- ¹ Title: Global patterns of forest autotrophic carbon fluxes
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Abstract

[very rough start:] Carbon fixation, allocation, and metabolism by trees set the basis for energy and material flows in forest ecosystems and define their interactions with Earth's changing climate. Here, we drew upon # records from the Global Forest Carbon Database (ForC), representing all major forest types 27 and the nine most significant forest autotrophic carbon flux (FACF) variables, to comprehensively explore 28 how C cycling in mature, undisturbed forests varies with latitude and climate on a global scale. We show 29 that, across all FACF variables analyzed, C cycling decreases linearly with latitude – a finding that confirms multiple previous studies but contradicts the idea that net primary productivity (NPP) of temperate forests 31 rivals that of tropical forests. The FACF variables generally increase in proportion to one another, with 32 few differences in allocation detectable at this global scale, but differed in that latitude explained a lower proportion of variation among subsidiary fluxes (in particular, woody aboveground NPP and belowground 34 NPP, BNPP). Climate explained a significant proportion (#-#%) of variation in all C fluxes analyzed (less for subsidiary fluxes), with temperature variables in general and mean annual temperature (MAT) in particular being the best predictors of FACF on this global scale. While other climate variables (e.q., XX)37 displayed significant correlation with FACF, none of them had significantly better explanatory power than MAT. The effects of temperature were modified by moisture availability, with reduced FACF under hot and dry conditions and sometimes under very high precipitation (especially for BNPP). FACF declined with temperature seasonality, but growing season length doesn't improve upon MAT as a predictor. Within the 41 growing season, the influence of climate on C cycling is smaller but still significant for a number of carbon fluxes. These findings clarify the big picture of how FACF varies with latitude and climate on a global scale. As we enter a period of accelerating climatic change, understanding of the fundamental climatic controls on FACF sets a foundation for understanding patterns of change.

46 Introduction

Carbon cycling in forests worldwide provides the energetic basis for sustaining the majority of Earth's terrestrial biodiversity, while strongly influencing atmospheric CO₂ and climate (Bonan, 2008). Forests' autotrophic carbon fluxes (FACF)-that is, carbon fixation, allocation, and metabolism by trees and other primary producers-sets the energy ultimately available to heterotrophic organisms (including microbes), in turn influencing their abundance (REFS) and possibly diversity (Waide et al., 1999) (REFS). FACF 51 influences all organic matter stocks in forest ecosystems and is linked to cycling of energy, water, and nutrients (REFS) (Piao et al., 2010). Critically, FACF also define forest interactions with Earth's changing climate. Over 69 Gt of CO₂ cycle through Earth's forests each year (Badgley et al., 2019), and in recent decades 54 their net C sequestration (~2.4 Gt C yr⁻¹) offset roughly 30% of anthropogenic fossil fuel emissions (Pan et al., 2011). As atmospheric carbon dioxide levels continue to rise, driving climate change, forests will play a critical role in shaping the future of Earth's climate (Cavaleri et al. 2015; Rogelj et al. 2018). However, our 57 ability to draw general macroscopic conclusions regarding global variation in FACF with respect to climate has been limited in that these analyses often mix forests that vary in stand age, disturbance history, and/or management status; do not always sufficiently parse related variables; and typically consider only one or a few variables at a time.

FACF decrease with latitude, but it remains unclear whether and how the shape of this relationship varies across fluxes. Studies agree that FACF are lowest in the boreal regions, and increase into the temperate regions (Luyssaert et al., 2007; Huston and Wolverton, 2009; Beer et al., 2010; Jung et al., 2011). However, 64 evidence is inconclusive on whether primary productivity continues to increase into the tropics, or whether it plateaus in temperate regions. Evidence for this is further complicated by the fact that different studies use different measures of productivity to explore these relationships. For example, modelling of global terrestrial 67 ecosystem gross primary productivity (GPP) through upscaling and calibration of eddy flux measurements indicates that GPP peaks in tropical forests (Beer et al., 2010; Jung et al., 2011; Badglev et al., 2019) ($Li \mathcal{E}$ 69 Xiao 2019). In contrast, some studies suggest that the highest values of net primary productivity (NPP) may be found in temperate forests (Luyssaert et al., 2007; Huston and Wolverton, 2009), while others find 71 NPP highest in the tropics and decreasing with latitude (Šímová and Storch, 2017). Other studies have chosen to focus exclusively on above-ground net primary productivity (ANPP), finding evidence of a weak 73 negative relationship between ANPP and latitude (Huston and Wolverton, 2009; Gillman et al., 2015).

The latitudinal gradient in FACF is primarily driven by climate, which is a significant driver of FACF across broad spatial scales (Luyssaert et al., 2007; Cleveland et al., 2011)(Hursh et al. 2017). The majority of studies have focused on exploring the relationships of FACF to mean annual temperature (MAT) and mean annual precipitation (MAP), as the most commonly reported site-level climate variables. While these fail to capture some important aspects of climate such as seasonality, they do describe broad trends in temperature and water availability, and therefore capture a substantial portion of global-scale variation in climate. There is strong evidence that both MAT and MAP show significant positive relationships with FACF (Chu et al., 2016). However, as with latitude, the shape of those relationships is not always clear, and, again, is complicated by the use of different measures of FACF across studies. Various measures of primary productivity {FACF?} saturate at high levels of MAP, though the saturation points identified vary from 1500mm (Luyssaert et al., 2007) up to 2445mm MAP (Schuur, 2003). Studies of the influence of MAT on productivity {FACF?} are less conclusive. Luyssaert et al. (2007) examined GPP and NPP and found that, while GPP increases linearly with MAT, NPP saturates at around 10°C MAT. In contrast,

Larjavaara and Muller-Landau (2012), find that increases in *GPP* saturate at approximately 25°C MAT, while Schuur (2003) shows that *NPP* increases linearly with temperature. Taylor et al. (2017) showed a positive interaction between MAT and MAP in shaping tropical forest productivity, such that high rainfall had a negative effect on productivity in cooler climates, compared to a positive effect in warmer climates.

{It would be good to add some more citations on soil respiration. I'm sure BBL can help.}

FACF can be influenced by many other factors as well, which often act across a range of scales, and may show interactive effects with each other (Cleveland et al., 2011). On a local scale, stand age (Litton et al., 2007; 94 Gillman et al., 2015), biodiversity (Liang et al., 2016), management (Šímová and Storch, 2017), nutrient 95 availability (Aragão et al., 2009), and altitude (Girardin et al., 2010; Malhi et al., 2017) all impact FACF. On a global scale, we expect that FACF are most strongly influenced by broad climatic gradients. There 97 is evidence that FACFs also respond to variables such as cloud cover (Taylor et al., 2017), solar radiation (Fyllas et al., 2017), and potential evapotranspiration (Kerkhoff et al., 2005) in potentially significant ways. Furthermore, MAT and MAP are very coarse measures of climate, and so fail to capture much variation 100 in climate on an intra-annual scale, including the effects of factors such as growing season length, number 101 of frost-free days, temperature seasonality, and dry season length. Some studies have suggested that the 102 apparently strong relationship between MAT and FACFs is actually a consequence of the correlation between 103 MAT and growing season length (Kerkhoff et al., 2005; Malhi, 2012; Michaletz et al., 2014, 2018). Kerkhoff et 104 al. (2005) and Michaletz et al. (2014) find that, within the growing season, there is no significant relationship 105 between primary productivity and MAT, suggesting that the effect of temperature is due to increased length 106 of growing season, rather than an inherent influence of temperature on FACF. 107

The recent development of a global forest carbon database synthesizing multiple variables and including 108 records of stand history (ForC; (Anderson-Teixeira et al., 2016, 2018)) opens up the possibility for a stan-109 dardized analysis of global scale variation in multiple FACF and the principle climatic drivers of these 110 patterns. In order to approach this broad topic, we simplify the major gaps in our knowledge to five broad 111 questions and corresponding hypotheses (Table 1). ** see issue #48** First, we ask how forest autotrophic 112 carbon fluxes (FACF) vary with latitude. We then test how these fluxes relate to MAT and MAP, and addi-113 tionally how they respond to other, less well-studied, climate variables. Finally, we consider the relationship 114 between FACF and seasonality, considering the role of seasonality in explaining variation in carbon fluxes, 115 and the influence of climate on FACF standardized by growing season length. We use a comprehensive 116 global database of forest carbon fluxes to address the above questions for nine carbon fluxes, allowing for an in-depth exploration of the effect of climate on FACF globally.

