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#### 1 Direct evidence for phosphorus limitation on Amazon forest productivity

Hellen Fernanda Viana Cunha<sup>1\*</sup>, Kelly M. Andersen<sup>2,3</sup>, Laynara Figueiredo Lugli<sup>1,4</sup>, 2 Flavia Delgado Santana<sup>1</sup>, Izabela Aleixo<sup>1</sup>, Anna Martins Moraes<sup>1</sup>, Sabrina Garcia<sup>1</sup>, 3 Raffaello Di Ponzio<sup>5</sup>, Erick Oblitas Mendoza<sup>1</sup>, Bárbara Brum<sup>1</sup>, Jéssica Schmeisk Rosa<sup>1</sup>, 4 Amanda L. Cordeiro<sup>6</sup>, Bruno Takeshi Tanaka Portela<sup>1</sup>, Gyovanni Ribeiro<sup>1</sup>, Sara 5 Deambrozi Coelho<sup>1</sup>, Sheila Trierveiler de Souza<sup>1</sup>, Lara Siebert Silva<sup>1</sup>, Felipe Antonieto<sup>1</sup>, 6 Maria Pires<sup>1</sup>, Ana Cláudia Salomão<sup>5</sup>, Ana Caroline Miron<sup>1</sup>, Rafael L. de Assis<sup>1,7</sup>, Tomas 7 F. Domingues<sup>8</sup>, Luiz E.O.C. Aragão<sup>3,9</sup>, Patrick Meir<sup>10</sup>, José Luis Camargo<sup>5</sup>, Antonio 8 Manzi<sup>1</sup>, Laszlo Nagy<sup>11</sup>, Lina M. Mercado<sup>3,12</sup>, Iain P. Hartley<sup>3</sup> and Carlos Alberto 9 Ouesada<sup>1</sup> 10

11 The productivity of rainforests growing on highly-weathered tropical soils is expected to be limited 12 by phosphorus (P) availability<sup>1</sup>. Yet, controlled fertilisation experiments have failed to demonstrate 13 a dominant role for P in controlling tropical forest net primary productivity (NPP). Recent syntheses have demonstrated that responses to N addition are as large as to P<sup>2</sup>, and adaptations to low P 14 15 availability appear to allow NPP to be maintained across major soil P gradients<sup>3</sup>. Thus, the extent to 16 which P availability limits tropical forest productivity is highly uncertain. The majority of the 17 Amazonia, however, is characterised by soils even more depleted in P than where most tropical 18 fertilisation experiments have previously taken place<sup>2</sup>. Thus, we established the first P, nitrogen (N), 19 and base cation addition experiment in an old growth Amazon rainforest, with the site's low soil P 20 content representative of ~60% of the basin. Here we show that NPP increased exclusively with P 21 addition. After 2 years, strong responses were observed in fine root (+29%) and canopy productivity 22 (+19%), but not stem growth. The direct evidence of P limitation of NPP suggests that P availability 23 may restrict Amazon forest responses to CO<sub>2</sub> fertilisation<sup>4</sup>, with major implications for future carbon 24 sequestration and forest resilience to climate change.

25 The inclusion of nutrient cycling in Earth systems models has substantially reduced predictions of future C uptake by vegetation under elevated atmospheric CO<sub>2</sub> (4,5,6,7). Furthermore, fundamental differences 26 27 between the cycles of nitrogen (N) and rock-derived elements such as P, mean that P limitation may place 28 a greater constraint on plant responses to  $CO_2$  fertilisation than N limitation<sup>8,9</sup>. During soil development<sup>10</sup>, 29 the weathering of rocks or parent material provides the major source of P for initial vegetation development. 30 Over millions of years, however, the parent material is gradually depleted, and available P, as well as rock-31 derived base cations such as calcium (Ca), magnesium (Mg) and potassium (K), may be lost via leaching 32 or made unavailable through occlusion by iron and aluminium-oxides, with organic forms of P becoming key pools in depleted and highly weathered systems<sup>10,11</sup>. Meanwhile, N tends to accumulate over time, with 33 34 inputs from biological fixation and atmospheric deposition exceeding N losses<sup>12</sup>. For these reasons, a long-35 standing paradigm in tropical ecology (the so-called P paradigm) has been that forest productivity on

<sup>&</sup>lt;sup>1</sup>Coordination of Environmental Dynamics, National Institute for Amazonian Research, Manaus, AM, Brazil. <sup>2</sup>Asian School of the Environment, Nanyang Technological University, Singapore. <sup>3</sup>Geography, College of Life and Environmental Sciences, University of Exeter, Amory Building, Rennes drive, Exeter, Devon, EX4 4RJ, United Kingdom. <sup>4</sup>TUM School of Life Sciences, Technical University of Munich, Freising 85354, Germany. <sup>5</sup>Biological Dynamics of Forest Fragment Project, National Institute for Amazonian Research, Manaus, AM, 69067-375, Brazil. <sup>6</sup>Colorado State University – CSU, Fort Collins, CO, USA. <sup>7</sup>Natural History Museum, University of Oslo, Norway. <sup>8</sup>Faculdade de Filosofia, Ciência e Letras de Ribeirão Preto, Universidade de São Paulo, Brazil. <sup>9</sup>National Institute for Space Research, São Jose dos Campos, São Paulo, Brazil. <sup>10</sup>School of Geosciences, University of Edinburgh, Edinburg, EH8 9AB, UK. <sup>11</sup>University of Campinas. <sup>12</sup>UK Centre for Ecology and Hydrology, Wallingford, UK. \*email:hfcunha.florestal@gmail.com

highly-weathered soils, such as in those in central Amazonia, is primarily limited by plant available P13, 36 37 with a potential secondary role of other rock-derived elements. Supporting this paradigm, seminal forest 38 ecology studies demonstrated very low levels of P and base cations in plant tissues in Amazonia<sup>14</sup>, and high 39 C:P ratios in litterfall of tropical forest more generally<sup>1</sup>. In Amazonia, greater wood productivity has also 40 been observed in forests growing on fertile soils in western Amazonia when compared to less fertile sites in central and eastern portions of the basin, with relationships with total soil phosphorus being strongest<sup>15,16</sup>. 41 42 However, across the Amazon basin, climatic and edaphic factors covary<sup>17</sup> influencing species distributions, standing forest biomass and turnover rates<sup>16</sup>. Thus, directly determining the extent to which soil fertility 43 44 controls tropical forest growth, and which elements are most important, remains a key knowledge gap<sup>18</sup>, 45 and addressing this is critical for understanding forest growth dynamics and predicting responses to  $CO_2$ 46 fertilisation19.

