- Title: Global patterns of forest autotrophic carbon fluxes
- Running head: Global patterns of forest carbon fluxes
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22 Abstract

Carbon (C) fixation, allocation, and metabolism by trees set the basis for energy and material flows in forest 23 ecosystems and define their interactions with Earth's changing climate. However, while many studies have considered variation in productivity with latitude and climate, we lack a cohesive synthesis on how forest carbon fluxes vary globally with respect to climate and one another. Here, we draw upon 1,319 records from the Global Forest Carbon Database (ForC), representing all major forest types and the nine most 27 significant autotrophic carbon fluxes, to comprehensively review how annual C cycling in mature, undisturbed forests varies with latitude and climate on a global scale. We show that, across all flux variables analyzed, C cycling decreases continuously with absolute latitude – a finding that confirms multiple previous studies and contradicts the idea that net primary productivity of temperate forests rivals that of tropical forests. C flux 31 variables generally displayed similar trends across latitude and multiple climate variables, with no differences 32 in allocation detected at this global scale. Temperature variables in general, and mean annual temperature or temperature seasonality in particular, were the single best predictors of C flux, explaining 19 - 71% of 34 variation in the C fluxes analyzed. The effects of temperature were modified by moisture availability, with C flux reduced under hot and dry conditions and sometimes under very high precipitation. Annual C fluxes increased with growing season length and were also influenced by growing season climate. These findings 37 clarify how forest C flux varies with latitude and climate on a global scale. In an era when forests will play a critical yet uncertain role in shaping Earth's rapidly changing climate, our synthesis provides a foundation for understanding global patterns in forest C cycling.

41 **Keywords:** carbon fluxes; carbon dioxide (CO₂); climate; forest; global; productivity; respiration; latitude

42 Introduction

Carbon (C) cycling in Earth's forests provides the energetic basis for sustaining the majority of Earth's terrestrial biodiversity and many human populations (Millennium Ecosystem Assessment, 2005), while strongly influencing atmospheric carbon dioxide (CO₂) and climate (Bonan, 2008). Forests' autotrophic C fluxes – that is, C 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 (Niedziałkowska et al., 2010; Zak et al., 1994) and possibly diversity (Chu et al., 2018; Waide et al., 1999). They are linked to cycling of energy, water, and nutrients and, critically, influence all C stocks and define forest interactions with Earth's changing climate. Each year, over 69 Gt of C cycle through Earth's forests (Badgley et al., 2019) – a flux more than seven times greater than that of recent anthropogenic fossil fuel emissions (9.5 Gt C yr⁻¹; Friedlingstein et al., 2019). As atmospheric CO₂ continues to rise, driving climate 52 change, forests will play a critical role in shaping the future of Earth's climate (Cavaleri et al., 2015; Rogel) et al., 2018). However, our understanding of global-scale variation in forest C cycling remains incomplete, in large part because it is pieced together from numerous studies, most considering only one or a few variables at a time, with various approaches for handling influential factors such as stand age, disturbance history, and management status (Gillman et al., 2015; Litton et al., 2007; Šímová & Storch, 2017). Forest C fluxes decrease with latitude (e.g., Luyssaert et al., 2007; Gillman et al., 2015; Li & Xiao, 2019). However, studies have differed in their conclusions regarding the shape of this relationship – quite possibly because of lack of standardization with respect to methodology and stand history. C flux and allocation vary with stand age, disturbance, and management (DeLucia et al., 2007; Fernandez-Martinez et al., 2014; Símová & Storch, 2017; Yu et al., 2014), making clear latitudinal patterns difficult to discern without standardization 62 of the dataset. Studies agree that gross primary productivity (GPP) increases continuously with decreasing latitude and is indisputably highest in tropical forests (Badgley et al., 2019; Beer et al., 2010; Jung et al., 2011; Li & Xiao, 2019; Luyssaert et al., 2007). However, this relationship is more ambiguous for subsidiary fluxes. Some studies have suggested that net primary productivity (NPP), or its aboveground portion (ANPP), exhibits a less distinct increase from temperate to tropical forests (Luyssaert et al., 2007) – or even a decrease (Huston & Wolverton, 2009, but see Gillman et al., 2015). A shallower increase in NPP than in GPP with decreasing latitude would align with the suggestion that tropical forests tend to have low carbon use efficiency (CUE= NPP/GPP; DeLucia et al., 2007; Anderson-Teixeira et al., 2016; Malhi, 2012) but contrast with recent findings of the opposite pattern (Collalti et al., 2020). Such differences among C fluxes in their relationship to latitude have profound implications for our understanding of the C cycle and its climate sensitivity (e.g., Collalti et al., 2020). However, until recently the potential to compare latitudinal

trends across C fluxes has been limited by lack of a large database with standardization for methodology, stand history, and management (Anderson-Teixeira et al., 2018).