Table 1: Summary of research questions, corresponding hypotheses, and results. Statistically significant support for/rejection of hypotheses is indicated with 'yes'/'no', parentheses indicate partial overall support/rejection of hypotheses across all FACF, and '-' indicates no significant relationship.

	Overall	Forest autotrophic carbon fluxes (FACF)									
Questions and hypotheses (with related references)		GPP	NPP	ANPP	$ANPP_{woody.stem}$	$ANPP_{foliage}$	BNPP	$BNPP_{fine.root}$	R_{auto}	$R_{auto-root}$	Support
Q1. How do FACF vary with latitude?											
H1.1. FACF decrease linearly with latitude. 1,2,3	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes	Fig. 2
Q2. How do FACF vary with MAT and MAP?											
H2.1. FACF increase with MAT. 1,4	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes	Figs. 4, S4, S5
$\mathrm{H}2.2.\ \mathrm{FACF}$ increase with precipitation. 1,4	(yes)	yes	yes	yes	-	yes	yes	yes	yes	yes	Figs. 4, S4, S5
H2.3. Temperature and precipitation interactively shape FACF. 5	(yes)	yes	yes	-	yes	-	yes	yes	yes	-	Fig. 3
Q3. How are FACF related to other climate variables?											
H3.1. FACF increase with PET.	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes	Figs. 4, S4, S5
H3.2. FACF increase with vapour pressure deficit.	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes	Figs. 4, S4, S5
H3.3. FACF increase with solar radiation.	(yes)	yes	yes	yes	yes	yes	yes	yes	yes	-	Figs. S4, S5
Q4. How does seasonality influence FACF?											
H4.1. FACF decrease with temperature seasonality.	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes	Figs. 4, S6, S7
H4.2. FACF decrease with precipitation seasonality.	-	-	-	-	-	-	-	-	-	-	Figs. S6, S7
H4.3. FACF increase with growing season length. 6,7,8	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes	Figs. 4, S6, S7
H4.4. Growing season length is a better predictor of FACF than MAT. $^{7.8}$	(no)	no	no	no	no	no	no	-	no	no	Table S4
Q5. When standardised by growing season length, how do FACF v	vary with	climat	te?								
H5.1. Growing season FACF increase with temperature. 8	(yes)	-	-	yes	-	yes	-	-	-	-	Figs. S8, S9
H5.2. Growing season FACF increase with PET.	(yes)	yes	yes	-	yes	-	yes	yes	-	-	Figs. S8, S9
H5.3. Growing season FACF increase with precipitation.	(yes)	-	-	yes	-	yes	-	-	-	-	Figs. S8, S9
H5.4. Growing season FACF increase with solar radiation.	(yes)	yes	yes	-	-	-	yes	yes	-	-	Figs. S8, S9

¹ Luyssaert et al. (2007) ² Gillman et al. (2015) ³ Simova and Storch (2017) ⁴ Schuur (2003) ⁵ Taylor et al. (2016) ⁶ Malhi (2012) ⁷ Michaletz et al. (2014) ⁸ Chu et al. (2016)

Materials and Methods

120 Forest carbon flux data

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This analysis focused on nine FACF included in the open-access ForC database (Table 2) (Anderson-Teixeira 121 et al., 2016, 2018). For C contains records of field-based measurements of forest carbon stocks and annual 122 fluxes, compiled from original publications and existing data compilations and databases. Associated data, 123 such as stand age, measurement methodologies, and disturbance history, are also included. The database 124 was significantly expanded since the publication of Anderson-Teixeira et al. (2018) through integration 125 with the Global Soil Respiration Database (Bond-Lamberty and Thomson, 2010). Additional targeted 126 literature searches were conducted to identify any further available data on the FACF analyzed here, with 127 particular focus on mature forests in temperate and boreal regions, which were not included in the review of 128 Anderson-Teixeira et al. (2016). We used ForC v3.0, archived on Zenodo with DOI 10.5281/zenodo.3403855. This version contained 29,730 records from 4,979 plots, representing 20 distinct ecozones across all forested 130 biogeographic and climate zones. 131

This analysis focused on mature forests with no known history of signficant disturbance or management. There is evidence that stand age influences patterns of FACF and carbon allocation in forest ecosystems, and can confound relationships between latitude and primary productivity (DeLUCIA et al., 2007; Gillman et al., 2015). To reduce any biasing effects of stand age, we included only stands of known age ≥ 100 years and those described by terms such as "mature", "intact", or "old-growth". Since management can alter observed patterns of FACF (Šímová and Storch, 2017), sites were excluded from analysis if they were managed, defined as plots that were planted, managed as plantations, irrigated, fertilised or including the term "managed" in their site description. Sites that had experienced significant disturbance within the past 100 years were also excluded. Disturbances that qualified sites for exclusion included major cutting or harvesting, burning, flooding, drought and storm events with site mortality >10% of trees. Grazed sites were retained.

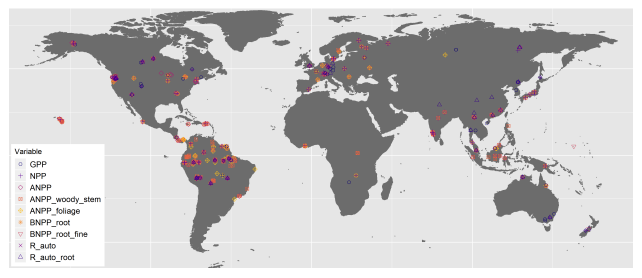


Figure 1: Map showing all data used in the analysis, coded by variable. Variables are plotted individually in Fig. S1.

Climate data

For C contains geographic coordinates associated with each measurement record and, when available, mean

Table 2: Definitions and sample sizes of FACF variables used in analysis. All variables are in units of Mg C $\rm ha^{-1}~\rm yr^{-1}$.

				Sample size		
Variable	Definition	Components included	Methodologies	records	geographi areas*	
GPP	Gross Primary Production	full ecosystem	flux partitioning of eddy-covariance; $NPP + R_{auto}$	243	49	
NPP	Net Primary Production	stem, foliage, coarse root, fine root, optionally others (e.g., branch, reproductive, understory)	$ANPP + BNPP$ (majority); $GPP\text{-}R_{auto}$	161	56	
ANPP	Above ground NPP	stem, foliage, optionally others (e.g., branch, reproductive, understory)	$ANPP_{woody-stem} + ANPP_{foliage}$ (+ others)	278	86	
$ANPP_{woody.stem}$	Woody stem growth component of $ANPP$	woody stems down to DBH \leq 10cm (no branch turnover)	stem growth measurements scaled to biomass using allometries $$	264	96	
$ANPP_{foliage}$	Foliage component of $ANPP$	foliage	litterfall collection (separated into components)	98	49	
BNPP	Below ground NPP	coarse and fine roots	coarse roots estimated indirectly using allometries based on above ground stem increment measures ; fine roots as below	101	48	
$BNPP_{fine.root}$	Fine root component of $BNPP$	fine roots	measurements combined one or more of the following: soil cores, minirhizotrons, turnover estimates, root ingrowth cores	88	41	
R_{auto}	Autotrophic respiration	foliage, stem, and root	chamber measurements of foliage and stem gas exchange + $R_{auto-root}$ (as below)	22	13	
$R_{auto-root}$	$u_{to-root}$ Root respiration (coarse and) fine roots		partitioning of total soil respiration (e.g., through root exclusion), scaling of root gas exchange; excluded alkali absoption and soda lime methods for measuring soil respiration	64	26	

^{*} Geographic areas group geographically proximate sites, defined using a hierarchical cluster analysis on the distance matrix of the sites, and a cutoff of 25km

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annual temperature (MAT) and mean annual precipitation (MAP) as reported in the primary literature (Anderson-Teixeira et al., 2018). Based on the geographic co-ordinates for each site, data on twelve climate variables—including MAT, MAP, temperature and precipitation seasonality, annual temperature range, solar radiation, cloud cover, annual frost and wet days, potential evapotranspiration (PET), aridity (MAP/PET), and vapor pressure deficit (VPD)-were extracted from five open-access climate datasets: WorldClim (Hijmans et al., 2005), WorldClim2 (Fick and Hijmans, 2017), the Climate Research Unit (CRU) time-series dataset v. 4.03 (Harris et al., 2014), the Global Aridity Index and Potential Evapotranspiration Climate Database (Trabucco and Zomer, 2019), and TerraClimate (Abatzoglou et al., 2018) (Table S1). From these data, we derived maximum VPD, defined as the VPD of the month with the largest deficit, and the number of water stress months, defined as the number of months annually where precipitation was lower than PET. Where site-level data was missing for MAT or MAP, we used values from the WorldClim dataset.