47 By minimising confounding factors, manipulation experiments can identify directly which specific elements limit forest productivity<sup>20</sup>. Although no large-scale N, P and base cation experiment has been 48 49 carried out in Amazonia until now, a recent synthesis study argued that there is as much evidence for N 50 limitation of tropical forest productivity as there is for P (ref 2). For example, in Costa Rica, P additions did not elicit any changes in litterfall and fine root productivity in two years after fertilisation<sup>21</sup>, and in 51 Panama, an increase in litter production with P addition was evident only 8 years after fertilisation<sup>22</sup>, with 52 53 initial responses stronger for N additions, at least in the rainy season<sup>23</sup>. Critically, previous nutrient 54 manipulation studies in primary tropical rain forests have mainly taken place where total soil P contents are 55 much greater than in central and eastern Amazonia (~443-1600 mg kg<sup>-1</sup> versus 70-120 mg kg<sup>-1</sup> in typical 56 Amazon Ferralsols). In Amazonia, fertilisation experiments have been carried out in secondary forests, but little evidence for strong P limitation has been observed<sup>24,25</sup>, with N availability found to be important 57 58 during initial forest recovery<sup>26,27</sup>. There have been fertilisation experiments in forests growing on soils with 59 P as low as in Amazonia in Cameroon<sup>28</sup> and Borneo<sup>29</sup>. These studies have also generally failed to provide 60 clear support for the P paradigm, with no positive effects of P addition being observed<sup>28</sup>, or with responses 61 to N being at least as large as those to  $P^{29}$ . However, the tree communities were very different to those 62 found across Amazonia, with fundamental differences in nutrient uptake strategies including contrasting mycorrhizal associations. Therefore, while previous fertilisation studies strongly question the ubiquity of P 63 64 limitation in tropical forests, their results cannot be extrapolated to Amazonian forests, especially those 65 growing on low fertility soils in central and eastern regions of the basin.

66 To address this major knowledge gap, in 2017 in lowland tropical evergreen rainforest near Manaus, Brazil, 67 we set up a large-scale fully factorial N, P and base cation-addition experiment (the Amazon Fertilisation Experiment-AFEX), manipulating 8 hectares of forest across 32 plots in four blocks<sup>30</sup>. The Ferralsols of 68 the study site have low concentrations of total P and base cations that are characteristic of up to 60% of 69 70 Amazon forest soils<sup>31</sup> (Fig. 1). To determine directly which nutrient(s) control Amazon forest productivity, 71 we measured the responses of fine root, stem wood, and litterfall production between 2017 and 2019 (see 72 Methods), making nearly 1500 measurements of canopy production, quantifying root productivity every 73 three months across 160 locations and measuring the growth of 4849 trees. Importantly, our base cation 74 treatment added the same amount of calcium as in the super-triple phosphate that was used in the P addition 75 treatment. Thus, comparisons between these treatments ensure that the effects of P can be isolated.

76 Annual NPP rapidly increased with the addition of P in a Central Amazon Forest. After two years of P 77 addition, annual NPP significantly increased by 1.16 Mg C ha<sup>-1</sup> yr<sup>-1</sup>, or 15.6% (+P (with P addition): 8.60  $\pm 0.33$  versus -P (without P addition): 7.44  $\pm 0.21$  Mg C ha<sup>-1</sup> yr<sup>-1</sup>; F<sub>1.27</sub> = 9.56 p = 0.005; Fig. 2a), due to 78 79 greater canopy and fine root productivity. No significant effects of N and base cation addition were 80 observed on total NPP or any of its components measured. The increase in NPP may have been driven by the increase in P availability stimulating GPP<sup>32</sup>, and/or through reductions in autotrophic respiration<sup>33</sup>. It 81 82 has been shown that forests growing on high fertility soils may produce biomass more efficiently and thus 83 show greater carbon use efficiency (CUE, the ratio of net carbon gain to gross carbon assimilated,

NPP/GPP)<sup>34</sup>. Although the direct causes of changes are not yet clear, our results clearly demonstrate that
NPP in this forest is limited by P alone. The observed increase in NPP with +P, and the lack of any N
response, strongly contrasts with a meta-analysis based on previous tropical forest fertilisation studies<sup>2</sup>,
with the lower levels of soil P in Amazonia likely explaining this contrast (Fig. 1). We have previously
observed that base cation addition affects root morphology and mycorrhizal colonisation<sup>30</sup>. Thus, while

89 base cation availability did not appear to limit NPP, they do appear to influence key belowground processes.

90 We observed a substantial 0.83 Mg C ha<sup>-1</sup> yr<sup>-1</sup>, or 19% (+P:  $5.19 \pm 0.15$  versus -P:  $4.36 \pm 0.12$  Mg C ha<sup>-1</sup> 91  $yr^{-1}$ ;  $F_{1,30} = 18.3, p < 0.001$ ; Fig. 2b), increase in canopy productivity. Investment in leaf production provides 92 a return revenue stream of photosynthate that can promote NPP of other tissues and can be used to acquire 93 other limiting resources<sup>35</sup>, such as light and nutrients. We observed weak evidence towards higher leaf area 94 index (LAI) with P addition over the first 1.5 years of the experiment (3.6% increase: +P:  $5.75 \pm 0.10$  versus 95 -P:  $5.55 \pm 0.15$  m<sup>2</sup> m<sup>-2</sup>; F<sub>1,27</sub> = 1.76, p = 0.20; Extended Data Figure 1), which may have had minor 96 contributions to enhanced rates of C gain. The increase in litterfall productivity in our site appears to result 97 from a decrease in leaf life span, which was estimated to have decreased by 10 to 20% following phosphorus 98 addition (+P:  $1.03 \pm 0.04$  versus -P:  $1.15 \pm 0.05$  yr;  $F_{1,30} = 4.08$ , p = 0.05 and +P:  $1.15 \pm 0.05$  versus -P: 99  $1.56 \pm 0.07$ ;  $F_{1,27} = 28.4$ , p = 0.0000127, analysis based on fresh and litter leaves, respectively – see methods; 100 Extended Data Figure 2). Therefore, the increases in leaf turnover appear important in driving the greater

101 canopy productivity in response to P addition, and so far no substantial LAI increment was observed.

