at this tropical site in the CN simulations. However, simulated BP and litter flux of C by JULES 652 C/CN are higher than in JULES-CNP but also overestimate the observations (litter flux of JULES C/CN: 1.18, 653 JULES CNP: 0.91 and obs 0.69 (kg C m<sup>-2</sup> yr<sup>1</sup>) and BP of JULES C/CN: 1.24, JULES CNP: 0.96 and obs1.14-654 1.31 (kg C m<sup>-2</sup> yr<sup>-1</sup>), respectively). By including the P cycling in JULES an excess C flux of 0.3 (kg C m<sup>-2</sup> yr<sup>-1</sup>) 655 is simulated, indicating a 24% P limitation to BP at this site according to JULES CNP, which represents a 29% 656 decrease in BP compared to JULES-C/CN. Consequently, the total vegetation C stock for models without P 657 inclusion is higher than the CNP version (+3% difference) due to the lack of representation of P limitation. The 658 simulated soil C stock in JULES C and JULES CN is also higher than in the CNP version (JULES C/CN: 13.93 659 vs. JULES CNP: 13.18 (kg C m<sup>-2</sup> yr<sup>-1</sup>)) and higher than the observations. Moreover, CUE in JULES C/CN 660 (eq.54) is higher than observations and JULES CNP version (JULES C/CN: 0.38 vs. JULES CNP: 0.31, obs: 661  $0.34 \pm 0.1$  (dimensionless).





fluxes (BP and Litter C) (in kg C m<sup>-2</sup> yr<sup>-1</sup>) and CUE under ambient CO<sub>2</sub>. Note that C

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### 670 **3.1.4 <u>Model sensitivity</u>** 671

672 The results indicate that among all the corresponding C and P pools and fluxes, the excess C flux – which 673 demonstrates P limitation to growth – shows the highest sensitivity to changes in C:P ratios, KP and 674 Kor-max, Kin-max. A decrease in plant C:P results in a large increase in excess C. This is due to the higher plant 675 P demand as a result of lower plant C:P ratios. An increase in the uptake factor and maximum sorbed organic 676 and inorganic P also results in an increase in excess C. This is due to the higher uptake demand through higher 677 uptake capacity (due to higher KP) and lower available P for uptake due to higher organic and inorganic sorbed P (due to higher K<sub>or-max</sub>, K<sub>in-max</sub>). Since the total P in the system is lower than the plant demand, the uptake 678 679 capacity and sorbed P, higher P limitation is placed on growth (decreasing BP) which results in an increase in 680 excess C and decrease in plant C, but also soil C which is a result of lower litter input (Figure 4). Total soil P 681 shows low sensitivity to changes in plant C:P and uptake factor but high sensitivity to maximum inorganic 682 sorbed P. Moreover, sorbed P shows middle to high sensitivity to maximum organic and inorganic sorbed P 683 respectively (Figure. S5). Nevertheless, organic and inorganic P adsorption coefficients (K<sub>sorp-or</sub>, K<sub>sorp-in</sub>) 684 show no sensitivity to C and P pools and fluxes. This is due to limiting the organic and inorganic P sorption 685 terms controlled only by maximum sorption, hence no effect applied by organic and inorganic adsorption 686 coefficients.

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Figure. 4- Model sensitivity test results and corresponding C and P pools and fluxes under ambient CO2.

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# 3.2 Model application under elevated CO<sub>2</sub>

# 694 **3.2.1** Simulated plant and soil C and P pools and fluxes -JULES CNP: eCO<sub>2</sub> vs ambient CO<sub>2</sub> 695

696 The eCO<sub>2</sub> simulation using JULES CNP yields a higher GPP compared to the ambient CO<sub>2</sub> (0.83 (kg C m<sup>-2</sup> yr<sup>-1</sup>) increase), as a result of CO2 fertilization. Moreover, due to the GPP increase, NPP and RESP follows the same 697 698 trend and increased compared to ambient CO<sub>2</sub> (NPP: 0.49 and RESP:0.3 (kg C m<sup>-2</sup> yr<sup>-1</sup>) increase) (Table 5). The 699 total simulated vegetation C pool increases under eCO<sub>2</sub> compared to ambient CO<sub>2</sub> (0.41 kg C m<sup>-2</sup>), hence the 700 estimated plant P (estimated as a fraction of C:P ratios) increases as well (+0.45 (g P m<sup>-2</sup>)) (Fig 6, Table 4). 701 Thus, the simulated plant P demand is higher, and as the total available soil P for uptake is limited, the simulated 702 excess C flux increases to  $0.51(\text{kg C m}^{-2} \text{ yr}^{-1})$ . Moreover, despite the higher NPP under eCO<sub>2</sub> compared to 703 simulated NPP under ambient CO<sub>2</sub>, due to the substantial increase in simulated excess C, the BP is similar to the 704 ambient CO<sub>2</sub> (2% difference). 705

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- 709 The simulated organic soil P under eCO<sub>2</sub> yields close to the ambient CO<sub>2</sub> (1.6 g P  $m^{-2}$ ) (Table 5). This is due to
- 710 the same parameterization of the output fluxes from this pool for eCO<sub>2</sub> and ambient CO<sub>2</sub>. The simulated pool of
- 711 inorganic P under eCO<sub>2</sub> decreases compared to the ambient CO<sub>2</sub> by 0.11 (g P m-<sup>2</sup>) due to the increased plant P
- 712 pools and slight increase in uptake (+0.13 %).
- 713 However, the simulated sorbed organic and inorganic soil P from eCO<sub>2</sub> are similar to those simulated under the
- 714 ambient CO<sub>2</sub> which is due to the same parameterizing of sorption function (maximum sorption capacity) from
- 715 the ambient CO<sub>2</sub> run as explained in calibration section. Moreover, the modelled occluded and weathered soil P 716 yield similar to those in the ambient CO<sub>2</sub> simulation (Table 5) which is due to the same prescribed observational
- 717 data that was used for this simulation.
- 718

#### 719 3.2.2 Comparison of JULES C, CN and CNP under elevated CO<sub>2</sub> 720

721 JULES C/CN show higher vegetation and soil C pools, BP and litter flux compared to JULES-CNP: (Table 6, 722 Figure. S2). Under eCO<sub>2</sub>, simulated NPP using JULES C-CN is 4.5% higher than JULES CNP and the BP with 723 JULES- C/CN is 96.8% higher than in JULES-CNP which simulates an excess C flux of 0.81 (kg C m<sup>-2</sup> yr<sup>-1</sup>) 724 equivalent to 46% P limitation under eCO<sub>2</sub>. As a result of P limitation and eCO<sub>2</sub>, the simulated CO<sub>2</sub> fertilization 725 effect estimated based on changes in biomass under ambient and eCO2 was reduced from 13% with JULES-726 C/CN to 10% JULES-CNP. Moreover, the CUE from JULES C/CN is 87.5% higher than the JULES CNP as a 727 728 729 730 result of high P limitation over biomass production.

Table 6. C pools and fluxes using JULES C/CN and difference in percentage with JULES CNP model under eCO2. A positive % means larger respective values simulated with JULES C and JULES CN than with JULES CNP (between period

731 2017-18).