Length of the growing season was estimated to the nearest month, where growing season months were defined 155 as months with mean minimum temperature $> 0.5^{\circ}$ C. We experimented with a definion of growing season months including a moisture index, defined as (MAT - PET)/PET, > -0.95 (Kerkhoff et al. 2005; see also Michaletz et al. 2014). However, we found that including a moisture index had no effect on the estimates of growing season length, and so chose to exclude it. (Becky, was it really no effect? or minimal?) Monthly data for PET, precipitation, and temperature from the CRU dataset v 4.03 (Harris et al., 2014), and solar radiation from WorldClim2 (Fick and Hijmans, 2017) were used to calculate mean monthly PET, precipitation, temperature and solar radiation during the growing season. Total growing season precipitation

and solar radiation were also calculated.

164 Analyses

The effects of latitude and climate on FACF were analysed using mixed effects models using the package 165 'lme4' (Bates et al., 2015) in R v.3.5.1 (?). The basic model for all analyses included a fixed effect of latitude or 166 climate and a random effect of plot nested within geographic area. Geographic areas-i.e., spatially clustered 167 sites—are defined within ForC using a hierarchical cluster analysis on the distance matrix of the sites and a 168 cutoff of 25km (Anderson-Teixeira et al., 2018). We experimented with inclusion of altitude as a fixed effect, 169 but excluded it from the final models because it added very little explanatory power-that is, the difference 170 in AIC (Δ AIC) relative to models excluding altitude was generally small (often Δ AIC<2). Hypotheses were accepted if the Δ AIC between a model including the fixed effect of interest and a corresponding null model 172 excluding that fixed effect exceeded 2.0. All R^2 values presented here are marginal R^2 values, and refer to 173 the proportion of variation explained by only the fixed effects, unless otherwise specified. Becky, are there any instances of this? Specific analyses are as described below. 175

We first examined the relationship between latitude and FACF (Q1; Table 1). We tested models with latitude as a linear and term (corresponding null: model without latitude) and as a second-order polynomial term (corresponding null: model with latitude as a linear term). If the shape of the relationship **made biological** sense, and was a significant improvement on the linear relationship (Δ AIC >2), we accepted the polynomial as the best model. We also examined relationships among fluxes across latitude, testing whether sums of component fluxes matched the larger fluxes and whether C allocation varied with latitude, as specified below.

To test whether trends in component fluxes across latitude sum to match those of larger fluxes, regression lines for smaller component fluxes were summed to generate new estimates of larger fluxes, which were then compared against the latitudinal regression of the larger flux. Confidence intervals for the larger flux were calculated using the 'bootMer' function from the lme4 package (Bates et al., 2015). This analysis was applied to following sets of fluxes: (1) $GPP = NPP + R_{auto}$, (2) NPP = ANPP + BNPP, and (3) $ANPP = ANPP_{foliage} + ANPP_{woody.stem}$. In addition, we estimated total belowground C flux (TBCF, not analyzed due to limited data) as $TBCF = BNPP + R_{root}$.

Variation in allocation to component carbon fluxes along latitudinal gradients was explored for the following pairings: GPP:NPP, ANPP:BNPP, and $ANPP_{foliage}:ANPP_{woody.stem}; ANPP_{foliage}:NPP,$ $ANPP_{woody.stem}:NPP, ANPP:NPP,$ and BNPP:NPP. For each set of paired fluxes, measurements taken at the same site and plot, and in the same year, were paired together, and the ratio of each pair of measurements calculated. The ratios were regressed against latitude and climate variables, using the linear model specified above. Cook's distance analyses were carried out for each of the models, and indicated that data from a few high-elevation sites were having a disproportionate influence on the regressions. To account for this, models were re-run using only data from sites < 1000m.

We next examined the relationships of FACF to climate variables (Q2-Q4; Table 1). As with latitude, we tested both linear and polynomial fits for each climate variable. We tested relationships of each FACF (Table 2) against each climate variable (Table S1), but focus presentation—including focal hypotheses (Table 1)—on climate variables that explained >20% of variation in FACF. Becky, please make this more specific. I drafted this based on the results, but its not very clear.

To investigate the potential joint and interactive effects of climate variables on carbon fluxes, multivariate models were also specified. We explored a large range of climate variable combinations and ultimately

decided to focus on MAT and MAP. We favored these over variables with similar explanatory power because 204 they are the most commonly reported climate variables and are recorded in ForC. MAT was consistently among the best individual predictors of FACF (Table S2), although it was occasionally out-performed by 206 closely correlated climate variables (Fig. S2). In combination with MAT as a primary climate variable, MAP 207 was consistently among the best secondary climate variables. Specifically, compared again a model including MAT x MAP, there were no other variables that, in interaction with MAT, improved AIC by >2.0 and R^2 209 by >0.05. In examinging the joint and interactive effects of climate variables, an additive model including 210 MAT and MAP as fixed effects was accepted when $\Delta AIC > 2$ relative to a null including only MAT as a fixed effect. An interactive model including an MAT x MAP interaction was accepted when $\Delta AIC > 2$ relative to 212 a null including MAT and MAP as fixed effects. 213

To test whether and how FACF varied with climate when standardised by growing season length (Q5), we first standardized FACF by dividing by growing season length (as defined above). We then tested for correlations between growing season length-standardised FACF against growing season climate variables.

For analyses on data within the growing season, only linear models were specified.

All analyses were conducted in R (Version). Code and results are archived on GitHub....

219 Results

In total, we analyzed 1228 records from nine FACF variables taken from forests that had experienced no major anthropogenic disturbances within the past 100 years. These records represented a total of 154 distinct geographic areas (Fig. 1, Table 2), across all forested biogeographic and climate zones.

223 How do FACF vary with latitude?

All major carbon fluxes decreased linearly with latitude (Fig. 2; Table S2). Latitude was a strong predictor for many of the carbon fluxes-particularly the larger fluxes (Table S2). Specifically, latitude explained 64% of variation in GPP (n = 254, p<0.0001), 50% in NPP (n = 114, p<0.0001) and 45% in ANPP (n = 259, p<0.0001). The FACF that were most poorly predicted by latitude were $BNPP_{root-fine}$ (R^2 =0.17) and $ANPP_{woody.stem}$ =0.18). For all FACF, the relationship with latitude was best predicted by the linear model (but see this issue).