Fine root productivity responded strongly to P addition, increasing by 0.35 Mg C ha<sup>-1</sup> yr<sup>-1</sup>, and had the 102 strongest relative increase of 29.4% in the top 30 cm of soil (+P:  $1.54 \pm 0.09$  versus -P:  $1.19 \pm 0.06$  Mg C 103 ha<sup>-1</sup> yr<sup>-1</sup>;  $F_{1,30} = 9.24$ , p = 0.005; Fig. 2b). The overall increase in fine root productivity over two years of 104 105 fertilisation, was greater compared to observations during the first 12 months (23.4% ref 30). Fine root productivity increased significantly in the top 10 cm of soil depth (+P:  $0.96 \pm 0.05$  versus -P:  $0.71 \pm 0.04$ 106 107 Mg C ha<sup>-1</sup> yr<sup>-1</sup>;  $F_{1,30} = 12.9$ , p = 0.001; Table S25-27), but below 10 cm, although fine root productivity was 108  $\sim 20\%$  greater following P addition, this difference was not statistically significant (+P: 0.58 ± 0.04 versus 109 -P:  $0.48 \pm 0.03$  Mg C ha<sup>-1</sup> yr<sup>-1</sup>; F<sub>1.30</sub> = 3.56, p = 0.069; Table S29-30). The greater fine root productivity in the upper soil layer may be due to the low mobility of P in the soil<sup>36</sup>, with most of the added P likely to 110 remain in the top 10 cm, where it can be rapidly taken up by roots<sup>30,37,38</sup>, or soil microbes. In a nearby site, 111 112 at least 40% of fine root productivity was shown to occur below 30 cm<sup>39</sup>. Thus, while it is unlikely that 113 reductions in productivity below 30 cm could have compensated for the increased root growth near the 114 surface, across the full rooting depth the overall stimulation of fine root production will probably have been 115 lower than 29%.

116 There is very limited information on fine root productivity responses to nutrient addition in old growth 117 tropical rainforests. In a fertilisation experiment in Panama, while fine root productivity was not measured 118 directly, K addition induced significant changes, decreasing fine root standing biomass, increasing fine root 119 turnover, and reducing root tissue density, leading to shifts toward the construction of fine roots with a 120 more acquisitive strategy<sup>40,41</sup>. In one of the few studies that measured root productivity responses to large-121 scale nutrient additions in the tropics, in a secondary tropical forest ( $\sim$ 30 years) in Costa Rica, the addition 122 of P did stimulate root productivity one year after fertilisation, but this appeared to be at the expense of 123 aboveground tissue production, with no overall effect of nutrient addition on total productivity<sup>42</sup>. The clear 124 increase in fine root productivity in our experiment also contrasts strongly with results observed in 125 temperate forests, where reductions in root productivity and soil respiration (less heterotrophic and 126 autotrophic respiration) have generally been observed following experimental fertilisation and alleviation 127 of N limitation43.

128 No significant effects of the nutrient addition were detectable on stem wood productivity (P:  $F_{1,24} = 0.001$ , 129 p = 0.97; cations:  $F_{1,27} = 0.01$ , p = 0.92; N:  $F_{1,26} = 0.003$ , p = 0.96). Mean stem wood productivity was 1.85 130  $\pm 0.39$  Mg C ha<sup>-1</sup> year<sup>-1</sup> (DBH > 10 cm). While plants that grow in high-fertility soils can increase the

concentration of nutrients in tissues, with the potential to promote growth<sup>44</sup>, species in low-fertility sites 131 may be adapted to allocate nutrients to tissues with higher P demand (more active), prioritising roots and 132 133 leaves, increasing photosynthetic and metabolic capacities, promoting ion uptake, tissue growth and 134 maintenance<sup>45</sup>. In addition, the advantage of higher woody biomass production occurs only if it provides a 135 competitive advantage over neighbouring trees (competition for light) or decreases the risk of mortality<sup>46</sup>. 136 The rapid responses to P addition observed for the canopy and fine roots are important and enhance our 137 understanding of nutrient limitation in Amazon forests, but longer-term monitoring of the experiment is required to determine whether the responses of different NPP components, and resource allocation, change 138 139 over time, and whether a stem wood productivity response becomes apparent.

140 While attributing variation in forest productivity to P availability across fertility gradients in Amazonian 141 has proven challenging due to confounding variation in tree species composition and both climatic and soil 142 physical factors, our results suggest that P availability may be critical in controlling geographical variation 143 in canopy and fine root productivity across the basin. Along a natural soil fertility gradient spanning the 144 Amazon Basin, fine root productivity, measured in the top 30 cm and extended to 1 m depth, increased on average by ~28% and canopy productivity also increased by ~28% from East (less fertile soils) to West 145 146 (high-fertility soils)<sup>47</sup>. Thus, after two years of P addition, the 29.4% stimulation in fine root productivity in our experiment is comparable to the difference in fine root productivity between Amazon regions with 147 148 contrasting soil fertility (Extended Data Table 1). The observed 19% increase in canopy productivity with 149 P addition (Fig. 2b) is lower than the 28% greater litterfall production in fertile Western forests of the basin 150 (Peru, Colombia), compared with low-fertility sites in Central and Eastern Amazonia (Brazil)<sup>47</sup> (Extended 151 Data Table 1). This may be explained by spatial variability representing the combination of direct P effects 152 as well as changes in the species present, with a greater dominance of fast-growing species with lower wood 153 density in the western Amazon<sup>16</sup>. However, overall, the similar magnitudes of the responses observed in 154 our experiment, in which confounding variation in climatological variables, other edaphic factors, and 155 species present has been minimised, to the patterns observed across major soil fertility gradients, strongly 156 suggest that P availability is a critical in controlling geographical variation in fine root and canopy 157 productivity across the basin.

Direct demonstration of limitation by P, rather than N, of NPP in a Central Amazon forest has major 158 159 implications for predicting forest responses to climate change and rising atmospheric CO<sub>2</sub>. In contrast to 160 the N cycle, the P cycle has no major gaseous phase, and aqueous losses are low<sup>9</sup>. Therefore, while 161 ecosystem N stocks can increase under elevated CO<sub>2</sub> if rates of biological fixation increase, or aqueous or 162 gaseous losses are reduced<sup>8</sup>, in ecosystems with highly weathered soils there is little opportunity for total P 163 stocks to change due the lack of inputs and outputs<sup>9</sup>. For this reason, P limitation may place a stronger constraint on forest responses to rising atmospheric CO<sub>2</sub> than N limitation, questioning the potential for 164 165 current high rates of C uptake in Amazonia to be maintained. Recent model projections demonstrated that 166 the inclusion of P in dynamic global vegetation models reduced predictions of C uptake and biomass 167 production in Amazon forests<sup>4</sup>, decreasing forest C sink, and contributing to more rapid global climate 168 change<sup>7</sup>. Furthermore, because the resistance of tropical forests to climate change depends on their ability 169 to respond positively to rising  $CO_2$  levels, if the responses to elevated  $CO_2$  are limited by P availability, 170 Amazon forests growing in low fertility soils may be more vulnerable than currently recognised<sup>48</sup>. Testing 171 this suggestion directly with experimental manipulations of atmospheric CO<sub>2</sub> in tropical rainforests remains 172 an urgent research priority, with the AmazonFACE (https://amazonface.inpa.gov.br/en/index.php) 173 experiment aiming to do just that. Overall, in contrast to recent meta-analyses and the results from experiments in different tropical regions, our results provide direct evidence for P availability controlling 174 forest productivity in the low fertility soils that characterise central and eastern Amazonia, with no evidence 175 176 for a role of N. This new understanding of the role of nutrient limitation in Amazon forests has critical 177 implications for current and future mitigation policies required to avoid the most dangerous consequences 178 of climate change.