Latitudinal gradients in forest C flux rates, along with altitudinal gradients (Girardin et al., 2010; Malhi et al., 2017; Muller-Landau et al., 2020), are driven primarily by climate, which is a significant driver of C fluxes across broad spatial scales (Cleveland et al., 2011; Luyssaert et al., 2007; Muller-Landau et al., 2020; Wei et al., 2010). However, there is little consensus as to the shapes of these relationships or the best predictor 79 variables. The majority of studies have focused on exploring the relationships of C fluxes to mean annual temperature (MAT) and precipitation (MAP), which are the most commonly reported site-level climate variables. C fluxes increase strongly with MAT on the global scale, but whether they saturate or potentially decrease at higher temperatures remains disputed. Some studies have detected no deceleration or decline in GPP (Luyssaert et al., 2007), NPP (Schuur, 2003), or root respiration (R_{root}; Piao et al., 2010; Wei et al., 2010) with increasing MAT. In contrast, others have found evidence of saturation or decline of C flux in the warmest climates: Luyssaert et al. (2007) found NPP saturating at around 10°C MAT; Larjavaara & Muller-Landau (2012) found that increases in GPP saturate at approximately 25°C MAT, and Sullivan et al. (2020) found that, within the tropics, woody stem productivity $(ANPP_{stem})$ decreases at the highest maximum temperatures. C fluxes generally saturate at high levels of MAP, though the saturation points identified vary widely (e.g., ~1000 - 2,445 mm yr⁻¹; Wei et al., 2010; Schuur, 2003). Interactions between MAT and MAP may also influence productivity (Yu et al., 2014); within the tropics, there is a positive interaction between MAT and MAP in shaping ANPP, such that temperature has a positive effect on productivity in moist climates, but a negative effect in dry climates (Taylor et al., 2017). There is also evidence that C fluxes also respond to climate variables such as seasonality of temperature and precipitation (Wagner et al., 2016), cloud cover (Taylor et al., 2017), solar radiation (Beer et al., 2010; Fyllas et al., 2017), and potential evapotranspiration (Kerkhoff et al., 2005); however, these are not typically assessed in global-scale analyses of annual forest C flux.

Mean annual temperature and precipitation do not capture intra-annual climate variation, including temperature and precipitation seasonality and growing season length. Most forests—even tropical evergreen—exhibit some seasonality in both climate and C flux (e.g., Wagner et al., 2014), and this seasonality influences annual C fluxes (Churkina et al., 2005; Fu et al., 2019; Keenan et al., 2014). In particular, growing season length has been linked to ANPP, NPP, GPP, and net ecosystem exchange of CO₂ (NEE, or the difference between GPP and ecosystem respiration; Kerkhoff et al., 2005; Churkina et al., 2005; Keenan et al., 2014; Michaletz et al., 2014; Zhou et al., 2016). However, the relative importance of growing season length, as opposed to climate within the growing season, remains debated. On one end of the spectrum, some studies have suggested that

the influence of temperature on C fluxes may be limited to determining the length of the frost-free growing 106 season, and that climate within the growing season has little influence on C fluxes because of plant adaptation and acclimatization to local climates (Enquist et al., 2007; Kerkhoff et al., 2005; Michaletz et al., 2018, 2014). 108 In support of this, Kerkhoff et al. (2005) and Michaletz et al. (2014) found no significant relationship between growing season temperature and ANPP or NPP standardized to a climate-defined growing season length 110 (but see Chu et al., 2016). The idea that growing season length is an important determinant of annual C flux also aligns with evidence that cross-site variation in NEE is strongly correlated with growing season 112 length (Churkina et al., 2005) and that warming-induced increases in growing season length are enhancing 113 forest GPP and C sequestration (Keenan et al., 2014; Zhou et al., 2016). On the other end of the spectrum, 114 climatic conditions within the growing season may exert a stronger influence on annual C fluxes than the 115 length of the growing season. This aligns with observations that in forests, NEE tends to be more closely tied to the maximum rate of CO₂ uptake than to the carbon uptake period (Fu et al., 2019; Zhou et al., 117 2016), and with numerous tree-ring analyses finding that annual growth is more closely controlled by peak growing season climate than by spring or fall conditions (e.g., Helcoski et al., 2019) REFS. Thus, the extent 119 to which growing season length controls global-scale variation in forest autotrophic C fluxes remains unclear. 120 The recent development of the Global Forest Carbon database (ForC), which synthesizes multiple variables 121 and includes records of stand history (Anderson-Teixeira et al., 2016, 2018), opens up the possibility for a 122 standardized analysis of global scale variation in multiple C fluxes and the principle climatic drivers of these patterns. In order to approach this broad topic, we organize the major gaps in our knowledge under five 124 broad review questions and corresponding predictions, many derived from the findings of previous studies (Table 1). First, we ask how nine forest autotrophic carbon fluxes in ForC vary with latitude (Q1). We 126 then test how these fluxes relate to MAT and MAP (Q2), and additionally how they respond to other, less 127 well-studied, climate variables (Q3). Finally, we consider the relationship between C flux and seasonality, 128 considering the role of seasonality in explaining variation in carbon fluxes (Q4), and the influence of climate 129 on C flux standardized by growing season length (Q5). Our analyses represent a major step forward in relation to previous work (e.g., Luyssaert et al., 2007) in that we examine global climatic trends in more 131 variables (9 vs. \leq 3), draw from a much larger database (>4 times more records analyzed), and control for the effects of stand age, disturbance, and management. 133

Table 1: Summary of review questions, corresponding expectations based on previous studies (when applicable), and results. Statistically significant support for/ rejection of hypotheses is indicated by checkmarks/ X's, and '-' indicates no significant relationship. Parentheses indicate partial overall support or rejection of hypotheses across all fluxes considered. Flux variables are defined in Table 2.