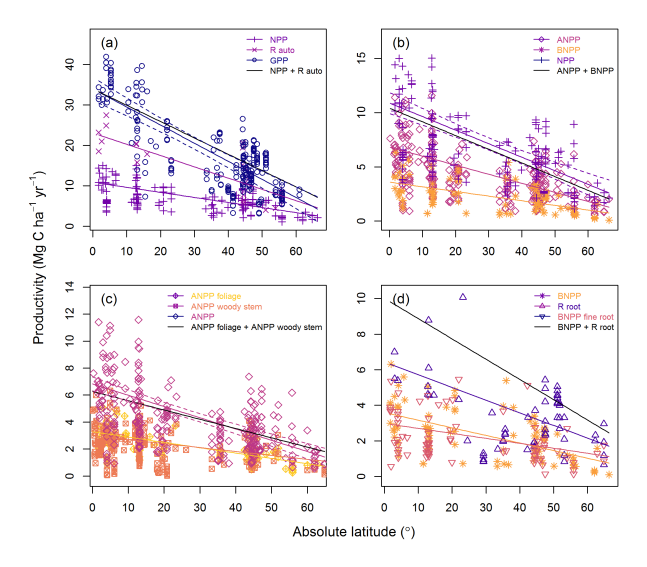


Figure 2: Latitudinal trends in forest autotropic carbon flux. Lines of best fit are plotted according to the best model selected during analysis. All regressions are significant (p < 0.05). Each panel shows major C fluxes together with component fluxes. Also plotted are predicted trends in the major C fluxes based on the sum of component fluxes. 95% confidence intervals are plotted for the major flux for comparison with predicted trends. In (d), which shows three belowground fluxes, the major flux, total belowground carbon flux, is one for which we have no data

In general, smaller component fluxes summed approximately to larger fluxes across the latitudinal gradient (Fig. 2). That is, modelled estimates of GPP, generated from the sum of NPP and R_{auto} ; NPP, generated from the sum of ANPP and BNPP; and ANPP, generated from the sum of $ANPP_{foliage}$ and $ANPP_{woody-stem}$, fell completely within the confidence intervals of the regressions of field estimates of GPP, NPP, and ANPP, respectively.

We found little evidence that allocation between fluxes varied substantially with latitude or climate (Fig. S3). Of the 7 FACF ratios regressed against latitude and three climate variables (MAT, MAP, temperature seasonality), there were only four signficant relationships, all with $R^2 \leq 0.2$ (Fig. S3). Specifically, the proportion of NPP allocated to $ANPP_{foliage}$ increased weakly with MAT ($R^2 = 0.20$), and the proportion of NPP allocated aboveground (ANPP) decreased weakly with latitude ($R^2 = 0.11$) and temperature

seasonality ($R^2 = 0.17$), while increasing with MAT ($R^2 = 0.11$). There were no significant results from regressing ratios of carbon fluxes against latitude, or against any of the climate variables.

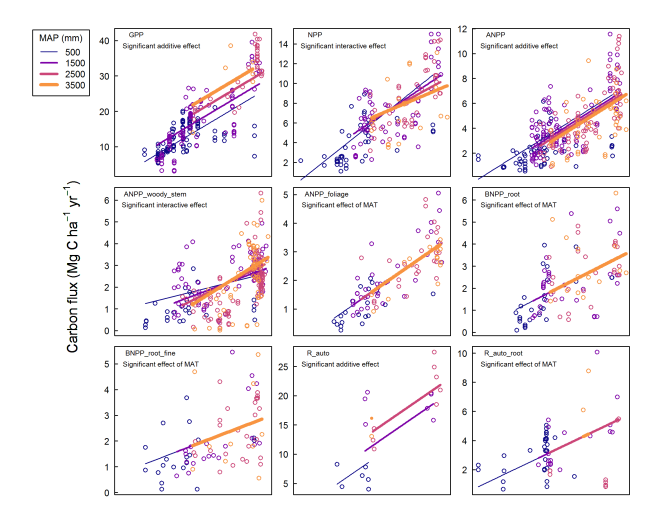
242 How do FACF relate to MAT and MAP?

All FACF increased linearly with MAT, and we found no support for a saturation point of FACF with MAT (all p<0.05; Figs. 3-4, S4-S5, Table S2). As with latitude, MAT tended to explain more variation in the larger FACF (GPP, NPP, ANPP, R_{auto}) and $ANPP_{foliage}$ (all $R^2 > 0.4$) than in subsidiary and belowground fluxes ($ANPP_{woody-stem}$, R_{root} , $BNPP_{root-fine}$; all $R^2 < 0.25$).

MAP was a significant (p<0.05) predictor of all FACF but $ANPP_{woody-stem}$ (Figs. 4a, S4-S5; Table S2). 247 However, it explained little variation: with the exception of R_{auto} , MAP explained at most 37% of variation 248 in FACF. For the majority of FACF, a polynomial model was the best fit. FACF generally increased with 249 precipitation, up until a saturation point at between 3000 and 4000mm annual precipitation, above which 250 they started to decrease (Figs. 4, S4-S5). The notable exception to this was GPP: the model indicated that 251 GPP continued to increase with precipitation up to measures of at least 5000mm annually (p<0.0001, R^2 252 0.33. Data above this point were not available, but the model trend indicated that the saturation point for 253 this model would be around 5000mm MAP. 254

There was a significant additive effect of MAT and MAP on GPP, ANPP and R_{auto} (Fig. 3, Table S3).

Accounting for MAT, MAP had a substantial positive effect on GPP and R_{auto} and a small negative effect on ANPP. There was a significant interactive effect between MAT and MAP for NPP and $ANPP_{woody-stem}$ (Fig. 3, Table S3). The interaction was negative for NPP and positive for $ANPP_{woody-stem}$. For $ANPP_{foliage}$, BNPP, $BNPP_{root-fine}$, and $R_{auto-root}$, MAP did not have a significant effect when accounting for MAT (Fig. 3, Table S3). For the variables which showed a significant interactive or additive effect between MAT and MAP, no other climate variable, in combination with MAT, significantly improved on that model. {need to confirm this given changes in MAT MAP results (or you could just drop the sentence.)}



Mean Annual Temperature (degrees)

Figure 3: Interactive effects of mean annual temperature and mean annual precipitation on FACF. For visualization purposes, data points are grouped into bins of 0 - 1000, 1001 - 2000, 2001 - 3000, and >3000mm mean annual precipitation, and lines of best fit models are plotted for mean annual precipitation values of 500, 1500, 2500, and 3500mm. All regressions are significant (p < 0.05).

264 How do FACF relate to other climate variables?

Our results indicated that FACF were most strongly explained by temperature at the global scale, with temperature-related climate variables coming out as strong predictors of FACF. In addition to MAT, several of its correlates (Fig. S2) were consistently identified as strong univariate predictors of FACF: temperature seasonality, annual temperature range, annual frost days, PET, and length of growing season (Figs. 4, S4-S7).

We found a significant relationship between C flux and potential evapotranspiration for all FACF. $ANPP_{foliage}$, $BNPP_{root-fine}$ and R_{root} increased linearly with PET; however, all other fluxes showed a polynomial relationship with PET (Fig. 4c, S4-5; Table S2). We found strong evidence for a saturation point or peak with PET: FACF tended to increase at values below 1000mm, before saturating between 1200

- 274 and 1700mm. There was also evidence that FACF begin to decrease at values above 1800mm PET.
- Vapour pressure deficit was a significant predictor of C flux for all FACF. $BNPP_{root-fine}$ showed a linear
- relationship with vapour pressure deficit ($R^2 = 0.07$, p<0.05), but all other fluxes showed a polynomial rela-
- tionship (Figs. 4d, S4-5; Table S2). FACF initially increased with vapour pressure deficit, before saturating
- $_{278}$ at around 0.8 kPa, after wich point they began to decrease.
- All fluxes, with the exception of R_{root} , showed a positive linear relationship with solar radiation (Figs. S4-S5,
- Table S2). Solar radiation explained a low proportion of variability in all FACF, explaining less than 20%
- of the variation in each flux, with the exception of R_{auto} ($R^2 = 0.26$, p<0.05).
- Annual wet days, cloud cover, and aridity were poor or non-significant explainers of variation in FACF,
- explaining less than 20% of the variation in each of the carbon fluxes (Figs. S4-S5; Table S2).