	GPP	NPP	BP	CUE	Litter C	Leaf C	Root C	Wood C	Soil C
JULES C/CN	4.1	1.85	1.85	45%	1.77	0.42	0.42	26.1	19.2
JULES CNP	3.9	1.77	0.94	24%	0.83	0.4	0.4	24.71	12.71
$\Delta C/CN$ : CNP	5.1%	4.5%	96.8%	87.5%	113.3%	5%	5%	5%	51.1%

### 732 733 3.2.2.1 Inter-models under elevated CO<sub>2</sub>

734 735 Following Fleischer et al., (2019), we report the simulated response to eCO<sub>2</sub> for year 1999 (initial: CO<sub>2</sub> effect) 736 and 1999-2013 (15 years: final effect) which are different than our evaluation period (2017-18). Using JULES C 737 and JULES CN under eCO<sub>2</sub>, simulated GPP and NPP during the 1<sup>st</sup> year increase by 30% and 61% respectively 738 and by 28% and 52% after 15 years (Figure. 5). However, using JULES CNP, eCO<sub>2</sub> increases simulated GPP, 739 NPP and BP responses during the 1st year by 29%,51% and 20% and by 28%, 43% and 7%, after 15 years 740 respectively.

741

742 Corresponding simulated CUE during the 1<sup>st</sup> year and 15 years shows an increase of 24% and 20% in response 743 to eCO<sub>2</sub> using JULES C/CN respectively. However, using JULES CNP, simulated CUE for the 1<sup>st</sup> and after 15 744 years is reduced by 7% and 17% in response to eCO<sub>2</sub>. 745

746 Simulated total biomass (leaf, fine root and wood C) ( $\Delta Cveg$ ) using JULES C/CN for the 1<sup>st</sup> and 15 years of

747 eCO<sub>2</sub> increases by 9% and 13% respectively. However, using JULES CNP ΔCveg only increases by 0.5% and

- 748 9% for 1<sup>st</sup> and 15 years of eCO<sub>2</sub>, respectively.
- 749
- 750



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**Figure. 5-** Relative effect of eCO<sub>2</sub> on simulated GPP, NPP, BP, CUE,  $\Delta Cveg$ , leaf C, wood C and fine root C, using three versions of JULES model in 1<sup>st</sup> (initial response) and 15 years periods (final response).

### 6 **3.3** Plant P Demand, uptake and <u>excess C</u> under ambient and elevated CO<sub>2</sub>

To understand further the CP-cycle dynamics, we studied the monthly averaged plant P demand and the relative
 (limited) P uptake (eq. <u>26</u>) under both ambient and elevated CO<sub>2</sub> conditions (Figure. 6).

761Under ambient CO2 condition the highest GPP is estimated at  $3.5\pm0.19 \text{ kg Cm}^{-2} \text{ month}^{-1}$  in July and the lowest762at  $2.06\pm0.61 \text{ kg Cm}^{-2} \text{ month}^{-1}$  in October (Figure. 6-a). The estimated WUE and SMCL in October is among the763lowest estimated monthly values at  $2.3\pm0.51 \text{ kg CO}_2/\text{kg H}_2\text{O}$  and  $526.2\pm31 \text{ kg m}^{-2}$  respectively (Figure. 6-c).764The highest P demand is estimated at  $0.4\pm0.02 \text{ g P m}^{-2} \text{ month}^{-1}$  in July and the lowest demand at  $0.2\pm0.08 \text{ g P m}^{-2}$ 765 $^2 \text{ month}^{-1}$  in October. Consequently, the highest and lowest uptake ( $0.32\pm0.01$  and  $0.19\pm0.07 \text{ g P m}^{-2} \text{ month}^{-1}$ ,766respectively). The excess C for the highest and lowest GPP and demand periods are estimated at  $0.4\pm15$  and767 $0.04\pm0.07 \text{ kg C m}^{-2} \text{ month}^{-1}$ , respectively.

769 However, similar to ambient CO<sub>2</sub>, under eCO<sub>2</sub> condition the highest estimated GPP is in July at 4.36±0.21 kg C 770 m<sup>-2</sup> month<sup>-1</sup> and lowest for October 3.02±0.75 kg C m<sup>-2</sup> month<sup>-1</sup> (Figure. 6-b). The estimated WUE and soil 771 moisture content (SMCL) for the lowest GPP period is among the lowest monthly estimated values at 3.5±0.74 772 kg CO<sub>2</sub>/kg H<sub>2</sub>O and 552±33 kg m<sup>-2</sup> for October respectively (Figure. 6-d). The highest P demand is estimated 773 for July at 0.51±0.02 g P m<sup>-2</sup> month<sup>-1</sup> with the uptake flux of 0.31±0.02 g P m<sup>-2</sup> month<sup>-1</sup> and the lowest demand 774 is estimated for October at 0.32±0.1 g P m<sup>-2</sup> month<sup>-1</sup> with the estimated uptake flux of 0.26±0.06 g P m<sup>-2</sup> month<sup>-1</sup> 775 <sup>1</sup>. The highest excess C flux is also for July at 1.01±0.17 kg C m<sup>-2</sup> month<sup>-1</sup> and lowest for October 0.27±0.29 kg 776 C m<sup>-2</sup> month<sup>-1</sup>, respectively. 777

However, despite the P limitation in both eCO<sub>2</sub> and ambient CO<sub>2</sub> conditions, the P uptake flux under eCO<sub>2</sub> is
higher than the ambient CO<sub>2</sub> condition. This is due to the higher WUE and increased SMCL (controlling uptake
capacity (eq. 27)) under eCO<sub>2</sub> condition, hence more water availability during the dry season to maintain

- 781 productivity and critically transport P to the plant (see eq. 27), compared to ambient CO<sub>2</sub> condition (Figure. 6-c
- 782 and d). Additionally, in JULES both the vertical discretisation (Burke, Chadburn and Ekici, 2017) and

783 mineralisation terms (Wiltshire et al., 2021) depend on the soil moisture and temperature. Thus, higher P

784 concentration and uptake under eCO<sub>2</sub> condition.

785



 $\begin{array}{c} \textbf{Figure. 6- Simulated monthly plant P demand and uptake (g P m^{-2} yr^{-1}), excess C and GPP (kg C m^{-2} yr^{-1}) under a) aCO_2} \\ \textbf{and b) eCO_2, water use efficiency (g m^{-2} yr^{-1}) under c) ambient CO_2 (aCO_2) and d) eCO_2 conditions. The grey area represents the standard deviation. \\ \textbf{790} \end{array}$ 

# 792 **3.4** Soil P pools profile under ambient CO<sub>2</sub> and elevated CO<sub>2</sub> 793

We explored the distribution of the inorganic and organic soil P and their sorbed fraction within the soil layers and under different CO<sub>2</sub> conditions (Figure. S3). Both the ambient and eCO<sub>2</sub> simulations have a close inorganic soil P distribution at the topsoil layer (0-30cm) (0.85 vs. 0.9 (g P m<sup>-2</sup>) respectively) as well as similar organic soil P distribution (0.85 vs 0.9 (g P m<sup>-2</sup>) respectively).

However, the organic soil P and sorbed forms of inorganic and organic soil P profiles are not changing
significantly between different sets due to the similar parameterization of the processes that control these pools
(processes which are related to the physical aspects of soils, hence not changing under eCO<sub>2</sub> condition) and the
same parameter values used for both ambient and eCO<sub>2</sub> runs.

804 Moreover, the soil P within 30cm soil depth for ambient and eCO<sub>2</sub> conditions is at 14.7 (g P m<sup>-2</sup>) and 14.56 (g P m<sup>-2</sup>) respectively, and the total ecosystem P for both ambient and eCO<sub>2</sub> conditions is at 35.97 (g P m<sup>-2</sup>).

- However, the slightly lower soil P in the eCO<sub>2</sub> condition is due to the higher plant P demand compared to the
- ambient condition, hence the higher allocated P vegetation (10%) under eCO<sub>2</sub> condition.