	Overall	Forest autotrophic carbon fluxes									
Review questions and hypothesized relationships		GPP	NPP	ANPP	$ANPP_{stem}$	$ANPP_{foliage}$	BNPP	$BNPP_{fine.root}$	R_{auto}	R_{root}	Support
Q1. How do C fluxes vary with latitude?											
continuous increase with decreasing latitude 1,2,3	\checkmark	✓	✓	✓	✓	✓	✓	✓	\checkmark	✓	Fig. 2
sign ficantly decelerating increase with decreasing latitude ^1,4 $$	×	×	×	×	×	×	×	×	×	×	Fig. 2
Q2. How do C fluxes vary with mean annual temperatu	re (MAT) and	precipit	tation (M	IAP)?						
continuous increase with MAT 1,5,6,7	\checkmark	\checkmark	✓	✓	✓	✓	✓	✓	\checkmark	✓	Figs. 3, 4, S4, S
increase with MAP up to \geq 2000 $\mathrm{mm}^{1,4,7}$	\checkmark	\checkmark	✓	\checkmark	✓	\checkmark	\checkmark	\checkmark	\checkmark	✓	Figs. 4, S4, S5
increase with MAT \times MAP ^8,9	-	-	×	-	✓	-	-	-	-	-	Fig. 3, Table S3
Q3. How are C fluxes related to other annual climate va	ariables?										
decelerating increase or unimodal relationship with PET	✓	\checkmark	✓	\checkmark	✓	✓	\checkmark	✓	✓	✓	Figs. 4, S4, S5
decelerating increase or unimodal relationship with $\rm VPD^{10}$	\checkmark	\checkmark	✓	\checkmark	✓	\checkmark	\checkmark	\checkmark	\checkmark	✓	Figs. 4, S4, S5
increase with solar radiation 11,12	(\checkmark)	✓	✓	✓	✓	\checkmark	✓	✓	✓	-	Figs. S4, S5
Q4. How does seasonality influence annual C fluxes?											
decrease with temperature seasonality	✓	\checkmark	✓	\checkmark	✓	✓	\checkmark	✓	✓	✓	Figs. 4, S6, S7
decrease with precipitation seasonality 13,14	-	-	-	-	×	-	-	-	-	-	Figs. S6, S7
increase with growing season length 15,16,17,18	✓	✓	✓	✓	✓	✓	\checkmark	✓	✓	✓	Figs. 4, S6, S7
stronger relationship to growing season length than $\mathrm{MAT}^{16,17}$	(×)	×	×	×	-	×	×	×	×	×	Table S4
Q5. When standardised by growing season length, how	do annua	d C flu	xes var	y with c	limate?						
increase with growing season temperature 17	(\checkmark)	-	-	✓	-	\checkmark	-	-	-	-	Figs. S8, S9
increase with growing season PET	(\checkmark)	✓	✓	-	✓	-	\checkmark	\checkmark	-		Figs. S8, S9
increase with growing season precipitation 18	(\checkmark)	-	-	\checkmark	-	✓	-	-	-	-	Figs. S8, S9
increase with growing season solar radiation	(√)	-	-	-	-	-	✓	✓	_	-	Figs. S8, S9

 $[\]begin{tabular}{ll} 1 Luyssaert et al. (2007) & 2 Gillman et al. (2015) & 3 Simova and Storch (2017) & 4 Huston & Wolverton (2009) & 5 Schuur (2003) & 6 Piao et al. (2010) & 7 Wei et al. (2010) & 8 Taylor et al. (2017) & 9 Muller-Landau et al. (2020) & 10 Smith et al. (2020) & 11 Fyllas et al. (2017) & 12 Nemani et al. (2003) & 13 Wagner et al. (2014) & 14 Wagner et al. (2016) & 15 Malhi (2012) & 16 Michaeltz et al. (2014) & 17 Chu et al. (2016) & 18 Fernandez-Martinez et al. (2014) &$

Materials and Methods

135 Forest carbon flux data

This analysis focused on nine C flux variables included in the open-access ForC database (Table 2; Anderson-136 Teixeira et al., 2016, 2018). For contains records of field-based measurements of forest carbon stocks and 137 annual fluxes, compiled from original publications and existing data compilations and databases. Associated 138 data, such as stand age, measurement methodologies, and disturbance history, are also included. The database was significantly expanded since the publication of Anderson-Teixeira et al. (2018) through integration with 140 the Global Soil Respiration Database (Bond-Lamberty & Thomson, 2010). Additional targeted literature searches were conducted to identify further available data on the fluxes analyzed here, with particular focus on 142 mature forests in temperate and boreal regions, which were not included in the review of Anderson-Teixeira et al. (2016). We used ForC v3.0, archived on Zenodo with DOI 10.5281/zenodo.3403855. This version 144 contained 29,730 records from 4,979 plots, representing 20 distinct ecozones across all forested biogeographic and climate zones. From this, we drew 1,319 records that met our criteria, as outlined below (Fig. 1). This analysis focused on mature forests with no known history of significant disturbance or management. 147 There is evidence that stand age influences patterns of C flux and allocation in forest ecosystems, and can confound relationships between latitude and primary productivity (DeLucia et al., 2007; Gillman et al., 149 2015). To reduce any biasing effects of stand age, we included only stands of known age ≥ 100 years and 150 those described by terms such as "mature", "intact", or "old-growth". Since management can alter observed 151 patterns of C cycling (Šímová & Storch, 2017), sites were excluded from analysis if they were managed, 152 defined as plots that were planted, managed as plantations, irrigated, fertilised or included the term "managed" 153 in their site description. Sites that had experienced significant disturbance within the past 100 years were 154 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.