179	1.	Vitousek, P.M. Litterfall, nutrient cycling, and nutrient limitation in tropical forests. <i>Ecology</i> <b>65</b> , 285, 208 (1084)
100	2	203-270 (1704).
101	Ζ.	wright, S. J. <i>et al.</i> Plant responses to return ation experiments in lowland, species rich, tropical $f_{abc}$ (2010)
182	2	The place of the p
183	3.	Turner, B.L <i>et al.</i> Pervasive phosphorus limitation of tree species but not communities in tropical
184		forests. <i>Nature</i> <b>0</b> , 1-4 (2018).
185	4.	Fleischer, K. et al. Amazon forest response to CO <sub>2</sub> fertilization depend on plant phosphorus
186		acquisition. Nat. Geosci. 12, 736-741 (2019).
187	5.	Goll, D. S. et al. Nutrient limitation reduces land carbon uptake in simulations with a model of
188		combined carbon, nitrogen and phosphorus cycling. <i>Biogeosciences</i> 9, 3547-3569 (2012).
189	6.	Sun, Y. et al. Diagnosing phosphorus limitation in natural terrestrial ecosystems in carbon cycle
190		models. <i>Earths future</i> <b>5</b> , 730-749 (2017).
191	7.	Zhang, Q. et al. Nitrogen and phosphorus limitations significantly reduce allowable CO2
192		emissions. Geohys Lett. 41, 632-637 (2014).
193	8.	Luo, Y., Hui, D. & Zhang, D. Elevated CO2 stimulates net accumulations of carbon and nitrogen
194		in land ecosystem: a meta analysis. Ecology 87, 53-63 (2006).
195	9.	Jordan, C.F. The nutrient balance of an Amazonian Rainforest. <i>Ecology</i> 63, 647-654 (1982).
196	10.	Walker, T.W. & Syers, J. K. The fate of phosphorus during pedogenesis. Geoderma 15, 1-19
197		(1976).
198	11.	Crews, T.E. et al. Changes in Soil Phosphorus Fractions and Ecosystem Dynamics across a long
199		chronosequence in Hawai. Ecology 76, 1408-1424 (1995).
200	12.	Hedin, L.O. et al. Nutrient losses over four million years of tropical forest development. Ecology
201		84, 2231-2255 (2003).
202	13.	Dalling, J.W. et al. Nutrient availability in Tropical Rain Forests. The paradigm of phosphorus
203		limitation. In Tropical Tree Physiology (Springer, 2016).
204	14.	Herrera, R.R & Medina, E. Amazon Ecosystems, their structure and functioning with particular
205		emphasis on nutrients. Interciencia 3, 223-231 (1978).
206	15.	Quesada, C.A. et al. Variations in chemical and physical properties of Amazon forest soils in
207		relation to their genesis. Biogeosciences 7, 1515-1541 (2010).
208	16.	Quesada, C.A. et al. Basin wide variations in Amazon forest structure and function are mediated
209		by both soils and climate. Biogeosciences 9, 2203-2246 (2012).
210	17.	Mercado, L. et al. Variations in Amazon forest productivity correlated with foliar nutrients and
211		modelled rates of photosynthetic carbon supply. Philosophical Transitions of the Royal Society
212		<b>366</b> , 3316-3329 (2011).
213	18.	Wright, S.J. 2019. Plant responses to nutrient addition experiments conducted in tropical forests.
214		<i>Ecol. Monogr.</i> <b>0</b> , 1-18 (2019).
215	19.	Yang, X. et al. The effects of phosphorus cycle dynamics carbon sources and sink in the Amazon
216		region: a modelling study using ELM v1. J. Geophys. Res. Biogeosci. 124, 3686-3698 (2019).
217	20.	Sollins, P. Factors influencing species composition in tropical lowland rain forest: Does soil
218		matter? <i>Ecology</i> <b>79</b> , 23-30 (1998).
219	21.	Alvarez-Clare, S. et al. A Direct Test of Nitrogen and Phosphorus Limitation to Net Primary
220		Productivity in a Lowland Tropical Wet Forest. <i>Ecology</i> <b>94</b> , 1540-1551 (2013).
221	22.	Wright, S J. et al. Potassium, phosphorus, or nitrogen limit root allocation, tree growth, or litter
222		production in a lowland tropical forest. <i>Ecology</i> <b>92</b> , 1616-162 (2011).
223	23.	Sayer <i>et al.</i> Variable responses of lowland tropical forest nutrient status to fertilization and litter
224		manipulation. <i>Ecosystems</i> <b>15</b> , 387-400 (2012).
225	24.	Ganade, G. & Brown, V. Succession in old pastures of Central Amazonia: Role of Soil Fertility
226		and Plant Litter. Ecology 83, 743-754 (2002).
227	25.	Markewitz, D. et al. Soil and tree response to P fertilization in a secondary tropical forest supported
228		by an Oxisol. <i>Biol Fertil Soils</i> <b>48</b> , 665-678 (2012).