## 808 809 **4. Discussion**

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- 811 Studies show the significant role of the tropical forests, and Amazonia in particular, in C uptake and regulating
- atmospheric CO<sub>2</sub> (Brienen *et al.*, 2015; Phillips *et al.*, 2017). As soil P availability is low in the majority of
   Amazonia (Ouesada *et al.*, 2012), the competition for nutrients by both plant and soil communities is high
- Amazonia (Quesada *et al.*, 2012), the competition for nutrients by both plant and soil communities is high
   (Llovd *et al.*, 2001). The responses of these communities to eCO<sub>2</sub> under P limited conditions remains uncertain
- $\begin{array}{l} 814 \\ \hline (Lloyd \ et \ al., 2001). \ The responses of these communities to eCO_2 under P limited conditions remains uncertain \\ \hline (Fleischer \ et \ al., 2019). \ These responses in P enabled models are represented in different ways regarding the \\ \end{array}$
- 815 (Fleischer *et al.*, 2019). These responses in P enabled models are represented in different ways regarding the 816 excess C which is not used for plant growth due to P limitation. Either growth is directly downregulated taking
- 817 the minimum labile plant C,N and P (Goll *et al.*, 2017), or photosynthesis is downregulated via  $V_{\text{cmax}}$  and  $J_{\text{max}}$

818 (Comins and McMurtrie, 1993; Yang et al., 2014; Zhu et al., 2016) and finally models like JULES CNP

- 819 downregulate NPP via respiration of excess carbon that cannot be used for growth due to plant nutrient
- 820 constraints (Haverd et al., 2018). The estimated CUE depends on the modelling approach. Models that down
- 821 regulate the photosynthetic capacity and GPP consequently (Comins and McMurtrie, 1993; Yang et al., 2014; Zhu et al., 2016), simulate a positive CUE response to CO<sub>2</sub> fertilization while models that down regulate the
- 822 823 NPP and respire the excess C (Haverd et al., 2018) simulate a negative CUE response (Fleischer et al., 2019)
- which is in line with the studies showing lower CUE when nutrient availability declines (Vicca et al., 2012). 824
- 825 However, this remains a major uncertainty in understanding the implication of P limitation on terrestrial 826 biogeochemical cycles.
- 827 Our new developments include major P processes in both plant and soil pools and can be applied to the Amazon
- 828 region using existing soil (Quesada et al., 2011) and foliar structural and nutrient (Fyllas et al., 2009) data for
- 829 parameterisation. Moreover, JULES CNP can be applied at the global scale and for future projections using
- 830 global soil P data (Sun et al., 2021) for model initialization and PFT-specific plant stoichiometries
- 831 (Zechmeister-Boltenstern et al. 2015), soil stoichiometries (Zechmeister-Boltenstern et al. 2015; Tipping et al.
- 832 (2016), sorption and weathering ratios (based on lithological class specific from the GliM lithological map
- 833 (Hartmann and Moosdorf, 2012) and soil shielding from Hartmann et al., (2014)). 834
- 835 4.1. Evaluation of model performance against observations
- 836

837 JULES-CNP could reproduce the magnitude of soil organic and inorganic P pools and fluxes. The relative 838 distribution of total organic P, total inorganic P and residue P fractions of total P in soils under Brazilian 839 Eucalyptus plantations (Costa et al., 2016) shows inorganic P fraction of 28% from total soil P which is close to 840 our estimation of 24% and organic P fraction of 30% from total soil P which is higher than our estimated 841 fraction of 18%. Thus, we may need to improve the process representation or parameters that control the organic 842 P concentration, such as litter flux and decomposition, soil organic P mineralization, and immobilization in the 843 future.

844 845 Our estimated maximum P uptake, which represents the actual available P for plant uptake (Goll et al., 2017), 846 for both ambient and eCO<sub>2</sub> conditions, is highly correlated with the plant P demand ( $R^2 = 0.96$  and 0.52 847 respectively). The plant P demand depends on the GPP changes which are reflected by the WUE (Hatfield and 848 Dold, 2019). Hence, under ambient CO<sub>2</sub>, JULES CNP simulates lower GPP and plant P demand during the dry 849 season than during the wet season. Sufficient P uptake during these periods results in the lowest P limitation, 850 thus the lowest simulated excess C. Nevertheless, under  $eCO_2$  the same pattern is simulated but a higher 851 availability of soil P due to the stomatal closure in the dry season. Hence, due to the plant's more efficient water 852 usage, the soil moisture in the dry season is higher (Xu et al., 2016) which impacts our capped P uptake flux (eq. 853 854 27) and increases the uptake capacity respectively.

855 Overall, JULES-CNP reproduced the observed C pools and fluxes which are in the acceptable ranges compared 856 to the measurements. However, using the JULES default  $V_{\text{cmax}}$  estimation method (eq. <u>40</u>), the model slightly underestimates the total GPP (2.9 kg C m<sup>-2</sup> yr<sup>-1</sup> vs. 3-3.5 kg C m<sup>-2</sup> yr<sup>-1</sup>). Therefore, in this version of the model, we used the improved  $V_{\text{cmax}}$  estimation method based on N and P (eq. <u>46</u>) which resulted a final estimated GPP 857 858 859 closer to the measurements  $(3.06 \text{ kg C m}^{-2} \text{ yr}^{-1})$ . 860

861 Our results show an increase in GPP (21%) in response to  $eCO_2$  which is higher than the average increase of 862 GPP reported in mature eucalyptus forests (11%), also growing under low P soils at the free air CO<sub>2</sub> enrichment 863 experiment (EucFACE) facility in Australia (Jiang et al., 2020). This can be related to the lower decrease of 864 biomass growth response estimated by JULES-CNP (-3%) compared to the measurements from mature forests 865 (-8%) (Ellsworth et al., 2017), due to the P limitation which showed to impact the above-ground biomass

- 866 growth response in mature forests (Körner et al., 2005; Ryan, 2013; Klein et al., 2016).
- 867

868 In order to estimate the biomass production (BP), we deducted the excess C fluxes from the NPP. Using JULES 869 C/CN models our estimated biomass productivity enhancement due to eCO<sub>2</sub> (49%) is in the middle range of the 870 reported various studies from different biomes by Walker et al., (2021). Moreover, our estimated difference of

871 BP between ambient and eCO<sub>2</sub> conditions (2%) is close to the estimated difference for mature forests (3%) 872 (Jiang et al., 2020).

873 A global estimation for tropical forests using CASACNP model which includes N and P limitations on

874 terrestrial C cycling, shows that NPP is reduced by 20% on average due to the insufficient P availability (Wang, 875 Law and Pak, 2010) which is close to our estimated P limitation of 24%. This finding is in line with

- 876 experimental study that shows a strong correlation between the total NPP and the soil available P (Aragão et al.,
- 877

- 878 similar to sites in pool soils (see ZAR-01 site in Aragão et al., (2009)). The estimated decrease of NPP in
- 879 response to eCO<sub>2</sub> as a result of P limitation is in line with the findings from CLM-CNP model at five tropical
- 880 forests (Yang et al., 2014) which indicates the CO<sub>2</sub> fertilization dependency on the processes that affect P
- 881 availability or uptake.
- 882

883 Our estimated CUE (0.31) is close to the estimation by Jiang et al. (2020) for mature forests ( $0.31\pm0.03$ ), as well 884 as to the measurement for our study site  $(0.34 \pm 0.1)$ . There is currently a lack of representation of stand age in 885 JULES-CNP which can significantly change this ratio (e.g. mature trees are less responsive to the nutrient 886 limitations) (De Lucia et al., 2007; Norby et al., 2016). However, a recent development of Robust Ecosystem 887 Demography (RED) model into JULES (Argles et al., 2020) and its integration into JULES-CNP in the future