Table 2: Definitions and sample sizes of carbon flux 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		geographic 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 - R_{auto}	161	56	
ANPP	Above ground NPP	stem, foliage, optionally others (e.g., branch, reproductive, understory)	$ANPP_{stem} + ANPP_{foliage}$ (+ others)	278	86	
$ANPP_{stem}$	Stem growth component of $ANPP$	woody stems down to DBH $\leq 10 \mathrm{cm}$ (no branch turnover)	stem growth measurements scaled to biomass using allometries $$	264	96	
$ANPP_{foliage}$	Foliage component of $ANPP$	foliage	litterfall collection, with separation 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_{root} (as below)	22	13	
R_{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



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

157 Climate data

For C contains geographic coordinates associated with each measurement record and, when available, MAT158 and MAP as reported in the primary literature (Anderson-Teixeira et al., 2018). Based on the geographic 159 co-ordinates for each site, data on twelve climate variables – including MAT, MAP, temperature seasonality 160 (i.e., standard deviation across months), precipitation seasonality (i.e., coefficient of variation across months), annual temperature range, solar radiation, cloud cover, annual frost and wet days, potential evapotranspiration 162 (PET), aridity (MAP/PET), and vapor pressure deficit (VPD) – were extracted from five open-access climate datasets: WorldClim (Hijmans et al., 2005), WorldClim2 (fick & Hijmans, 2017), the Climate 164 Research Unit time-series dataset (CRU TS v4.03 (Harris et al., 2014), the Global Aridity Index and Potential Evapotranspiration Climate Database (Trabucco & Zomer, 2019), and TerraClimate (Abatzoglou et al., 2018) 166 (Table S1). Definitions and methods used to calculate each variable are included in Table S1. From these 167 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. 169 Where site-level data was missing for MAT or MAP, we used values from the WorldClim dataset. 170

Length of the growing season was estimated to the nearest month, where growing season months were defined 171 as months with mean minimum temperature > 0.5°C. This is consistent with the previous studies whose 172 hypothesis we were evaluating (Kerkhoff et al., 2005; Michaletz et al., 2014). We experimented with a 173 definition of growing season months including a moisture index, defined as (MAT - PET)/PET > -0.95174 (Kerkhoff et al., 2005; see also Michaletz et al., 2014). However, we found that including a moisture index 175 had minimal effect on the estimates of growing season length for the sites included here, and so chose to 176 exclude it. Monthly data for PET, precipitation, and temperature from CRU v 4.03 (Harris et al., 2014) and solar radiation from WorldClim2 (fick & Hijmans, 2017) were used to calculate mean monthly PET, 178 precipitation, temperature and solar radiation during the growing season.

180 Analyses

The effects of latitude and climate on C fluxes were analysed using mixed effects models using the package 'lme4' (Bates et al., 2015) in R v.3.5.1 (R Core Team, 2020). The basic model for all analyses included a fixed effect of latitude or climate and a random effect of plot nested within geographic area. Geographic areas—i.e., spatially clustered sites—were defined within ForC using a hierarchical cluster analysis on the distance matrix of the sites and a cutoff of 25km (Anderson-Teixeira et al., 2018). We experimented with inclusion of altitude as a fixed effect, as productivity is known to decline with elevation (Muller-Landau et al., 2020), but excluded it from the final models because it added very little explanatory power – that is, the

difference in AIC (ΔAIC) relative to models excluding altitude was generally small (often ΔAIC <2). Effects
were considered significant when inclusion of the fixed effect of interest resulted in p \leq 0.05 under an ANOVA
test, and $\Delta AIC \geq$ 2.0 relative to a corresponding null model. All R^2 values presented here are marginal R^2 values, and refer to the proportion of variation explained by only the fixed effects. Specific analyses are as
described below.