229	26.	Davidson, E. et al. Nitrogen and Phosphorus limitation of biomass growth in a tropical secondary
230		forest. Ecological Applications 14, 150-163 (2004).
231	27.	Massad, T. et al. Interactions between fire, nutrients, and insect herbivores affect the recovery of
232		diversity in the southern Amazon. Oecologia 172, 219-229 (2013).
233	28.	Newbery, D.M. et al. Does low phosphorus supply limit seedling establishment and tree growth
234		in groves of ectomycorrhizal trees in a central African rainforest? New Phytol. 156, 297-311.
235	29.	Mirmanto, E. et al. Effects of nitrogen and phosphorus fertilization in a lowland evergreen
236		rainforest. Phil. Trans.R.Soc.Lond. 354, 1825-1829 (1999).
237	30.	Lugli, L.F. et al. Rapid responses of root traits and productivity to phosphorus and cation additions
238		in a tropical lowland forest in Amazonia. New Phytol. 230, 116-128 (2020).
239	31.	Quesada, C et al. Soils of Amazonia with particular reference to the rainfor sites. Biogeosciences
240		8, 1415-1440 (2011).
241	32.	Giardina, C. et al. Primary production and carbon allocation in relation to nutrient supply in a
242		tropical experiment forest. Glob Chang Biol 9, 1438-1450 (2003).
243	33.	Rowland, L. et al. Scaling leaf respiration with nitrogen and phosphorus in tropical forests across
244		two continents. New Phytol. 214, 1064-1077 (2017).
245	34.	Vicca, S. et al. Fertile forests produce biomass more efficiently. Ecol. Lett. 15, 520-526 (2012).
246	35.	Wright, I.J., Reich, P.B., Westoby, M. et al. The worldwide leaf economics spectrum. Nature 428,
247		821-826 (2004).
248	36.	Hinsinger, P. How do Plant roots acquire mineral nutrients? Chemical processes involved in the
249		rhizosphere. Adv. Agron. 64 (1998).
250	37.	Van Langehove, L. <i>et al.</i> Rapid root assimilation of added phosphorus in a lowland tropical
251		rainforest of French Guiana. Soil Biol. Biochem. 140, 107646 (2019).
252	38.	Martins, N.P et al. Fine roots stimulate nutrient release during early stages of litter decomposition
253		in a Central Amazon rainforest. <i>Plant Soil</i> <b>469</b> , 287-303 (2021).
254	39.	Cordeiro, A.L et al. Fine root dynamics vary with soil and precipitation in a low-nutrient tropical
255		forest in the Central Amazonia. <i>Plant Environment Interactions</i> <b>220</b> , 3-16 (2020).
256	40.	Yavitt, J. Soil fertility and fine root dynamics in response to four years of nutrient (N,P, K)
257		fertilization in a lowland tropical moist forest, Panamá. Austral Ecol. 36, 433-445 (2011).
258	41.	Wurzburger, N. & Wright, S.J. Fine root responses to fertilization reveal multiple nutrient
259		limitation in a lowland tropical forest. <i>Ecology</i> <b>96</b> , 2137-2146 (2015).
260	42.	Waring, B.G., Aviles, D.P., Murray, J.G. & Powers, J.S. Plant community responses to stand level
261		nutrient fertilization in a secondary tropical dry forest. <i>Ecology</i> <b>0</b> , 1-12 (2019).
262	43.	Jansens, I.A. <i>et al.</i> Reductions of forest soil respiration in response to nitrogen deposition. <i>Nat.</i>
263		<i>Geosci.</i> <b>3</b> , 315- 322 (2010).
264	44.	Alvarez Claire, S. et al. Do foliar, litter, and root nitrogen and phosphorus concentration reflect
265		nutrient limitation in a lowland tropical wet forest? <i>Plos one</i> <b>10</b> , 1-16 (2015).
266	45.	Bouma, T. Understanding Plant Respiration: Separating Respiratory Components versus a
267		Process-Based Aproach. In: Advances in Photosynthesis and Respiration. Vol 18, Springer,
268		Dordrecht, pp. 177-194 (2005).
269	46.	Malhi, Y. <i>et al.</i> Comprehensive assessment of carbon productivity, allocation and storage in three
270		Amazonian forests. <i>Glob Chang Biol.</i> <b>15</b> , 1255-1274 (2009).
271	47.	Aragão, L.E.O <i>et al.</i> Above and below ground net primary productivity across ten Amazonian
272		forests on contrasting soils. <i>Biogeosciences</i> <b>6</b> , 2759-2778 (2009).
273	48.	Cox. P.M. <i>et al.</i> Sensitivity of tropical carbon to climate change constrained by carbon dioxide
274		variability. <i>Nature</i> <b>494</b> , 341-344 (2013).
275	49.	Quesada, C.A & Lloyd, J. Soil-Vegetation Interactions in Amazonia. In Interactions Between
276		Biosphere, Atmosphere and Human Land Use in the Amazon Basin (Springer, 2016).
277	50.	Girardin, C.A.J et al. Seasonal trends of Amazonian rainforest phenology, net primary production.
278		and carbon allocation. Global Biogeochem. Cycles 30, 700-715 (2016).

279

280 Figure 1| Total soil phosphorus measured in primary forest plots across the Amazon Basin, showing

the low P concentration at our site and across central and eastern Amazonia. A fertility gradient across
 the basin is shown, with red circles showing the lowest concentration of total phosphorus and blue circles
 showing the highest concentration of total phosphorus. The two large scale fertilisation experiments in
 Central American *terra firme* tropical forest are also shown, highlighting the five to eighteen-fold greater

total phosphorus concentrations than in central Amazon. Total phosphorus concentrations are derived from
 Quesada and Lloyd 2016 (ref 49), except the values of Costa Rica<sup>21</sup> and Panama<sup>40</sup>. \*In Costa Rica, values
 are available only for the 0-10 cm soil depth. For the other sites, values are for 0-30 cm soil depth.

288 Figure 2 | The effect of N, P and base cation availability on total net primary productivity and its 289 components. a, The responses of total net primary productivity (NPP), representing the sum of NPP 290 components. Only the statistically significant P effects are shown for total NPP, as N, base cation and all 291 interactions had no effect (Table S2-4). b-d, The individual components of NPP where litterfall, stem wood 292 and fine root productivity are shown in green, brown and orange bars, respectively. b, Litterfall productivity 293 showed an increase with P addition only, and base cation (c) and N (d) are shown for comparison (Table 294 S6-8). b, In stem wood productivity there was no effect of any nutrient addition (Table S32-33). b, Fine 295 root productivity (0-30 cm) showed an increase with P addition only, and base cation (c) and N (d) are 296 shown for comparison (Table S21-23). Both 0-10 and 10-30 cm had higher fine root productivity with P 297 addition, but only the 0-10 cm layer had significantly different means. Means  $\pm$  1SE are presented, n=16 298 plots. The dotted lines represent the mean values for the control plots (no nutrients added; n=4 plots) for 299 comparison purposes. Linear mixed models were performed to evaluate responses in total NPP and its 300 components to added nutrients, where nutrient additions and their interactions were fixed effects and block was a random effect with the general full model formula lmer (response ~ N \* P \* Cations + (1|Block). 301 302 Only P addition remained in significant models after model simplification. All differences in mean values 303 between plots with and without added nutrients with p < 0.01 are indicated. Cation (c) and nitrogen (d) 304 panels for NPP components are added for comparison only.

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#### 329 METHODS

330 Site. This research was part of the Amazon Fertilisation Experiment (AFEX), a large-scale fertilisation 331 experiment installed in a lowland tropical forest, 80 km north of Manaus, Brazil, in Central Amazonia (lat  $2^{\circ}$  30 S, long 60° W) at one of the continuous old growth evergreen forests of the Biological Dynamics of 332 Forest Fragments Project (BDFFP)<sup>51</sup>. The experimental site is located in *terra firme* forest and has a high-333 334 species diversity, with about 280 plant species (≥10 cm DBH) per hectare<sup>52</sup>. The dominant tree families in 335 our site are Lecythidaceae, Sapotaceae, Fabaceae and Burseraceae, and the most abundant species are 336 Micrandropsis scleroxylon, Protium hebetatum, Eschweilera wachenheimii, Scleronema micranthum and 337 Eschweilera truncata.