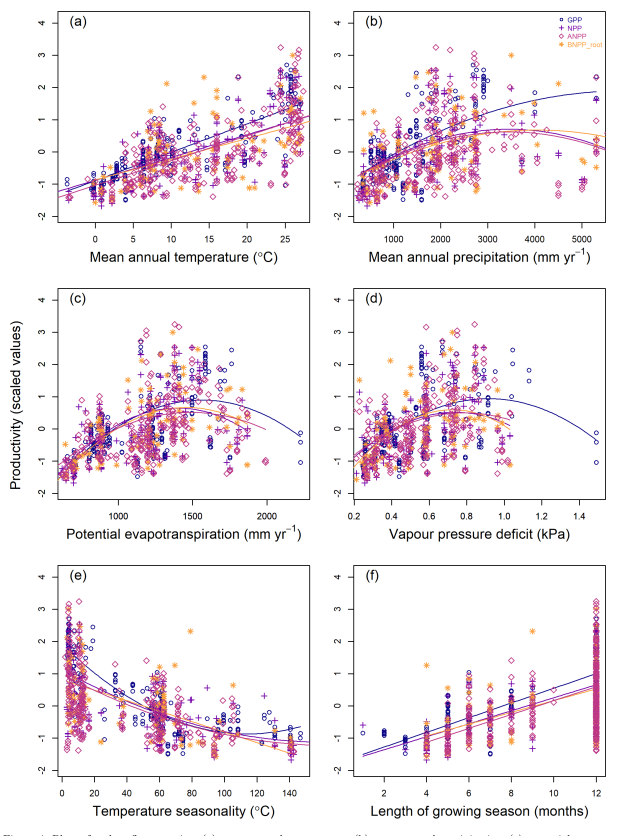


Figure 4: Plots of carbon fluxes against (a) mean annual temperature; (b) mean annual precipitation; (c) potential evapotranspiration, (d) vapour pressure deficit; (e) temperature seasonality; (f) length of growing season. For visualization purposes, data for each flux was rescaled with a mean of 0 and standard deviation of 1. Lines of best fit are plotted according to the best model selected during analysis (**see issue 47**). All regressions are significant (p < 0.05).

284 What is the role of seasonality in explaining FACF?

Temperature seasonality was a significant predictor of FACF. GPP, NPP, ANPP, and R_{root} exhibited a polynomial relationship with seasonality (all p<0.05; Figs. 4e, S6-7; Table S2). $ANPP_{foliage}$, $ANPP_{woody-stem}$ and R_{auto} decreased linearly with temperature seasonality (all p<0.05; Figs. 4e, S6-288 S7; Table S2). Temperature seasonality was strongly correlated with annual temperature range, which was likewise a similarly strong predictor of FACF (Table S2).FACF were highest where temperature seasonality = 0, and at an annual temperature range of 15°C or lower.

In contrast, there was no significant effect of precipitation seasonality on FACF, and both maximum vapour pressure deficit, and water stress months were poor or non-significant explainers of variation in FACF (Figs. S6-S7; Table S2).

We found a significant relationship between length of growing season and FACF, with all fluxes showing a linear increase with length of growing season (Figs. 4e, S6-S7; Table S2). Length of growing season was a strong predictor of FACF, explaining 51% of variation in GPP, 39% of variation in NPP, and 34% of variation in ANPP, but it was a weaker predictor than MAT for all fluxes analysed (Table S4).

298 Within the growing season, how do FACF vary with climate?

When FACF were standardized by growing season length, correlations with growing season climate—including 299 temperature, precipitation, solar radiation, and PET-were generally weak (Figs. S8-S9). Speficifally, ANPP 300 increased with growing season temperature ($R^2 = 0.10$, p<0.001) and precipitation ($R^2 = 0.04$, p<0.05). 301 Similarly, $ANPP_{foliage}$ increased slightly with growing season temperature ($R^2 = 0.16$, p<0.01) and pre-302 cipitation ($R^2 = 0.09$, p<0.05). Growing season solar radiation had a positive influence on GPP ($R^2 =$ 303 0.21, p<0.001), NPP ($R^2 = 0.21$, p<0.001), BNPP ($R^2 = 0.16$, p<0.001) and BNPP_{fine.root} ($R^2 = 0.12$, p<0.01). Growing season PET had a positive influence on GPP ($R^2 = 0.15$, p<0.01), NPP ($R^2 = 0.18$) 305 p<0.01), BNPP ($R^2=0.23$, p<0.0001), $BNPP_{fine.root}$ ($R^2=0.11$, p<0.05), and $ANPP_{woody-stem}$ ($R^2=0.11$), $R^2=0.11$, $R^2=0.11$, 306 = 0.06, p<0.05). {Becky, please verify/ edit the following: There were no other significant correlations between growing season length-standardized FACF (9 variables in Table 2) and growing season climate 308 (which variables?)}. 309

310 Discussion

Our analysis of a large global database (ForC) reveals how autotrophic carbon fluxes in mature forests vary 311 with latitude and climate on a global scale. We show that, across all nine FACF analyzed, C cycling decreases 312 continually with latitude (H1.1; Fig. 2)—a finding that confirms multiple previous studies but contradicts the 313 idea that productivity of temperate forests rivals that of tropical forests (Huston and Wolverton, 2009). The 314 FACF generally increase in proportion to one another (H1.2), with few differences in allocation detectable 315 at this global scale (Fig. S2) and with component fluxes summing appropriately to larger fluxes (Fig. 2). 316 indicating no major, systematic omissions or overestimations of flux components. However, climate explained 317 lower proportions of variability among subsidiary C fluxes (e.g., $ANPP_{woody}$, $BNPP_{fine.root}$, $R_{auto-root}$; 318 Fig. 2; Table S2). Latitudianal variation in FACF is primarily attributable to temperature-related variables 319 (H3, H4), particularly MAT (Figs. 3-4). Water availability is also influential, but generally of secondary 320 importance across the range represented in our database (Figs. 3-4). Temperature seasonality and growing 321 season length are closely correlated with MAT and are strong predictors of FACF (H4; Figs. 4e-f, S2, S6-S7), though growing season length doesn't improve upon MAT as a predictor. Within the growing season, the influence of climate on C cycling is smaller but still significant for a number of FACF (*H5*; Fig. S9; Table S4). These findings clarify the big picture of how FACF vary with latitude and climate on a global scale.

Past studies have differed in their conclusions regarding the relationship between FACF and latitude or its 326 correlates (Table 1, H1; REFS)—quite possibly because of lack of standardization with respect to stand 327 age and disturbance history. Our findings indicate that, among mature, undisturbed stands, FACF are unambiguously highest in the tropical regions, and the relationship is approximately linear (Fig. 2). This 320 contrasts with the suggestion that productivity of temperate forests is similar to that of tropical forests 330 (Huston and Wolverton, 2009). Compared to tropical forests, the temperate forest biome has experienced more widespread anthropogenic disturbance and has a larger fraction of secondary stands (Poulter et al. 2018-332 DOI:10.1594/PANGAEA.889943; Potapov et al. 2008. Mapping the World's Intact Forest Landscapes by 333 Remote Sensing. Ecology and Society 13 (2), 51.), so analyses comparing across latitudinal gradients without controlling for stand age risk confounding age with biome effects. In addition, because C allocation varies 335 with stand age (DeLUCIA et al., 2007) (See Nobby's comment in manuscript-draft_NK.pdf), age 336 differences may introduce systematic biases into analyses of FACF across latitude or global climatic gradients. 337 For example, woody productivity tends to be higher in rapidly aggrading secondary stands than in old-growth 338 forests, where proportionally more C is allocated to respiration (** Nobby AFM paper: Understanding 339 the controls over forest carbon use efficiency on small spatial scales: Effects of forest disturbance and tree diversity**) [*purpose for respiration/ other compenents (See Nobby's comment in manuscript-341 draft_NK.pdf)]. 342