- 888 can resolve this issue. Moreover under low P availability, all available P is considered to be adsorbed or taken 889 by plant and microbes for further consumption, with leaching considered to be minor within the time scales of
- 890 our study period (Went and Stark, 1968; Bruijnzeel, 1991; Neff, Hobbie and Vitousek, 2000).
- 891 Due to the strong fixation of P in the soil (Aerts & Chapin, 2000; Goodale, Laitha, Nadelhoffer, Bover, &
- 892 Jaworski, 2002), the P deposited is unlikely to be available to plants in the short term (de Vries et al., 2014), for
- 893 this reason this version of JULES CNP did not include P deposition. However both P deposition and leaching
- 894 are likely to have a very important role on sustaining the productivity of tropical forests in the Amazon over
- 895 longer time scales (Van Langenhove et al., 2020) and needs to be considered in future studies.
- 896 Moreover, biochemical mineralisation is not included in the current version of JULES CNP and it only accounts 897 for total mineralization. However, even the models which includes this process, show no significant difference
- 898 between total and biochemical mineralized P which can be due to complexity of identifying the inclination of 899 mineralization versus uptake (Martins et al., 2021).
- 900 Lastly, in order to capture plant internal nutrient impact on the C storage, the future work should focus on
- 901 implanting a recent developed Non-Structural Carbohydrate (NSC) model (SUGAR) (Jones et al., 2020) in 902 JULES-CNP.
- 903

#### 904 4.2. Inter-models comparison

905

906 The comparison of simulated GPP enhancement across JULES versions for the 1<sup>st</sup> year is within the middle 907 range of the 1<sup>st</sup> year CO<sub>2</sub> responses of the C/CN models studied by Fleischer et al., (2019) evaluating simulated 908 eCO<sub>2</sub> effects at a site in Manaus using the same meteorological forcing and methodology used in this study for 909 a range of DGVM's. However, comparison for 15 years of eCO<sub>2</sub>, shows that the simulated response with 910 JULES CNP is on the higher end of Fleischer et al., (2019) study which is due to the higher estimated biomass 911 growth by JULES CNP (Table S1). Similarly, using JULES CNP our estimated GPP enhancement is on the 912 higher end of model estimations in Fleischer et al., (2019). Moreover, comparing the GPP responses between 913 different versions of (JULES C/CN and CNP), the JULES CNP shows a slightly higher response to CO<sub>2</sub> 914 fertilization associated with the higher WUE changes (Xiao et al., 2013) (Figure. S4). This is due to the higher 915 sensitivity of the plant to water availability than the P availability in the P limited system (He and Dijkstra, 916 2014). Hence, under eCO<sub>2</sub> due to water-saving strategy of plants and stomatal closure (Medlyn et al., 2016), 917 simulated transpiration is decreased (Sampaio et al., 2021) and photosynthesis is enhanced compared ambient

- 918  $CO_2$ .
- 919 920 To that end, the monthly changes of WUE in JULES CNP are highly correlated to the GPP, hence the lowest
- 921 and highest WUE follow the same periods as GPP similar to responses captured with models studied by
- 922 Fleischer et al., (2019) (Table. S1).
- 923
- 924 Our estimated NPP enhancement using JULES C/CN models for both 1st and 15 years period is within the
- 925 middle range of the models in Fleischer et al., (2019). Nevertheless, JULES CNP response of BP is in the lower
- 926 band of the CNP models by Fleischer et al., (2019) and close to the estimations from CABLE (Haverd et al., 927
- 2018) and ORCHIDEE (Goll et al., 2017) models, which may be due to the similar representation of P processes
- 928 and limitation between these models. However, our results show a 29% decrease in NPP using JULES-CNP 929 compared to JULES-C/CN which is smaller than the differences between the CLM-CNP and CLM-CN versions
- 930 (51% decrease) (Yang et al., 2014). The lower estimated decrease in JULES highlights the need to further study
- 931 the fully corresponding plant C pools and fluxes to the changes in soil and plant P. Therefore, future work
- 932 should be focused on the improvement of the total P availability and the plant C feedbacks. Moreover, there are
- 933 other environmental factors such as temperature which shows a possible impact on the  $CO_2$  elevation and the
- 934 changes of NPP (Baig et al., 2015) which needs further improvement in our model.
- 935 The CUE estimations of 1st year and 15 years response to CO2 elevation from JULES C/CN are in the middle
- 936 range of C/CN models in Fleischer et al., (2019). However, the estimated CUE using JULES CNP for 1st and 15

937 years are in the low range of CNP models reported by Fleischer *et al.*, (2019) which is due to the same reason 938 discussed for NPP comparison.

939

Finally, our estimated total biomass enhancement ( $\Delta Cveg$ ) using JULES C/CN for the 1<sup>st</sup> and 15 years are in the middle range of C/CN models from Fleischer *et al.*, (2019) and in lower range of CNP models from Fleischer *et al.*, (2019) using JULES CNP. Nevertheless, while JULES-CNP includes the trait-based parameters (Harper *et al.*, 2016), other functions such as flexible C allocation and spatial variation of biomass turnover are still missing and future model improvement should be focused on their inclusion.

945

# 946 5. Conclusion

947

Land ecosystems are a significant sink of atmospheric CO<sub>2</sub>, ergo buffering the anthropogenic increase of this
 flux. While tropical forests contribute substantially to the global land C sink, observational studies show that a

950 stalled increase in carbon gains over the recent decade (Brienen et al., 2015; Hubau et al., 2020). However

951 modelling studies that lack representation of P cycling processes predict an increasing sink (Fernández-Martínez

952 *et al.*, 2019; Fleischer *et al.*, 2019). This is particularly relevant for efforts to mitigate dangerous climate change 953 and assumptions on the future efficacy of the land C sink. Therefore, in this study, we presented the full

955 and assumptions on the future efficacy of the fand C sink. Therefore, in this study, we presented the full 954 terrestrial P cycling and its feedback on the C cycle within the JULES framework. Our results show that the

955 model is capable of representing plant and soil P pools and fluxes at a site in Central Amazon. Moreover, the

956 model estimated a significant NPP limitation under ambient CO<sub>2</sub>, due to the high P deficiency at this site which

- 957 is representative of Central Amazon, and elevated CO<sub>2</sub> resulted in a further subsequent decrease in the land C
- 958 sink capacity relative to the model without P limitation. While our study is a step toward the full nutrient cycling
- 959 representation in ESMs, it can also help the empirical community to test different hypotheses (i.e., dynamic
- 960 allocation and stoichiometry) and generate targeted experimental measurements (Medlyn et al., 2015).
- 961

962 <u>Code availability</u>

- 963 The modified version of JULES vn5\_5 and the P extension developed for this paper are freely available on Met
   964 Office Science Repository Service:
- 965 <u>https://code</u>.metoffice.gov.uk/svn/jules/main/branches/dev/mahdinakhavali/vn5.5\_JULES\_PM\_NAKHAVALI/
- 966 after registration (http://jules-lsm.github.io/access\_req/JULES\_access.html) and completion of software license
- 967 form. Codes for compiling model available at: (<u>https://doi</u>.org/10.5281/zenodo.5711160). Simulations were
- 968 conducted using two sets of model configurations (namelists): ambient CO<sub>2</sub> condition
- 969 (https://doi.org/10.5281/zenodo.5711144) and elevated CO<sub>2</sub> condition
- 970 (<u>https://doi</u>.org/10.5281/zenodo.5711150).
- 971 *Data availability*
- 972 The model outputs related to the results in this paper are provided on Zenodo repository
- 973 (https://doi.org/10.5281/zenodo.5710898). All the R scripts used for processing the model outputs and
- 974 producing results in form of table or figures are provided on Zenodo repository
- 975 (https://doi.org/10.5281/zenodo.5710896).
- 976 Author contributions. MAN, LMM, SS, SEC, CAQ, AJW, IAP, KMA and DBC developed the model, per-
- 977 formed simulations and analysis. CAQ, FVC, RP, LFL, KMA, GR, LS, ACMM, JSR, RA and JLC provided the
- 978 measurements for the model parasitisation and evaluation. MAN, LMM, SS, IAP, SEC, FVC, RP, LFL, KMA
- 979 and DBC contributed in writing the manuscript.
- 980
- 981 *Competing interests.* The authors declare no competing interests
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# 993 References:

- 995 Aerts, R. and Chapin, F. S. (1999) 'The Mineral Nutrition of Wild Plants Revisited: A Re-evaluation of
- 996 Processes and Patterns', Advances in Ecological Research, 30(C), pp. 1–67. doi: 10.1016/S0065-
- 997 2504(08)60016-1.
- Anav, A. *et al.* (2013) 'Evaluating the land and ocean components of the global carbon cycle in the CMIP5 earth
   system models', *Journal of Climate*, 26(18), pp. 6801–6843. doi: 10.1175/JCLI-D-12-00417.1.
- 1000 Åragão, L. E. O. C. et al. (2009) 'Above- and below-ground net primary productivity across ten Amazonian
- 1001 forests on contrasting soils', *Biogeosciences Discussions*, 6(1), pp. 2441–2488. doi: 10.5194/bgd-6-2441-2009.
- 1002 Araĝo, L. E. O. C. *et al.* (2009) 'Above- and below-ground net primary productivity across ten Amazonian
- 1003 forests on contrasting soils', *Biogeosciences*, 6(12), pp. 2759–2778. doi: 10.5194/bg-6-2759-2009.
- 1004 Araújo, A. C. *et al.* (2002) 'Comparative measurements of carbon dioxide fluxes from two nearby towers in a 1005 central Amazonian rainforest: The Manaus LBA site', *Journal of Geophysical Research*, 107(D20), p. 8090.
- 1006 doi: 10.1029/2001JD000676.
- Argles, A. P. K. *et al.* (2020) 'Robust Ecosystem Demography (RED version 1.0): A parsimonious approach to
  modelling vegetation dynamics in Earth system models', *Geoscientific Model Development*, 13(9), pp. 4067–
  4089. doi: 10.5194/gmd-13-4067-2020.
- 1010 Arora, V. K. *et al.* (2020) 'Carbon–concentration and carbon–climate feedbacks in CMIP6 models and their
- 1011 comparison to CMIP5 models', *Biogeosciences*, 17(16), pp. 4173–4222. doi: 10.5194/bg-17-4173-2020.
- 1012 Baig, S. et al. (2015) 'Does the growth response of woody plants to elevated CO2 increase with temperature? A
- 1013 model-oriented meta-analysis', *Global Change Biology*, 21(12), pp. 4303–4319. doi: 10.1111/gcb.12962.
- 1014 Baker, T. R. et al. (2004) 'Variation in wood density determines spatial patterns in Amazonian forest biomass',
- 1015 Global Change Biology, 10(5), pp. 545–562. doi: 10.1111/j.1365-2486.2004.00751.x.
- 1016 Bentsen, M. et al. (2013) 'The Norwegian Earth System Model, NorESM1-M Part 1: Description and basic
- evaluation of the physical climate', *Geoscientific Model Development*, 6(3), pp. 687–720. doi: 10.5194/gmd-6687-2013.
- 1019 Best, M. J. et al. (2011) 'The Joint UK Land Environment Simulator (JULES), model description Part 1:
- 1020 Carbon fluxes and vegetation dynamics', *Geoscientific Model Development*, 4(3), pp. 701–722. doi:
- 1021 10.5194/gmd-4-701-2011.
- 1022 Bradford, M. A. and Crowther, T. W. (2013) 'Carbon use efficiency and storage in terrestrial ecosystems', *New* 1023 *Phytologist*, 199(1), pp. 7–9. doi: 10.1111/nph.12334.
- Brienen, R. J. W. et al. (2015) 'Long-term decline of the Amazon carbon sink', Nature, 519(7543), pp. 344-
- 1025 348. doi: 10.1038/nature14283.
- Bruijnzeel, L. A. (1991) 'Nutrient input—output budgets of tropical forest ecosystems: A review', *Journal of Tropical Ecology*, 7(1), pp. 1–24. doi: 10.1017/S0266467400005010.
- 1028 Burke, E. J., Chadburn, S. E. and Ekici, A. (2017) 'A vertical representation of soil carbon in the JULES land
- 1029 surface scheme (vn4.3-permafrost) with a focus on permafrost regions', Geoscientific Model Development,
- 1030 10(2), pp. 959–975. doi: 10.5194/gmd-10-959-2017.
- 1031 Castanho, A. D. A. *et al.* (2013) 'Improving simulated Amazon forest biomass and productivity by including
- spatial variation in biophysical parameters', *Biogeosciences*, 10(4), pp. 2255–2272. doi: 10.5194/bg-10-2255-1033
  2013.
- 1034 Chapin, F. S. *et al.* (2011) *Principles of Terrestrial Ecosystem Ecology*. Springer New York (Biomedical and 1035 Life Sciences). Available at: https://books.google.co.uk/books?id=68nFNpceRmIC.
- 1036 Chave, J. *et al.* (2014) 'Improved allometric models to estimate the aboveground biomass of tropical trees',
- 1037 *Global Change Biology*, 20(10), pp. 3177–3190. doi: 10.1111/gcb.12629.
- 1038 Clark, D. B. et al. (2011) 'The Joint UK Land Environment Simulator (JULES), model description Part 2:
- 1039 Carbon fluxes and vegetation dynamics', *Geoscientific Model Development*, 4(3), pp. 701–722. doi: 1040 10.5194/gmd-4-701-2011.
- 1041 Collatz, G. J. et al. (1991) 'Physiological and environmental regulation of stomatal conductance, photosynthesis
- 1042 and transpiration: a model that includes a laminar boundary layer', Agricultural and Forest Meteorology, 54(2-
- 1043 4), pp. 107–136. doi: 10.1016/0168-1923(91)90002-8.
- 1044 Collatz, G., Ribas-Carbo, M. and Berry, J. (1992) 'Coupled Photosynthesis-Stomatal Conductance Model for
- 1045 Leaves of C4 Plants', *Functional Plant Biology*, 19(5), p. 519. doi: 10.1071/pp9920519.
- 1046 Comins, H. N. and McMurtrie, R. E. (1993) 'Long-Term Response of Nutrient-Limited Forests to CO"2
- 1047 Enrichment; Equilibrium Behavior of Plant-Soil Models', *Ecological Applications*, 3(4), pp. 666–681. doi:
- 1048 10.2307/1942099.
- 1049 Costa, M. G. et al. (2016) 'Labile and Non-Labile Fractions of Phosphorus and Its Transformations in Soil
- 1050 under Eucalyptus', pp. 1–15. doi: 10.3390/f7010015.
- 1051 DeLuca, T. H., Keeney, D. R. and McCarty, G. W. (1992) 'Effect of freeze-thaw-events on mineralization of
- 1052 soil nitrogen', *Biol. Fertil. Soils*, 14, pp. 116–120. doi: 10.1007/BF00336260.