The mean annual air temperature is c. 26  $^{\circ}C^{53}$ , and the mean annual precipitation is 2400 mm with a dry 338 339 season from June to October, when monthly precipitation can reach less than than 100 mm<sup>54</sup>. Above ground 340 biomass (AGB) was estimated to be  $322 \pm 54$  Mg ha<sup>-1</sup> (tree individuals  $\geq 10$  diameter at breast height -DBH) with mean wood density of 0.67 g cm<sup>-3 55</sup>. Local soils are geric Ferrasols (WRB Soil Classification) 341 also known as Oxisols (USDA Soil Taxonomy) <sup>56,57</sup>. The soils are deep ( $\geq$  400 cm) with good particle 342 aggregation, friable and with low subsoil bulk density  $(0.8 - 1.2 \text{ g cm}^{-3})^{58}$ , typically acidic (pH ~ 4.1), with 343 344 low concentrations of nutrients such as P (total  $P = 87.5 \text{ mg kg}^{-1}$ ), calcium (Ca) (0.034 cmolc kg<sup>-1</sup>), and K 345  $(0.066 \text{ cmolc kg}^{-1})$ . The soil texture of the site is 7.69% sand, 14.75% silt, and 77.55% clay.

Experimental design. AFEX is composed of thirty-two 50 m x 50 m plots distributed across four blocks
separated by at least 200 m<sup>30</sup>. Each of the four blocks comprises eight plots, which are separated by at least
50 m, representing eight treatments applied in a fully factorial design: control (with no addition of
nutrients), N, P, CATIONS (Ca, Mg, K), N+P, N+CATIONS, P+CATIONS and N+P+CATIONS.

Fertilisation consists of 125 kg ha<sup>-1</sup> year<sup>-1</sup> of N as urea (CO(NH<sub>2</sub>)<sub>2</sub>), 50 kg ha<sup>-1</sup> year<sup>-1</sup> of P as triple 350 superphosphate  $(Ca(H_2PO_4)_2)$  and base cations with 160 kg ha<sup>-1</sup> year<sup>-1</sup> as dolomitic limestone  $(CaMg(CO_3)_2)$ 351 352 for Ca and Mg plus 50 kg ha<sup>-1</sup> year<sup>-1</sup> as potassium chloride (KCl) for K. Annual doses of N, P and K are similar to the Panama fertilisation experiment, in order to facilitate comparisons<sup>59</sup>, while the addition rates 353 of Ca within the base cation treatment equals the addition rate of Ca in the triple superphosphate, allowing 354 355 us to directly determine the effect of the added P. Nutrient additions are split into three equal applications 356 over the course of each wet season, with nutrients added every year since May 2017. The results presented 357 here correspond to forest growth after 2 years of field measurements.

Fine root productivity. The productivity of fine roots was measured every three months using the ingrowth core method as described in detail in Lugli et al. (2021). In each plot, the five ingrowth cores were bulked into a composite sample per plot, divided into depths of 0-10 cm and 10-30 cm, and roots were removed from the soil core by hand in the field over a period of 60 minutes, which was split into 15 minutes time intervals. Subsequently, fine roots (<2 mm diameter) were cleaned, dried at 60 °C until constant mass and weighed.</p>

Different curve types were fitted to the first 60 minutes of manual root extraction and used to predict the
 pattern of extraction up to 180 minutes<sup>30,60</sup>.

We used the census from November 2017 to September 2019, comprising two years of data collection (Year 1: November 2017 to Sept 2018 and Year 2: Dec 2018 to Sep 2019 in a total of 8 ingrowth core collections). Total fine root productivity (0-30 cm) was summed for both years and the annual mean root productivity was obtained dividing the root productivity by two. To convert root productivity from biomass to C, we used C data from the root tissues carried out in the study area<sup>30</sup>, in which the average C

371 concentration was 43.94%. Fine root productivity was expressed in Mg C ha<sup>-1</sup> year<sup>-1</sup>.

**Stem wood productivity.** To calculate stem wood productivity, the stem diameter of all identified trees with a diameter at breast height (DBH)  $\geq$  10 cm were recorded annually at the end of the wet season (May) from 2017 - 2019. An allometric equation specific for tropical moist forest<sup>61</sup> was applied to convert tree DBH (cm), species wood density (g cm<sup>-3</sup>) and a bioclimatic parameter (*E*) in woody biomass. The equation has the following expression:

377 AGB =  $\exp(-2.024 - 0.896E + 0.920 \ln (WD) + 2.795 \ln (D) - 0.0461 [\ln (D)^2])$ 

This is the slightly modified Eq 7 of Chave et al. 2014 given by the biomass package, where woody biomass
can be inferred in the absence of height measurements. The bioclimatic parameter (E) is a measure of
environmental stress<sup>61</sup> related to climatic water deficit, temperature seasonality and precipitation
seasonality, inferred when the site coordinates were given (lat 2° 40'S; long 60° W).

Wood density was estimated for each species from the *getWoodDensity* function from R *biomass* package using the global wood density database as a reference<sup>62,63</sup>, ideally assigned to species, but to genus level where species-level wood density data were not available. Of the total number of individuals, 55.1% of the wood densities were obtained at the species level, 37.1% at the genus level and for the remaining 7.9% of the individuals, we assumed the average wood density of the plot, because species was not identified or was absent in the database.

388 Stem wood productivity was calculated as the change in stem biomass of surviving trees added to the 389 biomass of the recruited individuals divided by the census length. For 4600 tree individuals, we selected a 390 census length of two years (2017-2019) and for 249 trees where one census was missing (e.g.: tree not 391 measured in 2017, recruited in 2018 census, measurement error), annual productivity was calculated using 392 one year interval (2017-2018 or 2018-2019). Recruitment was the inclusion of new individuals who reached 393 10 cm of DBH in the 2019 inventory (42 trees). 22 trees with DBH > 15 cm in 2019 that were not measured 394 in at least two censuses were not considered in the analyses. For 38 trees that died in 2019, productivity 395 was calculated by the difference in biomass between 2018 and 2017.

396 The change in biomass was then summed over all trees  $\geq 10$  cm DBH in each plot (2500 m<sup>2</sup>) and 397 extrapolated to estimate the change in biomass per hectare. To convert biomass values into C, we assumed 398 that dry stem biomass corresponds to 50% C<sup>64</sup> and stem wood productivity was expressed in Mg C ha<sup>-1</sup> year-1. To avoid or minimise potential errors, we used some parameters to check for quality control of the 399 400 data. We used data that fell inside both of the following criteria: diametric growth smaller than 4 cm yr<sup>1</sup> 401 and a negative growth limit of -0.5 cm across the census intervals. Small negative DBH increments were 402 included to accommodate measurement error and also because trees may shrink by a small amount due to 403 hydrostatic effects in times of drought<sup>65</sup>.

Litterfall Productivity. Litterfall production was estimated by sampling litterfall every fifteen days in five
 litter traps (0.25 m<sup>2</sup>) placed 1 m above the ground within the central area of each plot (30 x 30 m). Litterfall
 includes leaves, twigs and thin branches with diameter < 2 cm, reproductive material (flowers, fruits and</li>
 seeds), residues (other fractions not identified) and insect frass that were oven-dried at 65 °C to constant
 mass and weighed.