We first examined the relationship between latitude and C fluxes (Q1; Table 1). We tested models with 193 latitude as a first-order linear, second-order polynomial, and logarithmic term. For brevity, we henceforth refer to first-order linear models as "linear" and second-order polynomial models as "polynomial". We selected as 195 the best model that with the highest Δ AIC relative to a null model with no fixed term, with the qualification that a polynomial model was considered an improvement over a linear model only if it reduced the AIC value 197 by 2.0 or more. In addition, pairwise comparisons of R^2 values were carried out for a selection of pairs of C 198 fluxes to test for differences among variables in the proportion of variation explained by latitude and climate. 199 Models were run on data from sets of sites that were common to each pair, in order to account for variation 200 in the number of data points included. To standardise for variation in degrees of freedom across model types, 201 only linear and logarithmic models were included in the pairwise analysis. 202

To test whether trends in component fluxes across latitude sum to match those of larger fluxes, regression lines 203 for smaller component fluxes were summed to generate new estimates of larger fluxes. Because no fluxes were 204 significantly better predicted by a logarithmic or polynomial fit than by a linear fit, we used linear fits for all 205 fluxes in this analysis. We then determined whether these summed predictions fell within the 95% CI for the larger flux across the entire latitudinal range. Confidence intervals for the line of best fit for the larger flux were 207 estimated using the 'bootMer' function, a parametric bootstrapping method for mixed models (Bates et al., 2015). This function carried out 2000 simulations estimating the line of best fit, using quantiles at 0.025 and 209 0.975 to estimate 95% CIs. This analysis was applied to the following sets of fluxes: (1) $GPP = NPP + R_{auto}$ (2) NPP = ANPP + BNPP, and (3) $ANPP = ANPP_{foliage} + ANPP_{stem}$. In addition, we estimated total 211 belowground C flux (TBCF, not analyzed due to limited data) as $TBCF = BNPP + R_{root}$. 212

We next examined the relationships of C fluxes to climate variables (Q2-Q4; Table 1). We tested first-order linear, second-order polynomial, and logarithmic fits for each climate variable. Again, polynomial fits were considered superior to first-order linear fits only if inclusion of a second-order polynomial term resulted in $\Delta \text{AIC} \geq 2.0$ relative to a first-order linear model. We tested relationships of each C flux (Table 2) against each climate variable (Table S1). Variables which were not significant explanatory variables or which explained <20% of variation in C fluxes are only presented in SI.

Linear models were used to investigate the potential joint and interactive effects of MAT and MAP on carbon fluxes (Q2; Table 1). An additive model including MAP in addition to MAT was accepted when Δ AIC >2 relative to a null including only MAT as a fixed effect. An interactive model containing a MAT x 221 MAP interaction was accepted when $\Delta AIC > 2$ relative to a null including MAT and MAP as fixed effects. Variation in allocation to component carbon fluxes was explored for three groupings: (1) $GPP = NPP + R_{auto}$, (2) NPP = ANPP + BNPP, and (3) $ANPP = ANPP_{foliage} + ANPP_{stem}$. For each group, measurements 224 taken at the same site and plot, and in the same year, were grouped together. For groups (1) and (2), where 2 of the 3 flux measurements were available for a given site, plot, and year, these measurements were 226 used to calculate the third. We then calculated the ratio of each pair of component fluxes $(NPP:R_{auto};$ $ANPP:BNPP;ANPP_{foliage}:ANPP_{stem}).$ The logs of these ratios were regressed against latitude, MAT, 228 MAP, and temperature seasonality, using the linear models specified above. Cook's distance analyses were 229 carried out for each of the models, and extreme outliers removed.

To test whether and how C fluxes varied with climate when standardised by growing season length (Q5;
Table 1), we first standardized all annual C fluxes by dividing by growing season length (as defined above).
We then derived four variables to describe growing season climate, specifically growing season temperature,
precipitation, solar radiation, and PET (Table S1). We tested for correlations between these standardised
fluxes and growing season climate variables, using only first-order linear models.

All analyses were conducted in R v.3.5.1 (R Core Team, 2020). Code and data necessary to reproduce all results are available through GitHub (https://github.com/forc-db/Global_Productivity) and archived in Zenodo (DOI: TBD).

239 Results

In total, we analyzed 1,319 records from nine forest autotrophic C flux variables taken from forests that had experienced no major anthropogenic disturbances within the past 100 years. These records represented a total of 255 plots in 154 distinct geographic areas across all forested biogeographic and climate zones (Figs. 1, S1; Table 2).

244 Q1. How do C fluxes vary with latitude?

All major carbon fluxes decreased with latitude (Fig. 2; Table S2). Latitude was a strong predictor for many of the carbon fluxes, particularly the larger fluxes (Table S2, S6). Latitude explained 64% of variation in GPP (n = 243, p<0.0001), 50% in NPP (n = 161, p<0.0001) and 44% in ANPP (n = 278, p<0.0001). The C fluxes that were most poorly predicted by latitude were $BNPP_{fine.root}$ (n = 88, p< .01, R^2 =0.17) and

 249 $ANPP_{stem}$ (n = 264, p<0.0001, R^2 =0.18). The relationship with latitude was best fit by the first-order linear model, with the exception of NPP and R_{root} , for which a logarithmic model was a slightly – but not significantly – better fit.