- 1053 Ellsworth, D. S. *et al.* (2017) 'Elevated CO2 does not increase eucalypt forest productivity on a low-phosphorus 1054 soil', *Nature Climate Change*, 7(4), pp. 279–282. doi: 10.1038/nclimate3235.
- 1055 Elser, J. J. et al. (2007) 'Global analysis of nitrogen and phosphorus limitation of primary producers in
- 1056 freshwater, marine and terrestrial ecosystems', *Ecology Letters*, 10(12), pp. 1135–1142. doi: 10.1111/j.1461-1057 0248.2007.01113.x.
- 1057 Fernández-Martínez, M. *et al.* (2019) 'Global trends in carbon sinks and their relationships with CO2 and
- 1059 temperature', *Nature Climate Change*, 9(1), pp. 73–79. doi: 10.1038/s41558-018-0367-7.
- 1060 Fleischer, K. *et al.* (2019) 'Amazon forest response to CO2 fertilization dependent on plant phosphorus
- 1061 acquisition', *Nature Geoscience*. doi: 10.1038/s41561-019-0404-9.
- 1062 Friedlingstein, P. et al. (2006) 'Climate-carbon cycle feedback analysis: Results from the C4MIP model
- 1063 intercomparison', Journal of Climate, 19(14), pp. 3337–3353. doi: 10.1175/JCLI3800.1.
- 1064 Friedlingstein, P. et al. (2019) 'Comment on "The global tree restoration potential", Science. doi:
- 1065 10.1126/science.aay8060.
- Fyllas, N. M. *et al.* (2009) 'Basin-wide variations in foliar properties of Amazonian forest: phylogeny, soils and climate', *Biogeosciences*, 6(11), pp. 2677–2708. doi: 10.5194/bg-6-2677-2009.
- 1068 Gentile, R. et al. (2012) 'Effects of long-term exposure to enriched CO2 on the nutrient-supplying capacity of a
- 1069 grassland soil', *Biology and Fertility of Soils*, 48(3), pp. 357–362. doi: http://dx.doi.org/10.1007/s00374-011-1070 0616-7.
- 1071 Goll, D. S. et al. (2017) 'A representation of the phosphorus cycle for ORCHIDEE (revision 4520)',
- 1072 Geoscientific Model Development, 10(10), pp. 3745–3770. doi: 10.5194/gmd-10-3745-2017.
- 1073 Harper, A. B. et al. (2016) 'Improved representation of plant functional types and physiology in the Joint UK
- 1074 Land Environment Simulator (JULES v4.2) using plant trait information', Geoscientific Model Development,
- 1075 9(7), pp. 2415–2440. doi: 10.5194/gmd-9-2415-2016.
- 1076 Hatfield, J. L. and Dold, C. (2019) 'Water-use efficiency: Advances and challenges in a changing climate',
- 1077 Frontiers in Plant Science, 10(February), pp. 1–14. doi: 10.3389/fpls.2019.00103.
- Haverd, V. *et al.* (2018) 'A new version of the CABLE land surface model (Subversion revision r4601)
- incorporating land use and land cover change, woody vegetation demography, and a novel optimisation-based
   approach to plant coordination of photosynthesis', *Geoscientific Model Development*, 11(7), pp. 2995–3026.
- doi: 10.5194/gmd-11-2995-2018.
  He, M. and Dijkstra, F. A. (2014) 'Drought effect on plant nitrogen and phosphorus: A meta-analysis', *New*
- He, M. and Dijkstra, F. A. (2014) 'Drought effect on plant nitrogen and phosphorus: A meta-analysis', *New Phytologist*, 204(4), pp. 924–931. doi: 10.1111/nph.12952.
- 1084 Hedley, M. J., Stewart, J. W. B. and Chauhan, B. S. (1982) 'Changes in Inorganic and Organic Soil Phosphorus
- 1085 Fractions Induced by Cultivation Practices and by Laboratory Incubations', *Soil Science Society of America*
- 1086 Journal, 46(5), pp. 970–976. doi: https://doi.org/10.2136/sssaj1982.03615995004600050017x.
- 1087 Hou, E. et al. (2019) 'Quantifying Soil Phosphorus Dynamics: A Data Assimilation Approach', Journal of
- 1088 *Geophysical Research: Biogeosciences*, 124(7), pp. 2159–2173. doi: 10.1029/2018JG004903.
- 1089 Hou, E. et al. (2020) 'Global meta-analysis shows pervasive phosphorus limitation of aboveground plant
- 1090 production in natural terrestrial ecosystems', *Nature Communications*, 11(1), pp. 1–9. doi: 10.1038/s41467-020-1091 14492-w.
- 1092 Hubau, W. et al. (2020) 'Asynchronous carbon sink saturation in African and Amazonian tropical forests',
- 1093 Nature, 579(7797), pp. 80-87. doi: 10.1038/s41586-020-2035-0.
- Hungate, B. a et al. (2003) 'Nitrogen and Climate Change', Science, 302(November), pp. 1512–1513.
- 1095 Jenkinson, D. S. et al. (1990) 'The turnover of organic carbon and nitrogen in soil', The Royal Society,
- 1096 329(1255). doi: https://doi.org/10.1098/rstb.1990.0177.
- 1097 Jenkinson, D. S. and Coleman, K. (2008) 'The turnover of organic carbon in subsoils. Part 2. Modelling carbon
- 1098 turnover', *European Journal of Soil Science*, 59(2), pp. 400–413. doi: 10.1111/j.1365-2389.2008.01026.x.
- 1099 Ji, D. et al. (2014) 'Description and basic evaluation of Beijing Normal University Earth System Model (BNU-
- 1100 ESM) version 1', Geoscientific Model Development, 7(5), pp. 2039–2064. doi: 10.5194/gmd-7-2039-2014.
- 1101 Jiang, M. et al. (2019) 'Towards a more physiological representation of vegetation phosphorus processes in land
- 1102 surface models', *New Phytologist*, 222(3), pp. 1223–1229. doi: 10.1111/nph.15688.
- 1103 Jiang, M. et al. (2020) 'The fate of carbon in a mature forest under carbon dioxide enrichment', Nature,
- 1104 580(7802), pp. 227–231. doi: 10.1038/s41586-020-2128-9.
- Johnson, M. O. *et al.* (2016) 'Variation in stem mortality rates determines patterns of above-ground biomass in
- 1106 Amazonian forests: implications for dynamic global vegetation models', *Global Change Biology*, 22(12), pp.
- 1107 3996–4013. doi: 10.1111/gcb.13315.
- 1108 Jones, S. et al. (2020) 'The impact of a simple representation of non-structural carbohydrates on the simulated
- 1109 response of tropical forests to drought', *Biogeosciences*, 17(13), pp. 3589–3612. doi: 10.5194/bg-17-3589-2020.
- 1110 Kattge, J. et al. (2009) 'Quantifying photosynthetic capacity and its relationship to leaf nitrogen content for
- 1111 global-scale terrestrial biosphere models', *Global Change Biology*, 15(4), pp. 976–991. doi:
- 1112 https://doi.org/10.1111/j.1365-2486.2008.01744.x.