This paragraph needs work-including re-assessment of results, interpretation, writing We show 343 that FACF are broadly consistent in their responses to climate drivers on the global scale (with the exception 344 of some differences in MAT-MAP interactions; Fig. 3), with no pronounced trends in C allocation among 345 the variable pairs tested (Figs. 2, S3). Although variation in allocation has been observed along gradients 346 of elevation (Moser et al., 2011) and water availability (Newman et al., 2006)—along with non-climatic axes 347 of stand age (Litton et al., 2007), nutrient availability (Litton et al., 2007; Gill and Finzi, 2016), and 348 forest structure (Taylor et al., 2019)-little variation in relation to climate is apparent at the global scale 349 within ForC, which contains the bulk of relevant data. Our conclusion, then, is that hypothesized gradients 350 in allocation along global climate gradients cannot currently be supported for mature forests, although 351 data quantity and standardization is currently insufficent to rule out the possibility that such trends exist. bbl: remove following sentence? Of particular interest and significance are the relationships amongst 353 GPP, net primary productivity (NPP and its components, particularly $ANPP_{woody-stem}$), and respiration 354 (R_{auto}) and components). There have been suggestions that tropical forests tend to have low carbon use 355 efficiency ($CUE = NPP/GPP = (GPP-R_{auto}/GPP)$, which are based on observations of low CUE in old-356 growth tropical forests relative to (mostly younger) extratropical forests (DeLUCIA et al., 2007; Malhi, 2012; 357 Anderson-Teixeira et al., 2016), but our analysis suggests that these low values might more appropriately be attributed to the fact that these forests are old than to their tropical climate. Indeed, CUE is known 359 to decline with forest age (DeLUCIA et al., 2007; Collalti and Prentice, 2019), but appears to be roughly 360 independent of GPP (Litton et al., 2007). Among our sites with relevant data, there is indication that 361 CUE or $ANPP_{woody-stem}/GPP$ increase with latitude (Fig. S3). Additional measurements with careful 362 methodological standardization across a consistent set of mature forest sites will be necessary to determine 363 whether any climate-driven gradients in allocation exist at the global scale.

One interesting observation was that climate tends to explain more variation in the major fluxes (GPP,NPP, R_{auto} - latitude $R^2 \geq 48\%$) than in subsidiary fluxes (latitude $R^2 < 27\%$ for $BNPP_{fine.root}$, $R_{auto-root}$, $ANPP_{woody-stem}$) (Fig. 2; Table S2). There are two, non-exclusive, potential explanations for 367 this. First, it may be that methodological variation is larger relative to flux magnitude for some of the 368 subsidiary fluxes. Belowground fluxes in particular are difficult to quantify, and measurement methods for the belowground fluxes considered here may be measured through fundamentally different approaches 370 (e.g., minirhizotrons, ingrowth cores, or sequential coring for $BNPP_{root-fine}$; root exclusion, stable isotope 371 tracking, or gas exchange of excised roots for $R_{auto-root}$), and sampling depth is variable and often 372 insufficient to capture the full soil profile. $ANPP_{woody-stem}$, which is also poorly explained by latitude or 373 climate, is more straightforward to measure but is subject to variability introduced by differences such as 374 biomass allometries applied and minimum plant size sampled (bbl: cite e.g. Huntzinger?). However, methodological variation and uncertainty affect all of fluxes considered here-not necessarily any less than 376 the aforementioned, and some of the larger fluxes that vary more strongly with respect to climate (ANPP,377 NPP) are estimated by summing uncertain component fluxes. Second, differences among variables in the 378 proportion of variation explained by climate may be attributable to more dicrect climatic control over GPP 379 than subsidiary fluxes. That is, subsidiary fluxes may be shaped by climate both indirectly through its 380 influence on GPP and respiration and directly through any climatic influence on C allocation, as well as 381 many other local- and regional-scale factors (**REFS**). 382

The latitudinal gradient in FACF (Fig. 2) is driven primarily by temperature-related climate variables, 383 and secondarily by moisture availability (Table 1, H2-H3; Figs. 3-4). Because many climate variables covary across the latitudinal gradient (Fig. S2), because climatic drivers affect forest carbon flux on much 385 shorter time scales than can be captured by annual climate summary variables, and because both climatic 386 conditions and C flux vary intra- and inter-annually around the long-term means, it is not appropriate to attempt to identify any one mean annual climate variable as a mechanistic driver of FACF. However, 388 it remains informative to consider these relationships. We find that temperature-related climate variables 389 $(MAT, \text{ temperature seasonality}, \dots \mathbf{LIST})$ explain the highest proportion of variability in FACF, and among these, MAT is generally the best predictor-perhaps because site-specific MAT is recorded for the majority 391 of sites in ForC, whereas other variables are extracted from global gridded data products (Table S1). The 392 effects of temperature are modified by moisture availability, with reduced FACF under hot and dry conditions 393 (i.e., high PET, high deficit; Fig. 4c-d) and sometimes under very high precipitation (Figs. 3, 4b). Negative 394 effects of very high precipitation on FACF have been observed previously (REFS) and are attributable 395 to nutrient and light limitations (REFS). Thus, although temperature and water availability jointly and interactively drive global-scale patterns of FACF. 397

FACF are negatively correlated with temperature seasonality (Table 1, H4; Fig. 4e), and is minimal during 398 cold- or dry- dormant seasons. To account for this, a number of analyses seeking to characterize global-399 scale effects of climate on productivity have examined the relationship of C flux per month of the growing 400 season with growing season climatic conditions (Table 1, H5; REFS). We found that the sort of simple 401 metric needed to define growing season at a global scale was uncertain for temperature and problematic for 402 moisture (WORK ON THIS). A temperature-defined growing season length had stong positive correlation 403 with FACF (Fig. 4f), but explained less variation than MAT. Dividing FACFs by growing season length to yield FACF per growing season month removed the majority of climate-related variation, supporting 405 the idea that the latitudinal gradient in FACF is attributable more to shorter growing seasons at high 406 latitudes than to inherently lower rates of photosynthesis or respiration by high-latitude forests (/ Enquist

et al. 2007 GCB- but check). However, there remained a number of significant correlations with growing 408 season climatic conditions, suggesting that climatic conditions remain influential within the growing season. We conclude that while correcting for growing season length takes analyses a step closer to mechanistic 410 linkage of instantaneous C flux rates to environmental conditions, it remains very crude relative to the the 411 timescales on which climate affects plant metabolism and does not advance statistical predictive power. Rather, mechanistic accounting for climatic effects on global FACF patterns requires models representing 413 physiologically meaningful timescales (e.q., refs). 414

Our analysis clarifies how FACF vary with latitude and climate on a global scale, with some important 415 implications for how forest carbon cycling relates to climate and, by extension, how it is likely to respond to climatic warming. Our findings show that higher temperatures with similar moisture availability result in a 417 generalized acceleration of FACF (Figs. 2-3). This is consistent with observations of continental- to global-418 scale increases in GPP (Li & Xiao 2019) and ANPPwoodustem (Brienen et al., 2015; Hubau et al. 2020), along with some C cycle components not considered here—tree mortality (Brienen et al., 2015; McDowell et 420 al. 2018), soil respiration (Bond Lamberty & Thompson 2010), and heterotrophic soil respiration (Bond-421 Lamberty et al. 2018). However, increasing C flux rates are not universal (e.g., Rutishauser et al., 2019, 422 MORE REFS). This is likely because factors other than rising temperatures are at play, including changes 423 to other aspects of cliamte, atmospheric pollution (CO₂, SO₂, NO_x), and local disturbances. Morevoer, 424 forest ecosystem responses to climatic changes outside the temperature range to which forest communities are adapated and acclimatized will not necessarily parallel responses across geographic gradients in 426 climate. Nevertheless, as we enter a period of accelerating climatic change, understanding of the fundamental 427 climatic controls on FACF sets a foundation for understanding patterns of change.

misc content for discussion

- the observed positive interaction between MAT and MAP for $ANPP_woody$ is consistent with the Taylor 430 et al. (2017) analysis showing such an interaction for ANPP in tropical forests. Similar to their analysis, 431 we find a cross-over point at ~ 20 C. However, we don't find such an interaction for ANPP, and we show 432 a contrasting negative interaction for NPP. Some of this is may be stochastic driven by composition 433 of the dataset, and the interactions we observe are not internally consistent.
- consistent with Hofhansl et al. 2015 (verify), we found a slight tendency for warmer sites to have 435 higher aboveground allocation 436
- -results are consistent with Muller-Landau et al., in review 437