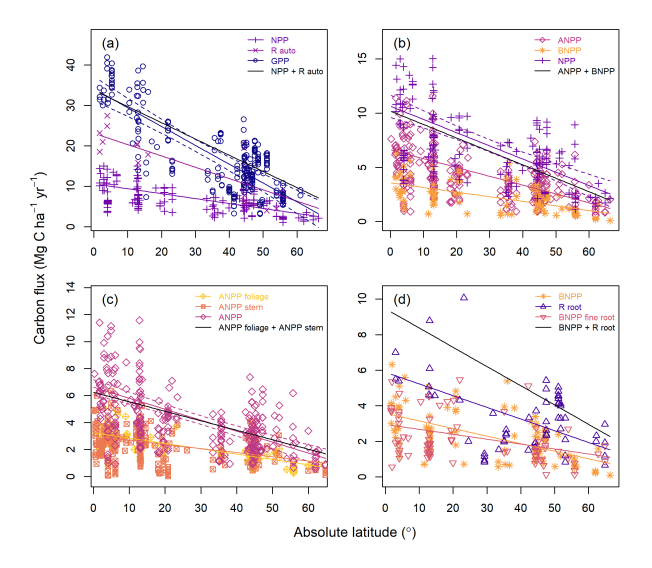


Figure 2: Latitudinal trends in forest autotropic carbon flux. Plotted are linear models, all of which were significant (p < 0.05) and had AIC values within 2.0 of the best model (for two fluxes, logarithmic fits were marginally better; Table S2). Each panel shows major C fluxes together with component 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, has insufficient data (n=9) to support a regression

Smaller component fluxes summed approximately to larger fluxes across the latitudinal gradient (Fig. 2).

That is, modeled 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 $ANPP_{stem}$, fell

almost completely within the confidence intervals of the regressions of field estimates of GPP, NPP, and

- 256 ANPP, respectively.
- ²⁵⁷ We found no evidence of systematic variation in C allocation with latitude or climate (Fig. S3). Of 12
- relationships tested (3 ratios among C flux variables regressed against latitude, MAT, MAP and temperature
- seasonality), none were significant (all p>0.05).
- 260 Q2. How do C fluxes relate to MAT and MAP?
- All fluxes increased with MAT (all p<0.05; Figs. 3-4, S4-S5, Table S2). For eight of the nine fluxes,
- 262 this relationship was linear. For BNPP the best fit was a lognormal fit, though this was not significant
- $_{263}$ (Δ AIC <2). As with latitude, MAT tended to explain more variation in the larger fluxes (GPP, NPP,
- $_{264}$ ANPP, R_{auto}) and ANPP_{foliage} (all $R^2 > 0.4$) than in subsidiary and belowground fluxes (ANPP_{stem},
- R_{root} , $BNPP_{fine.root}$; all $R^2 < 0.25$; Table S6).
- ₂₆₆ MAP was a significant (p<0.05) predictor of all fluxes (Figs. 4a, S4-S5; Table S2). However, it explained
- little variation: with the exception of R_{auto} , MAP explained at most 25% of variation in C flux. All fluxes
- increased with MAP up to at least 2000 mm, above which responses were variable (Figs. 4, S4-S5).
- There was a significant additive effect of MAT and MAP on GPP, ANPP and R_{auto} (Fig. 3, Table S3), and
- a significant interactive effect between MAT and MAP for NPP and $ANPP_{stem}$ (Fig. 3, Table S3). The
- interaction was negative for NPP and positive for $ANPP_{stem}$. For $ANPP_{foliage}$, BNPP, $BNPP_{fine.root}$,
- and R_{root} , MAP did not have a significant effect when accounting for MAT (Fig. 3, Table S3).

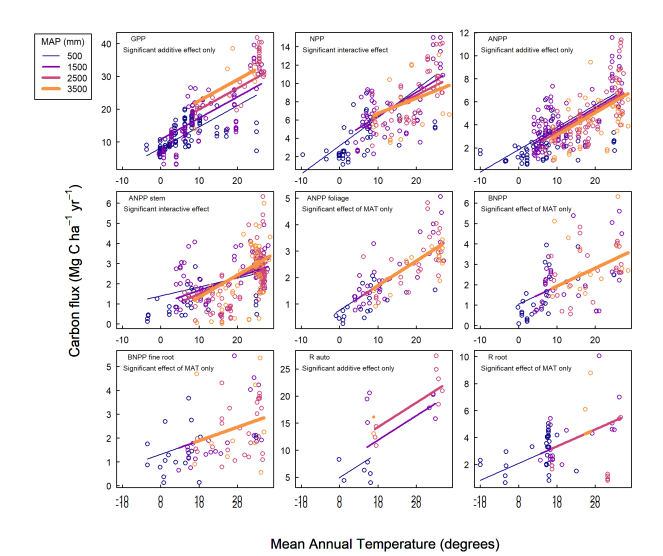


Figure 3: Interactive effects of mean annual temperature and precipitation on annual forest carbon fluxes. For visualization purposes, data points are grouped into bins of 0 - 1000, 1001 - 2000, 2001 - 3000, and >3000mm mean annual precipitation, and lines of best fit models are plotted for mean annual precipitation values of 500, 1500, 2500, and 3500mm. Significance is defined as p < 0.05.

273 Q3. How do C fluxes relate to other annual climate variables?

All C flux variables showed a significant relationship with annual PET. The relationship was logarithmic for $ANPP_{foliage}$, $BNPP_{fine.root}$ and R_{root} , and polynomial for all other fluxes (Fig. 4c, S4-5; Table S2). We found strong evidence for a saturation point or peak with PET: C fluxes tended to increase at values below 1000mm, before saturating between 1200 and 1700mm. There was also evidence that some C fluxes begin to decrease at values above 1800mm PET.