We used data from the census of July 2017 to June 2019, where this period comprises two years. Litterfall productivity in g m<sup>-2</sup> day<sup>-1</sup> was extrapolated to Mg ha<sup>-1</sup> year<sup>-1</sup> and the average was obtained considering two years of collection (Moraes et al, in prep; Supplementary material). Litter material was estimated to be 50%
C, based on mean values in our site, to convert biomass productivity into C productivity and it was also expressed in Mg C ha<sup>-1</sup> year<sup>-1</sup>.

Leaf area index (LAI). A LAI-2200C (LI-COR Biotechnology, Lincoln, Nebraska USA) was used to 414 415 measure LAI inside the central 30 m x 30 m of each plot. Sixteen measurement points were made in each 416 plot, on a grid with an even spacing of 10 m. Measurements made on these 16 points per plot were averaged to represent plot means. The data were collected from 6 am to 5 pm, avoiding recording data between 12:00 417 418 and 2:00 pm, to avoid direct sun. The LAI-2200C requires an above canopy reading for reference, and in 419 our case the optical sensor was placed in a clearing to log automatically while the operator collected 420 manually below the canopy. The sensors were always placed in the same compass direction (both in the 421 west in the morning and east in the afternoon) and we used a view cap of  $45^{\circ}$  in the sensors to remove the 422 operator from the sensor's view. The sensors were matched before the data collection. The raw data were 423 analysed using the FV2200 software, where LAI was obtained (m<sup>2</sup> one sided foliage area/ m<sup>2</sup> ground area) 424 and computed with 4 rings. These four rings read radiation in 4 angles, which are 7°, 23°, 38° and 53°. The 425 data were collected during 10 to 13 October 2017, 22 to 25 March 2018, 07 to 10 August 2018 and between 426 29 October and 02 November 2018. LAI was based on these 4 collections, and was transformed to a single

- 427 value representing the mean LAI over one year.
- 428 Total Productivity. We calculated total productivity, using the following equation:
- 429  $NPP_{total} = NPP_{fineroots} + NPP_{stem} + NPP_{litterfall}$
- 430 All terms are expressed in Mg C ha<sup>-1</sup> year<sup>-1</sup>.

431 Leaf residence time. This parameter was calculated by dividing the leaf biomass by annual leaf fall productivity (from July 2017 to July 2018) in Mg dry biomass ha<sup>-1</sup> yr<sup>1</sup> (<sup>66</sup>). Leaf biomass was calculated by 432 dividing the mean LAI of four campaigns (10 to 13 October 2017, 22 to 25 March 2018, 07 to 10 August 433 434 2018 and between 29 October and 02 November 2018) by specific leaf area (SLA). The SLA was included 435 in two approaches: 1) Obtained from a census in October 2018, from about 8 individuals per plot from canopy dominant trees (-P:  $83.36 \pm 1.83$  cm<sup>2</sup> g<sup>-1</sup> and +P:  $88.02 \pm 2.49$  cm<sup>2</sup> g<sup>-1</sup>, -CATIONS:  $85.61 \pm 2.25$ 436  $cm^2 g^{-1}$  and +CATIONS: 85.77 ± 2.28  $cm^2 g^{-1}$ , -N: 85.54 ± 2.67  $cm^2 g^{-1}$  and +N: 85.85 ± 1.76  $cm^2 g^{-1}$ , based 437 438 on mean values in our site; Andersen et al, unpublished) 2) Obtained from sampling in litter traps (-P: 439  $162.50 \pm 26$  g m<sup>-2</sup> and +P:  $128.75 \pm 11$  g m<sup>-2</sup>). Transformations from LMA to SLA were made when 440 necessary. The numerator, leaf biomass in g m<sup>2</sup> was extrapolated to Mg ha<sup>-1</sup>. The denominator, leaf fall 441 productivity was based on 24 collections, and was transformed to a single value representing the mean leaf 442 fall productivity over one year.

443 Data analyses. Linear mixed models were used to test the effect of added nutrients and their interaction in 444 the factorial design N\*P\*base cations. The model simplification method used to find the best model was 445 the step function in *ImerTest* package, based on the drop1 function which systematically drops fixed factors 446 in order of the model hierarchy<sup>67</sup>. We started with the full model including all nutrients and their interaction, 447 and followed a stepwise backward elimination on non-significant effects based on chi square test comparing 448 two consecutive models. When dropping interaction effects significantly changed the model fit, they were 449 retained in the model and the elimination process was completed. When all fixed effects were dropped from 450 the model, the intercept was accepted as the final model. A probability <0.05 was adopted to determine 451 significance. Results are reported for the best fit model in the text and figures. The denominator degrees of 452 freedom were estimated using the Satterthwaite approximation. The four blocks were used as random factors and the response variables were fine root, stem wood, litterfall productivity, total productivity, leaf 453 area index and leaf residence time. All models were run using *lme4* and *lmerTest* R packages<sup>68</sup>. We tested 454

the assumptions for normality and homogeneity of variance to meet assumptions for linear models, using the Shapiro-Wilk and Levene tests. Since no interactions between nutrients were found, all plots where a specific nutrient was not added (i.e. – P, n = 16) are compared to all plots where that nutrient was added (i.e., +P, n = 16)<sup>22,30</sup>. Original datasets from this study are publically available (Moraes et al. 2020<sup>69</sup>, Cunha et al. 2021a<sup>70</sup>, Cunha et al. 2021b<sup>71</sup>, Cunha et al. 2021c<sup>72</sup>). Compiled datasets and R scripts used for

460 statistical analyses, figures and tables are available at https://github.com/kmander7/Paper-AFEX-NPP.

461 Data availability. Data that support the findings of this study have been deposited in NERC Environmental 462 Information Data Centre at (https://doi.org/10.5285/b3a55011-bf46-40f5-8850-86dc8bc4c85d) for root 463 biomass, at (https://doi.org/10.5285/c2587e20-ba4a-4444-8ce9-ccdec15b0aa3) for tree census, at (https://doi.org/10.5285/c0294ec9-45d6-464c-b543-ce9ece9fd968) for litterfall production and at 464 465 (https://doi.org/10.5285/6e70665f-b558-4949-b42a-49fbaec7e7cc) for leaf area index. Global Wood 466 Density Database can be requested from http://datadrayad.org/handle/10255/dryad.235. Plot mean datasets 467 for all response variables and AFEX plot treatment identifications are available at 468 https://github.com/kmander7/Paper-AFEX-NPP.