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434

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References

- Abatzoglou, J. T., Dobrowski, S. Z., Parks, S. A., and Hegewisch, K. C. (2018). TerraClimate, a high-
- resolution global dataset of monthly climate and climatic water balance from 1958–2015. Scientific Data,
 5(1):170191.
- Anderson-Teixeira, K. J., Wang, M. M. H., McGarvey, J. C., Herrmann, V., Tepley, A. J., Bond-Lamberty,
- B., and LeBauer, D. S. (2018). For C: a global database of forest carbon stocks and fluxes. Ecology,
- 99(6):1507-1507.
- Anderson-Teixeira, K. J., Wang, M. M. H., McGarvey, J. C., and LeBauer, D. S. (2016). Carbon dynamics of
- mature and regrowth tropical forests derived from a pantropical database (TropForC-db). Global Change
- Biology, 22(5):1690–1709.
- 450 Aragão, L. E. O. C., Malhi, Y., Metcalfe, D. B., Silva-Espejo, J. E., Jimenez, E., Navarrete, D., Almeida,
- S., Costa, A. C. L., Salinas, N., Phillips, O. L., Anderson, L. O., Alvarez, C., Baker, T. R., Goncalvez,
- P. H., Huaman-Ovalle, J., Mamani, M., Meir, P., Monteagudo, A., Patino, S., Penuela, M. C., Prieto, A.,
- Quesada, C. A., Rozas-Davila, A., Rudas, A., Silva, J. A., and Vasquez, R. (2009). Above and belowground
- net primary productivity across Amazonian forests on contrasting soils. *Biogeosciences*, (6):2759–2778.
- Badgley, G., Anderegg, L. D. L., Berry, J. A., and Field, C. B. (2019). Terrestrial gross primary production:
- Using NIR v to scale from site to globe. Global Change Biology, 25(11):3731–3740.
- Bates, D., Mächler, M., Bolker, B., and Walker, S. (2015). Fitting Linear Mixed-Effects Models Using lme4.
- Journal of Statistical Software, 67(1).
- Beer, C., Reichstein, M., Tomelleri, E., Ciais, P., Jung, M., Carvalhais, N., Rodenbeck, C., Arain, M. A.,
- Baldocchi, D., Bonan, G. B., Bondeau, A., Cescatti, A., Lasslop, G., Lindroth, A., Lomas, M., Luyssaert,
- S., Margolis, H., Oleson, K. W., Roupsard, O., Veenendaal, E., Viovy, N., Williams, C., Woodward, F. I.,
- and Papale, D. (2010). Terrestrial Gross Carbon Dioxide Uptake: Global Distribution and Covariation
- with Climate. Science, 329(5993):834–838.
- Bonan, G. B. (2008). Forests and Climate Change: Forcings, Feedbacks, and the Climate Benefits of Forests.
- Science, 320(5882):1444-1449.
- 466 Bond-Lamberty, B. and Thomson, A. (2010). A global database of soil respiration data. Biogeosciences,
- 7(6):1915–1926.
- ⁴⁶⁸ Chu, C., Bartlett, M., Wang, Y., He, F., Weiner, J., Chave, J., and Sack, L. (2016). Does climate directly
- influence NPP globally? Global Change Biology, 22(1):12–24.
- Cleveland, C. C., Townsend, A. R., Taylor, P., Alvarez-Clare, S., Bustamante, M. M. C., Chuyong, G.,
- Dobrowski, S. Z., Grierson, P., Harms, K. E., Houlton, B. Z., Marklein, A., Parton, W., Porder, S., Reed,
- S. C., Sierra, C. A., Silver, W. L., Tanner, E. V. J., and Wieder, W. R. (2011). Relationships among
- net primary productivity, nutrients and climate in tropical rain forest: a pan-tropical analysis: Nutrients,
- climate and tropical NPP. Ecology Letters, 14(9):939–947.
- 475 Collalti, A. and Prentice, I. C. (2019). Is NPP proportional to GPP? Waring's hypothesis 20 years on. Tree
- Physiology, 39(8):1473–1483.

- DeLUCIA, E. H., Drake, J. E., Thomas, R. B., and Gonzalez-Meler, M. (2007). Forest carbon use efficiency: is respiration a constant fraction of gross primary production? *Global Change Biology*, 13(6):1157–1167.
- Fick, S. E. and Hijmans, R. J. (2017). WorldClim 2: new 1-km spatial resolution climate surfaces for
 global land areas: NEW CLIMATE SURFACES FOR GLOBAL LAND AREAS. International Journal
 of Climatology, 37(12):4302-4315.
- 482 Fyllas, N. M., Bentley, L. P., Shenkin, A., Asner, G. P., Atkin, O. K., Díaz, S., Enquist, B. J., Farfan-Rios,
- W., Gloor, E., Guerrieri, R., Huasco, W. H., Ishida, Y., Martin, R. E., Meir, P., Phillips, O., Salinas,
- N., Silman, M., Weerasinghe, L. K., Zaragoza-Castells, J., and Malhi, Y. (2017). Solar radiation and
- functional traits explain the decline of forest primary productivity along a tropical elevation gradient.
- Ecology Letters, 20(6):730-740.
- Gill, A. L. and Finzi, A. C. (2016). Belowground carbon flux links biogeochemical cycles and resource-use efficiency at the global scale. *Ecology Letters*, 19(12):1419–1428.
- Gillman, L. N., Wright, S. D., Cusens, J., McBride, P. D., Malhi, Y., and Whittaker, R. J. (2015). Latitude,
 productivity and species richness: Latitude and productivity. Global Ecology and Biogeography, 24(1):107–
 117.
- Girardin, C. A. J., Malhi, Y., Aragão, L. E. O. C., Mamani, M., Huaraca Huasco, W., Durand, L., Feeley,
- 493 K. J., Rapp, J., Silva-Espejo, J. E., Silman, M., Salinas, N., and Whittaker, R. J. (2010). Net primary
- productivity allocation and cycling of carbon along a tropical forest elevational transect in the Peruvian
- Andes: NET PRIMARY PRODUCTIVITY FROM ANDES TO AMAZON. Global Change Biology. 16(12):3176–3192.
- Harris, I., Jones, P., Osborn, T., and Lister, D. (2014). Updated high-resolution grids of monthly climatic
- $_{\rm 498}$ $\,$ observations the CRU TS3.10 Dataset: UPDATED HIGH-RESOLUTION GRIDS OF MONTHLY CLI-
- MATIC OBSERVATIONS. International Journal of Climatology, 34(3):623–642.
- Hijmans, R. J., Cameron, S. E., Parra, J. L., Jones, P. G., and Jarvis, A. (2005). Very high resolution
 interpolated climate surfaces for global land areas. *International Journal of Climatology*, 25(15):1965–1978.
- Huston, M. A. and Wolverton, S. (2009). The global distribution of net primary production: resolving the paradox. *Ecological Monographs*, 79(3):343–377.
- Jung, M., Reichstein, M., Margolis, H. A., Cescatti, A., Richardson, A. D., Arain, M. A., Arneth, A.,
- Bernhofer, C., Bonal, D., Chen, J., Gianelle, D., Gobron, N., Kiely, G., Kutsch, W., Lasslop, G., Law,
- B. E., Lindroth, A., Merbold, L., Montagnani, L., Moors, E. J., Papale, D., Sottocornola, M., Vaccari,
- 508 F., and Williams, C. (2011). Global patterns of land-atmosphere fluxes of carbon dioxide, latent heat,
- and sensible heat derived from eddy covariance, satellite, and meteorological observations. Journal of
- Geophysical Research, 116:G00J07.
- Kerkhoff, A. J., Enquist, B. J., Elser, J. J., and Fagan, W. F. (2005). Plant allometry, stoichiometry and the
- ${}_{512} \hspace{0.5cm} \text{temperature-dependence of primary productivity: Plant allometry, stoichiometry and productivity.} \hspace{0.5cm} \hspace{0.5cm} \textit{Global}$
- Ecology and Biogeography, 14(6):585–598.