- 1113 Keller, M. et al. (2004) 'Ecological research in the Large-scale Biosphere-Atmosphere Experiment in
- 1114 Amazonia: Early results', Ecological Applications, 14(4 SUPPL.), pp. 3–16. doi: 10.1890/03-6003.
- 1115 Klein, T. et al. (2016) 'Growth and carbon relations of mature Picea abies trees under 5 years of free-air CO2 1116 enrichment', Journal of Ecology, 104(6), pp. 1720–1733. doi: 10.1111/1365-2745.12621.
- 1117
- Koch, A., Hubau, W. and Lewis, S. L. (2021) 'Earth System Models Are Not Capturing Present-Day Tropical 1118 Forest Carbon Dynamics', Earth's Future, 9(5), pp. 1-19. doi: 10.1029/2020EF001874.
- 1119 Körner, C. et al. (2005) 'Ecology: Carbon flux and growth in mature deciduous forest trees exposed to elevated
- 1120 CO2', Science, 309(5739), pp. 1360-1362. doi: 10.1126/science.1113977.
- 1121 Van Langenhove, L. et al. (2020) 'Atmospheric deposition of elements and its relevance for nutrient budgets of 1122 tropical forests', Biogeochemistry, 149(2), pp. 175-193. doi: 10.1007/s10533-020-00673-8.
- 1123 Lapola, D. M. and Norby, R. (2014) 'Assessing the effects of increased atmospheric CO 2 on the ecology and
- 1124 resilience of the Amazon forest', Science plan et implementation strategy, AMAZON FAC.
- 1125 LeBauer, D. and Treseder, K. (2008) 'Nitrogen Limitation of Net Primary Productivity', Ecology, 89(2), pp. 1126 371-379.
- 1127 Lloyd, J. et al. (2001) 'Should Phosphorus Availability Be Constraining Moist Tropical Forest Responses to
- 1128 Increasing CO2 Concentrations?', in Global Biogeochemical Cycles in the Climate System. Elsevier, pp. 95-1129 114. doi: 10.1016/B978-012631260-7/50010-8.
- 1130 Long, M. C. et al. (2013) 'Twentieth-century oceanic carbon uptake and storage in CESM1(BGC)', Journal of 1131 Climate, 26(18), pp. 6775–6800. doi: 10.1175/JCLI-D-12-00184.1.
- 1132 De Lucia, E. H. et al. (2007) 'Forest carbon use efficiency: Is respiration a constant fraction of gross primary
- 1133 production?', Global Change Biology, 13(6), pp. 1157–1167. doi: 10.1111/j.1365-2486.2007.01365.x.
- 1134 Lugli, L. F. (2013) Estoque de nutrientes na serrapilheira fina e grossa em função de fatores edáficos em
- 1135 florestas do Amazonas, Brasil. Instituto Nacional de Pesquisas da Amazônia - INPA. Available at: 1136 https://repositorio.inpa.gov.br/handle/1/5028.
- 1137 Lugli, L. F. et al. (2020) 'Multiple phosphorus acquisition strategies adopted by fine roots in low-fertility soils 1138 in Central Amazonia', Plant and Soil, 450(1-2), pp. 49-63. doi: 10.1007/s11104-019-03963-9.
- 1139 Lugli, L. F. et al. (2021) 'Rapid responses of root traits and productivity to phosphorus and cation additions in a 1140 tropical lowland forest in Amazonia', New Phytologist, 230(1), pp. 116-128. doi: 10.1111/nph.17154.
- 1141 Luo, Y. et al. (2004) 'Progressive nitrogen limitation of ecosystem responses to rising atmospheric carbon
- 1142 dioxide', BioScience, 54(8), pp. 731-739. doi: 10.1641/0006-3568(2004)054[0731:PNLOER]2.0.CO;2.
- 1143 Malhi, Y. et al. (2004) 'The above-ground coarse wood productivity of 104 Neotropical forest plots', Global
- 1144 Change Biology, 10(5), pp. 563–591. doi: 10.1111/j.1529-8817.2003.00778.x.
- 1145 Malhi, Y. et al. (2006) 'The regional variation of aboveground live biomass in old-growth Amazonian forests',
- 1146 Global Change Biology, 12(7), pp. 1107–1138. doi: 10.1111/j.1365-2486.2006.01120.x.
- 1147 Malhi, Y. et al. (2009) 'Comprehensive assessment of carbon productivity, allocation and storage in three
- 1148 Amazonian forests', Global Change Biology, 15(5), pp. 1255–1274. doi: 10.1111/j.1365-2486.2008.01780.x.
- 1149 Malhi, Y. (2012) 'The productivity, metabolism and carbon cycle of tropical forest vegetation', Journal of 1150 *Ecology*, 100(1), pp. 65–75. doi: 10.1111/j.1365-2745.2011.01916.x.
- 1151 Malhi, Y., Doughty, C. and Galbraith, D. (2011) 'The allocation of ecosystem net primary productivity in
- 1152 tropical forests', Philosophical Transactions of the Royal Society B: Biological Sciences, 366(1582), pp. 3225-
- 1153 3245. doi: 10.1098/rstb.2011.0062.
- 1154 Martins, N. P. et al. (2021) 'Fine roots stimulate nutrient release during early stages of leaf litter decomposition
- 1155 in a Central Amazon rainforest', Plant and Soil, 469(1-2), pp. 287-303. doi: 10.1007/s11104-021-05148-9.
- 1156 Medlyn, B. E. et al. (2015) 'Using ecosystem experiments to improve vegetation models', Nature Climate 1157 Change, 5(6), pp. 528-534. doi: 10.1038/nclimate2621.
- 1158 Medlyn, B. E. et al. (2016) 'Using models to guide field experiments: a priori predictions for the CO2 response
- 1159 of a nutrient- and water-limited native Eucalypt woodland', Global Change Biology, 22(8), pp. 2834–2851. doi: 1160 10.1111/gcb.13268.
- 1161 Mercado, L. M. et al. (2011) 'Variations in Amazon forest productivity correlated with foliar nutrients and
- 1162 modelled rates of photosynthetic carbon supply', Philosophical Transactions of the Royal Society B: Biological 1163 Sciences, 366(1582), pp. 3316–3329. doi: 10.1098/rstb.2011.0045.
- 1164 Mitchard, E. T. A. (2018) 'The tropical forest carbon cycle and climate change', *Nature*, 559(7715), pp. 527–
- 1165 534. doi: 10.1038/s41586-018-0300-2.
- 1166 Neff, J. C., Hobbie, S. E. and Vitousek, P. M. (2000) 'Nutrient and mineralogical control on dissolved organic
- 1167 C, N and P fluxes and stoichiometry in Hawaiian soils', *Biogeochemistry*, 51(3), pp. 283-302. doi:
- 1168 10.1023/A:1006414517212.
- 1169 Norby, R. J. et al. (2016) 'Model-data synthesis for the next generation of forest free-air <scp>CO</scp>2
- 1170 enrichment ( <scp>FACE</scp> ) experiments', New Phytologist, 209(1), pp. 17–28. doi: 10.1111/nph.13593.
- 1171 Nordin, A., Högberg, P. and Näsholm, T. (2001) 'Soil nitrogen form and plant nitrogen uptake along a boreal
- 1172 forest productivity gradient', Oecologia, 129(1), pp. 125-132. doi: 10.1007/s004420100698.