469 Code availability. The R code used to find the best model for each variable is available in the
 470 Supplementary material. R scripts used to generate the Supplementary material are available at
 471 https://github.com/kmander7/Paper-AFEX-NPP

- 472 51. Laurance, W.F. *et al.* An Amazonian rainforest and its fragments as a laboratory of global change. *Biol.*473 *Rev.* 93, 223-247 (2018).
- 52. De Oliveira, A. & Mori, S.A. A central Amazonia terra firme forest. I. High tree species richness on
  poor soils. *Biodivers. Conserv.* 8, 1219-1244 (1999).
- 476 53. Ferreira, S. J. F., Luizão, F. J., & Dallarosa, R. L. G. Precipitação interna e interceptação da chuva em
- 477 floresta de terra firme submetida à extração seletiva de madeira na Amazônia Central. *Acta Amaz.* 35, 55478 62 (2005).
- 479 54. Tanaka, L. D. S., Satyamurty, P., & Machado, L. A. T. Diurnal variation of precipitation in central
  480 Amazon Basin. *Int J Climatol.* 34, 3574-3584 (2014).
- 481 55. Duque, A. *et al.* Insights into regional patterns of Amazonian forest structure and dominance from three
  482 large terra firme forest dynamics plots. *Biodivers. Conserv.* 26, 669-686 (2017).
- 483 56. Quesada C.A *et al.* Soils of Amazonia with particular reference to the rainfor sites. *Biogeosciences* 8, 1415-1440 (2011).
- 485 57. Quesada, C.A. *et al.* Variations in chemical and physical properties of Amazon forest soils in relation
  486 to their genesis. *Biogeosciences* 7, 1515-1541 (2010).
- 58. Martins, D. L. *et al.* Soil induced impacts on forest structure drive coarse wood debris stocks across
  central Amazonia. *Plant Ecol. & Divers.* 8, 229-241 (2014).
- 489 59. Wright, S. J. *et al.* Potassium, phosphorus, or nitrogen limit root allocation, tree growth, or litter
  490 production in a lowland tropical forest. *Ecology* 92, 1616-1625 (2011).
- 491 60. Metcalfe, D.B. *et al.* A method for extracting plant roots from soil which facilitates rapid sample 492 processing without compromising measurent accuracy. *New Phytol.* **174**, 697-703 (2007).
- 493 61. Chave, J. et al. Improved allometric to estimate the above ground biomass of tropical trees. Glob
- 494 *Chang Biol.* **20**, 3177-3190 (2014).
- 495 62. Chave J. et al. Towards a worldwide wood economics spectrum. Ecol. Lett. 12, 351-366 (2009).
- 496 63. Zanne, A. E., Lopes Gonzales, G.,Gomes, D. A *et al.* Global wood density database. Dryad Digital
  497 Repository, http://datadrayad.org/handle/10255/dryad.235 (2009).
- 498 64. Higuchi, N. & Carvalho, J. A. Fitomassa e conteúdo de carbono de espécies arbóreas da Amazônia.
- 499 Anais do Seminário: Emissão e Sequestro de  $CO_2$  Uma Nova Oportunidade de Negócios para o Brasil
- **500 12**, 7-153 (1994).
- 501 65. Brienen, R. J.W., Philips, O. L. & Zagt, R J. Long term decline of the Amazon carbon sink. *Nature* 519,
- **502** 344-348 (2015).

- 503 66. Malhado, A.C.M. *et al.* Seasonal leaf dynamics in an Amazonian tropical forest. *Forest Ecol Manag.*504 258, 1161-1165.
- 505 67. Kuznetsova, A., Brockhoff, P.B & Christensen, R.H.B. ImerTest Package: Tests in Linear Mixed 506 Effects Models. *Journal of Statistical Software* 82, 1-26.
- 507 68. Bates, D., Marcher, M., Bolker, B. M. & Walker, S C. Fitting Linear Mixed Effects Models using lme4.
  508 *J. Stat. Softw.* 67, 1-48 (2015).
- 69. Moraes, A.C.M. *et al.* Fine litterfall production and nutrient composition data from a fertilized site in
  Central Amazon, Brazil. NERC Environmental Information data Centre. (2020).
- 511 70. Cunha, H.F.V. *et al.* Fine root biomass in fertilised plos in the Central Amazon, 2017-2019. NERC
- 512 Environmental Information data Centre. (2021a).
- 513 71. Cunha, H.F.V. et al. Tree census and diameter increment in fertilised plos in the Central Amazon, 2017-
- 514 2020. NERC Environmental Information data Centre. (2021b).
- 515 72. Cunha, H.F.V. *et al.* Leaf area index (LAI) in fertilised plos in the Central Amazon, 2017-2018. NERC
  516 Environmental Information data Centre. (2021c).
- 517

**Extended Data Figure 1** Nutrient addition effects on Leaf area index. LAI was measured over four field campaigns across treatments in a lowland forest in Central Amazon. Each panel represents mean  $\pm$ 1SE LAI with (+) or without (-) the addition of specific nutrients (phosphorus addition (a); base cation addition (b); nitrogen addition (c)), based on the average LAI across the four field campaigns, n= 16 plots. No significant differences among the means were detected in linear mixed models for any of the nutrients. The dotted lines represent the mean values for the control plots (no nutrients added; n = 4 plots) for comparison purposes.

525

526 Extended Data Figure 2| Nutrient addition effects on Leaf residence time (LRT). Leaf residence time 527 (yr) across treatments in a lowland forest in Central Amazon. Two separate measures of specific leaf area 528 were used in the leaf residence time calculations based on: 1) fresh canopy leaves of common families 529 represented across all plots sampled for a photosynthesis campaign (a-c); 2) composite leaf litter collected in the plots (d-f). Leaf residence time showed a decrease with P addition only (a, d) for both LRT estimates, 530 531 with cations (**b**, **e**) and N (**c**, **f**) being shown for comparison. Means  $\pm 1$ SE are presented, n= 16 plots. Linear 532 mixed models were performed to evaluate responses in leaf residence time to added nutrients. The dotted 533 lines represent the mean values for the control plots (no nutrients added; n = 4 plots) for comparison 534 purposes.

535 Extended Data Table 1 NPP comparisons along the Basin. Total P (mg kg<sup>-1</sup>), N (%) and sum of base cations (SB in cmol<sub>c</sub> kg<sup>-1</sup> refer to the sum of Ca+Mg+K+Na), canopy, fine roots and stem wood net primary 536 537 productivity (Mg C ha<sup>-1</sup> yr<sup>-1</sup>), from low fertility soils in eastern Amazonian sites (CAX 03, MAN 05, CAX 538 06) and more fertile soils in western sites (TAM 05, AGP 02, TAM 06) according to their total soil P 539 concentrations. Components of net primary productivity are derived from Aragão et al. 2009. Aragão et al. 540 2009 presents fine root productivity to 1 m, so we have extended our data to 1 m by dividing by 0.6, based 541 on the study of Cordeiro et al. 2020 that demonstrated that 40% of fine root productivity was located below 542 30 cm at a nearby site on the same soil type. The percentage indicates the magnitude of differences between

543 more fertile and least fertile sites.