Mean annual VPD was a significant predictor of all C fluxes. $ANPP_{foliage}$, $BNPP_{fine.root}$ and R_{root} showed

- ²⁸⁰ a logarithmic relationship with VPD, but all other fluxes showed a polynomial relationship (Figs. 4d, S4-5;
- Table S2). C fluxes initially increased with VPD, before saturating at around 0.8 kPa, after which point
- they began to decrease.
- All fluxes, with the exception of R_{root} , showed a significant positive relationship with solar radiation (Figs.
- 284 S4-S5, Table S2). Solar radiation explained a low proportion of variability (<30%) in all C fluxes.
- Annual wet days, cloud cover, and aridity were poor or non-significant predictors of variation in C fluxes,
- 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. All regressions are significant (p < 0.05).

- 287 Q4. What is the role of seasonality in explaining C fluxes?
- ²⁸⁸ Variables describing temperature seasonality temperature seasonality, annual temperature range, annual
- frost days, and length of growing season were strongly correlated with both latitude and MAT (all $r \ge 0.2$;
- ²⁹⁰ Fig. S2), and were consistently identified as strong univariate predictors of C fluxes (Figs. 4, S4-S7).
- All fluxes decrease with increasing temperature seasonality, though the shape of this relationship varies (all
- p<0.05; Figs. 4e, S6-7; Table S2). Temperature seasonality was strongly correlated with annual temperature
- ²⁹³ range, which was likewise a similarly strong predictor of C fluxes (Table S2). C fluxes were highest where
- temperature seasonality = 0, and at an annual temperature range of 15° C or lower (i.e., in the tropics).
- In contrast, there was no significant effect of precipitation seasonality on C fluxes at this global scale. Both
- maximum vapour pressure deficit and water stress months were poor or non-significant predictors of variation
- in C fluxes (Figs. S6-S7; Table S2).
- We found a significant relationship between length of growing season and C fluxes, with all fluxes showing a
- positive relationship with length of growing season (Figs. 4e, S6-S7; Table S2). Length of growing season was
- a strong predictor of C fluxes, explaining 53% of variation in GPP, 38% of variation in NPP, and 34% of
- variation in ANPP (all p<0.05; Table S2), but it was a weaker predictor than MAT for all fluxes analysed
- 302 (Table S4).
- 303 Q5. Within the growing season, how do C fluxes vary with climate?
- When annual C fluxes were standardized by growing season length (in integer number of months), correlations
- with growing season climate were generally weak (Figs. S8-S9). ANPP increased with growing season
- temperature ($R^2 = 0.09$, p<0.001) and precipitation ($R^2 = 0.04$, p<0.05). Similarly, $ANPP_{foliage}$ increased
- slightly with growing season temperature ($R^2 = 0.16$, p<0.01) and precipitation ($R^2 = 0.09$, p<0.05). Growing
- season solar radiation was positively correlated with on BNPP ($R^2 = 0.17$, p<0.001) and BNPP_{fine.root}
- $(R^2 = 0.13, p < 0.01)$. Growing season PET had a positive influence on GPP ($R^2 = 0.15, p < 0.01$), NPP ($R^2 = 0.15, p < 0.01$).
- $_{310} = 0.07, \, \text{p} < 0.01), \, BNPP \, (R^2 = 0.23, \, \text{p} < 0.0001), \, BNPP_{fine.root} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, ANPP_{stem} \, (R^2 = 0.10, \, \text{p} < 0.05), \, \text{and} \, R^2 = 0.00, \, \text{and} \, R^2 + 0.00, \, \text{and} \, R^2 + 0.00, \, \text{and} \, R^2 + 0$
- $_{311} = 0.06$, p<0.05). All other relationships were non-significant.

2 Discussion

- Our analysis of a large global database (ForC) clarifies how autotrophic C fluxes in mature forests vary
- with latitude and climate on a global scale. We show that, across all nine variables analyzed, annual C
- 315 flux decreases continually with latitude (Fig. 2), a finding that confirms multiple previous studies and
- contradicts the idea that productivity of temperate forests rivals or even exceeds that of tropical forests

(Huston & Wolverton, 2009; Luyssaert et al., 2007). At this global scale, C fluxes increase approximately in 317 proportion to one another, with component fluxes summing appropriately to larger fluxes and no detectable differences in allocation across latitude or climates (Figs. 2, 4, S3). Similarly, we show broad - albeit not 319 complete - consistency of climate responses across C fluxes, with the observed latitudinal variation primarily attributable to temperature and its seasonality (Figs. 3-4). Water availability is also influential, but of 321 secondary importance across the climate space occupied by forests (Figs. 3-4). Contrary to prior suggestions 322 that the majority of variation in C cycling is driven primarily by the length of the growing season (Enquist et 323 al., 2007; Kerkhoff et al., 2005; Michaletz et al., 2014), we find modest explanatory power of growing season 324 length and small but sometimes significant influences of growing season climate (Figs. 4f,S6-S9). Together, 325 these findings yield a unified understanding of climate's influence on forest C cycling. 326