- 1173 Pan, Y. et al. (2011) 'A Large and Persistent Carbon Sink in the World's Forests', Science, 333(6045), pp. 988-
- 1174 993. doi: 10.1126/science.1201609.
- 1175 Perakis, S. S. and Hedin, L. O. (2002) 'Nitrogen loss from unpolluted South American forests mainly via
- 1176 dissolved organic compounds', Nature, 415(6870), pp. 416-419. doi: 10.1038/415416a.
- 1177 Phillips, O. L. et al. (2004) 'Pattern and process in Amazon tree turnover, 1976-2001', Philosophical
- 1178 Transactions of the Royal Society B: Biological Sciences, 359(1443), pp. 381–407. doi: 10.1098/rstb.2003.1438.
- 1179 Phillips, O. L. et al. (2017) 'Carbon uptake by mature Amazon forests has mitigated Amazon nations' carbon
- 1180 emissions', Carbon Balance and Management, 12(1), pp. 1-9. doi: 10.1186/s13021-016-0069-2.
- 1181 Quesada, C. A. et al. (2010) 'Variations in chemical and physical properties of Amazon forest soils in relation to 1182 their genesis', Biogeosciences, 7(5), pp. 1515-1541. doi: 10.5194/bg-7-1515-2010.
- 1183 Quesada, C. A. et al. (2011) 'Soils of Amazonia with particular reference to the RAINFOR sites',
- 1184 Biogeosciences, 8(6), pp. 1415–1440. doi: 10.5194/bg-8-1415-2011.
- 1185 Ouesada, C. A. et al. (2012) 'Basin-wide variations in Amazon forest structure and function are mediated by
- 1186 both soils and climate', *Biogeosciences*, 9(6), pp. 2203–2246. doi: 10.5194/bg-9-2203-2012.
- 1187 Reed, S. C., Yang, X. and Thornton, P. E. (2015) 'Incorporating phosphorus cycling into global modeling
- 1188 efforts: A worthwhile, tractable endeavor', New Phytologist, 208(2), pp. 324-329. doi: 10.1111/nph.13521.
- 1189 Ryan, M. G. (2013) 'Three decades of research at Flakaliden advancing whole-tree physiology, forest ecosystem
- 1190 and global change research', Tree Physiology, 33(11), pp. 1123-1131. doi: 10.1093/treephys/tpt100. 1191
- Sampaio, G. et al. (2021) 'CO2physiological effect can cause rainfall decrease as strong as large-scale 1192
- deforestation in the Amazon', Biogeosciences, 18(8), pp. 2511–2525. doi: 10.5194/bg-18-2511-2021.
- 1193 Sanchez, P. A. (1977) 'Properties and Management of Soils in the Tropics', Soil Science, 124(3). Available at:
- 1194 https://journals.lww.com/soilsci/Fulltext/1977/09000/Properties and Management of Soils in the Tropics.12. 1195 aspx.
- 1196 Sardans, J., Rivas-Ubach, A. and Peñuelas, J. (2012) 'The C:N:P stoichiometry of organisms and ecosystems in
- 1197 a changing world: A review and perspectives', Perspectives in Plant Ecology, Evolution and Systematics, 14(1), 1198 pp. 33-47. doi: 10.1016/j.ppees.2011.08.002.
- 1199 Schimel, D., Stephens, B. B. and Fisher, J. B. (2015) 'Effect of increasing CO2 on the terrestrial carbon cycle',
- 1200 Proceedings of the National Academy of Sciences of the United States of America, 112(2), pp. 436-441. doi: 1201 10.1073/pnas.1407302112.
- 1202 Shen, J. et al. (2011) 'Phosphorus dynamics: From soil to plant', Plant Physiology, 156(3), pp. 997-1005. doi: 1203 10.1104/pp.111.175232.
- 1204 Sitch, S. et al. (2008) 'Evaluation of the terrestrial carbon cycle, future plant geography and climate-carbon
- 1205 cycle feedbacks using five Dynamic Global Vegetation Models (DGVMs)', Global Change Biology, 14(9), pp. 1206 2015–2039. doi: 10.1111/j.1365-2486.2008.01626.x.
- 1207 Stephenson, N. L. and Van Mantgem, P. J. (2005) 'Forest turnover rates follow global and regional patterns of 1208 productivity', Ecology Letters, 8(5), pp. 524-531. doi: 10.1111/j.1461-0248.2005.00746.x.
- 1209 Sun, Y. et al. (2021) 'Global evaluation of the nutrient-enabled version of the land surface model ORCHIDEE-
- 1210 CNP v1.2 (r5986)', Geoscientific Model Development, 14(4), pp. 1987–2010. doi: 10.5194/gmd-14-1987-2021. 1211 Vicca, S. et al. (2012) 'Fertile forests produce biomass more efficiently', Ecology Letters, 15(6), pp. 520-526.
- 1212 doi: 10.1111/j.1461-0248.2012.01775.x.
- 1213 Vitousek, P. M. et al. (1997) 'Human Domination of Earth Ecosystems', Science, 278(5335), p. 21. Available 1214 at: http://www.cheric.org/research/tech/periodicals/view.php?seq=257860.
- 1215 Vitousek, P. M. et al. (2010) 'Terrestrial phosphorus limitation: Mechanisms, implications, and nitrogen-
- 1216 phosphorus interactions', Ecological Applications, 20(1), pp. 5-15. doi: 10.1890/08-0127.1.
- 1217 Vitousek, P. M. and Howarth, R. W. (1991) 'Nitrogen limitation on land and in the sea: How can it occur?', 1218 Biogeochemistry, 13(2), pp. 87-115. doi: 10.1007/BF00002772.
- 1219 Walker, A. P. et al. (2021) 'Integrating the evidence for a terrestrial carbon sink caused by increasing
- 1220
- atmospheric CO2', New Phytologist, 229(5), pp. 2413-2445. doi: 10.1111/nph.16866.
- 1221 Walker, T. W. and Syers, J. K. (1976) 'The fate of phosphorus during pedogenesis', Geoderma, 15(1), pp. 1–19. 1222 doi: 10.1016/0016-7061(76)90066-5.
- 1223 Wang, Y. P., Houlton, B. Z. and Field, C. B. (2007) 'A model of biogeochemical cycles of carbon, nitrogen, and

1224 phosphorus including symbiotic nitrogen fixation and phosphatase production', Global Biogeochemical Cycles, 1225 21(1), pp. 1–15. doi: 10.1029/2006GB002797.

- 1226 Wang, Y. P., Law, R. M. and Pak, B. (2010) 'A global model of carbon, nitrogen and phosphorus cycles for the 1227 terrestrial biosphere', Biogeosciences, 7(7), pp. 2261–2282. doi: 10.5194/bg-7-2261-2010.
- 1228 Went, F. W. and Stark, N. (1968) 'Mycorrhiza', BioScience, 18(11), pp. 1035–1039. doi: 10.2307/1294552.
- 1229 Wiltshire, A. J. et al. (2021) 'Jules-cn: A coupled terrestrial carbon-nitrogen scheme (jules vn5.1)',
- 1230 Geoscientific Model Development, 14(4), pp. 2161–2186. doi: 10.5194/gmd-14-2161-2021.
- 1231 Xiao, J. et al. (2013) 'Carbon fluxes, evapotranspiration, and water use efficiency of terrestrial ecosystems in
- 1232 China', Agricultural and Forest Meteorology, 182–183, pp. 76–90. doi: 10.1016/j.agrformet.2013.08.007.

- 1233 Xu, Z. et al. (2016) 'Elevated-CO2 response of stomata and its dependence on environmental factors', Frontiers 1234 in Plant Science, 7(MAY2016), pp. 1-15. doi: 10.3389/fpls.2016.00657.
- 1235 Yang, X. et al. (2013) 'The distribution of soil phosphorus for global biogeochemical modeling',
- 1236 Biogeosciences, 10(4), pp. 2525–2537. doi: 10.5194/bg-10-2525-2013.
- 1237 1238 Yang, X. et al. (2014) 'The role of phosphorus dynamics in tropical forests - a modeling study using CLM-
- CNP', Biogeosciences, 11(6), pp. 1667–1681. doi: 10.5194/bg-11-1667-2014.
- 1239 Yang, X. and Post, W. M. (2011) 'Phosphorus transformations as a function of pedogenesis: A synthesis of soil
- 1240 phosphorus data using Hedley fractionation method', Biogeosciences, 8(10), pp. 2907-2916. doi: 10.5194/bg-8-
- 1241 2907-2011.
- 1242 Zaehle, S. and Dalmonech, D. (2011) 'Carbon-nitrogen interactions on land at global scales: Current
- 1243 understanding in modelling climate biosphere feedbacks', Current Opinion in Environmental Sustainability,
- 1244 3(5), pp. 311–320. doi: 10.1016/j.cosust.2011.08.008.
- 1245 Zaehle, S. and Friend, A. D. (2010) 'Carbon and nitrogen cycle dynamics in the O-CN land surface model: 1.
- 1246 Model description, site-scale evaluation, and sensitivity to parameter estimates', Global Biogeochemical Cycles, 1247 24(1), pp. 1–13. doi: 10.1029/2009GB003521.
- 1248 Zhu, Q. et al. (2016) 'Multiple soil nutrient competition between plants, microbes, and mineral surfaces: model
- 1249 development, parameterization, and example applications in several tropical forests', Biogeosciences, 13(1), pp. 1250 341-363. doi: 10.5194/bg-13-341-2016.