- Larjavaara, M. and Muller-Landau, H. C. (2012). Temperature explains global variation in biomass among humid old-growth forests: Temperature and old-growth forest biomass. *Global Ecology and Biogeography*, 21(10):998–1006.
- Liang, J., Crowther, T. W., Picard, N., Wiser, S., Zhou, M., Alberti, G., Schulze, E.-D., McGuire, A. D.,
- Bozzato, F., Pretzsch, H., de Miguel, S., Paquette, A., Herault, B., Scherer-Lorenzen, M., Barrett, C. B.,
- Glick, H. B., Hengeveld, G. M., Nabuurs, G.-J., Pfautsch, S., Viana, H., Vibrans, A. C., Ammer, C.,
- Schall, P., Verbyla, D., Tchebakova, N., Fischer, M., Watson, J. V., Chen, H. Y. H., Lei, X., Schelhaas,
- M.-J., Lu, H., Gianelle, D., Parfenova, E. I., Salas, C., Lee, E., Lee, B., Kim, H. S., Bruelheide, H.,
- ⁵²² Coomes, D. A., Piotto, D., Sunderland, T., Schmid, B., Gourlet-Fleury, S., Sonke, B., Tavani, R., Zhu, J.,
- Brandl, S., Vayreda, J., Kitahara, F., Searle, E. B., Neldner, V. J., Ngugi, M. R., Baraloto, C., Frizzera,
- L., Ba azy, R., Oleksyn, J., Zawi a-Nied wiecki, T., Bouriaud, O., Bussotti, F., Finer, L., Jaroszewicz,
- B., Jucker, T., Valladares, F., Jagodzinski, A. M., Peri, P. L., Gonmadje, C., Marthy, W., OBrien, T.,
- Martin, E. H., Marshall, A. R., Rovero, F., Bitariho, R., Niklaus, P. A., Alvarez-Loayza, P., Chamuya,
- N., Valencia, R., Mortier, F., Wortel, V., Engone-Obiang, N. L., Ferreira, L. V., Odeke, D. E., Vasquez,
- R. M., Lewis, S. L., and Reich, P. B. (2016). Positive biodiversity-productivity relationship predominant
- in global forests. *Science*, 354(6309):aaf8957–aaf8957.
- Litton, C. M., Raich, J. W., and Ryan, M. G. (2007). Carbon allocation in forest ecosystems. Global Change
 Biology, 13(10):2089–2109.
- Luyssaert, S., Inglima, I., Jung, M., Richardson, A. D., Reichstein, M., Papale, D., Piao, S. L., Schulze,
- E. D., Wingate, L., Matteucci, G., Aragao, L., Aubinet, M., Beer, C., Bernhofer, C., Black, K. G.,
- Bonal, D., Bonnefond, J. M., Chambers, J., Ciais, P., Cook, B., Davis, K. J., Dolman, A. J., Gielen, B.,
- Goulden, M., Grace, J., Granier, A., Grelle, A., Griffis, T., Grünwald, T., Guidolotti, G., Hanson, P. J.,
- Harding, R., Hollinger, D. Y., Hutyra, L. R., Kolari, P., Kruijt, B., Kutsch, W., Lagergren, F., Laurila,
- T., Law, B. E., Le Maire, G., Lindroth, A., Loustau, D., Malhi, Y., Mateus, J., Migliavacca, M., Misson,
- L., Montagnani, L., Moncrieff, J., Moors, E., Munger, J. W., Nikinmaa, E., Ollinger, S. V., Pita, G.,
- Rebmann, C., Roupsard, O., Saigusa, N., Sanz, M. J., Seufert, G., Sierra, C., Smith, M. L., Tang, J.,
- Valentini, R., Vesala, T., and Janssens, I. A. (2007). CO ₂ balance of boreal, temperate, and tropical
- forests derived from a global database. Global Change Biology, 13(12):2509–2537.
- Malhi, Y. (2012). The productivity, metabolism and carbon cycle of tropical forest vegetation: Carbon cycle of tropical forests. *Journal of Ecology*, 100(1):65–75.
- Malhi, Y., Girardin, C. A. J., Goldsmith, G. R., Doughty, C. E., Salinas, N., Metcalfe, D. B., Huaraca Huasco,
- W., Silva-Espejo, J. E., del Aguilla-Pasquell, J., Farfán Amézquita, F., Aragão, L. E. O. C., Guerrieri,
- R., Ishida, F. Y., Bahar, N. H. A., Farfan-Rios, W., Phillips, O. L., Meir, P., and Silman, M. (2017). The
- variation of productivity and its allocation along a tropical elevation gradient: a whole carbon budget
- perspective. New Phytologist, 214(3):1019–1032.
- Michaletz, S. T., Cheng, D., Kerkhoff, A. J., and Enquist, B. J. (2014). Convergence of terrestrial plant production across global climate gradients. *Nature*, 512(7512):39–43.
- Michaletz, S. T., Kerkhoff, A. J., and Enquist, B. J. (2018). Drivers of terrestrial plant production across
 broad geographical gradients. Global Ecology and Biogeography, 27(2):166–174.

- Moser, G., Leuschner, C., Hertel, D., Graefe, S., Soethe, N., and Iost, S. (2011). Elevation effects on the carbon budget of tropical mountain forests (S Ecuador): the role of the belowground compartment: ELEVATION EFFECTS ON FOREST CARBON CYCLING. Global Change Biology, 17(6):2211–2226. 555
- Newman, G. S., Arthur, M. A., and Muller, R. N. (2006). Above- and Belowground Net Primary Production 556 in a Temperate Mixed Deciduous Forest. Ecosystems, 9(3):317–329.
- Pan, Y., Birdsey, R. A., Fang, J., Houghton, R., Kauppi, P. E., Kurz, W. A., Phillips, O. L., Shvidenko, A., 558
- Lewis, S. L., Canadell, J. G., Ciais, P., Jackson, R. B., Pacala, S. W., McGuire, A. D., Piao, S., Rautiainen, 559
- A., Sitch, S., and Hayes, D. (2011). A Large and Persistent Carbon Sink in the World's Forests. Science,
- 333(6045):988-993. 561

557

- Piao, S., Luyssaert, S., Ciais, P., Janssens, I. A., Chen, A., Cao, C., Fang, J., Friedlingstein, P., Luo, Y., and 562 Wang, S. (2010). Forest annual carbon cost: a global-scale analysis of autotrophic respiration. Ecology, 563 91(3):652-661. 564
- Schuur, E. A. G. (2003). PRODUCTIVITY AND GLOBAL CLIMATE REVISITED: THE SENSITIVITY 565 OF TROPICAL FOREST GROWTH TO PRECIPITATION. Ecology, 84(5):1165–1170.
- Taylor, P. G., Cleveland, C. C., Soper, F., Wieder, W. R., Dobrowski, S. Z., Doughty, C. E., and Townsend, 567 A. R. (2019). Greater stem growth, woody allocation, and aboveground biomass in Paleotropical forests 568 than in Neotropical forests. Ecology, 100(3):e02589.
- Taylor, P. G., Cleveland, C. C., Wieder, W. R., Sullivan, B. W., Doughty, C. E., Dobrowski, S. Z., and 570 Townsend, A. R. (2017). Temperature and rainfall interact to control carbon cycling in tropical forests. 571 Ecology Letters, 20(6):779-788. 572
- Trabucco, A. and Zomer, R. J. (2019). Global Aridity Index and Potential Evapo-Transpiration (ET0) 573 Climate Database v2. page 10. 574
- Waide, R. B., Willig, M. R., Steiner, C. F., Mittelbach, G., Gough, L., Dodson, S. I., Juday, G. P., and Parmenter, R. (1999). The Relationship Between Productivity and Species Richness. Annual Review of 576 Ecology and Systematics, 30(1):257-300. 577
- Šímová, I. and Storch, D. (2017). The enigma of terrestrial primary productivity: measurements, models. scales and the diversity-productivity relationship. *Ecography*, 40(2):239–252. 579