Our findings indicate that, among mature, undisturbed stands, forest C fluxes are unambiguously highest 327 in the tropical regions, and the relationship with both latitude and MAT is approximately linear (Table 1, 328 Q1,Q2; Figs. 2, 4). This contrasts with the suggestion that C fluxes (e.g., NPP, ANPP, ANPP_{stem}) of 329 temperate forests are similar to or even greater than that of tropical forests (Huston & Wolverton, 2009; 330 Luyssaert et al., 2007). Previous indications of this pattern may have been an artifact of differences in 331 stand age across biomes. Compared to tropical forests, the temperate forest biome has experienced more widespread anthropogenic disturbance and has a larger fraction of secondary stands (Potapov et al., 2008; 333 Poulter et al., 2018; Yu et al., 2014), so analyses comparing across latitudinal gradients without controlling for stand age risk confounding age with biome effects. Because carbon allocation varies with stand age 335 (Anderson-Teixeira et al., 2013; DeLucia et al., 2007; Doughty et al., 2018; Yu et al., 2014), age differences may introduce systematic biases into analyses of C fluxes across latitude or global climatic gradients. For 337 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 and non-woody productivity (DeLucia et 339 al., 2007; Doughty et al., 2018; Kunert et al., 2019; Piao et al., 2010). Thus, findings that temperate forest 340 productivity rivals that of tropical forests are likely an artifact of different forest ages across biomes. 341

We show that C fluxes are broadly consistent in their responses to climate drivers on the global scale, with no trends in C allocation among the variable pairs tested (Figs. 2, S3). This parallels the observation that C allocation across multiple C fluxes varies little with respect to climate along a steep tropical elevational gradient (Malhi et al., 2017; but see Moser et al., 2011), and is not surprising given that carbon allocation within forest ecosystems is relatively constrained (Enquist, 2002; Litton et al., 2007; Malhi et al., 2011). We find no significant trend in the allocation of *GPP* between production and respiration across latitude or climate (*NPP:Rauto*; Fig. S3), counter to the idea that tropical forests have anomalously low *CUE*

(Anderson-Teixeira et al., 2016; DeLucia et al., 2007; Malhi, 2012), which is predicted by most models (Collabti et al., 2020). In contrast, Collabti et al. (2020) found that forest production efficiency increased with temperature—a finding that is consistent in direction with the insignificant trends observed here (Fig. S3). 351 Previously observed differences in CUE between old-growth tropical forests relative to (mostly younger) extratropical forests are likely an artifact of comparing stands of different age, as CUE is known to decline 353 with forest age (Collalti et al., 2020; DeLucia et al., 2007; Piao et al., 2010). Another previously observed pattern for which we find no support is a tendency for belowground C allocation to decrease with increasing 355 temperature (Gill & finzi, 2016; Moser et al., 2011); rather, we observe no trends in allocation between 356 ANPP and BNPP across latitudes. Failure to detect significant trends in C allocation with respect to 357 climate in this analysis does not imply that none exist; rather, it suggests that, at this global scale, differences 358 are subtle and/or that more careful methodological standardization and/or more data is required to detect them. 360

Despite the broad consistency of climate responses across C fluxes, climate explains lower proportions of 361 variability among some of the subsidiary C fluxes (e.g., ANPP_{stem}, BNPP, BNPP_{fine.root}; Fig. 2; Tables S2, S6). There are two, non-exclusive, potential explanations for this. First, it may be that methodological 363 variation is larger relative to flux magnitude for some of the subsidiary fluxes. Belowground fluxes in particular are difficult to quantify, and measurement methods for the belowground fluxes considered here may use 365 fundamentally different approaches in different sites (e.g., minirhizotrons, ingrowth cores, or sequential coring for $BNPP_{fine.root}$; root exclusion, stable isotope tracking, or gas exchange of excised roots for R_{root}), and 367 sampling depth is variable and often insufficient to capture the full soil profile. $ANPP_{stem}$, which is also poorly explained by latitude or climate, is more straightforward to estimate but subject to variability introduced by 369 methodological differences including minimum plant size sampled and choice of biomass allometries (Clark 370 et al., 2001). That said, methodological variation and uncertainty affect all of fluxes considered here, and 371 some of the larger fluxes that vary more strongly with respect to climate (ANPP, NPP) are estimated by 372 summing uncertain component fluxes. Second, differences among variables in the proportion of variation 373 explained by climate may be attributable to more direct climatic control over GPP than subsidiary fluxes. 374 That is, subsidiary fluxes may be shaped by climate both through its influence on GPP and through its 375 influence on CUE and C allocation. 376

Temperature and its seasonality were the primary drivers of C fluxes on the global scale (Table 1, Q2, Q4; Figs. 2-4), consistent with a long legacy of research identifying temperature as a primary driver of forest ecosystem C cycling (e.g., Lieth, 1973; Luyssaert et al., 2007; Wei et al., 2010). We find little evidence of any non-linearity in temperature's influence on C fluxes. The relationship of all fluxes to MAT as an individual

driver were best described by a linear function (Table S2) – with the exception of BNPP, whose response to MAT was close to linear (Fig. 4a). This result contrasts with the hypothesis that fluxes saturate with MATbelow approximately 25°C MAT (Huston & Wolverton, 2009; Luyssaert et al., 2007). It remains possible 383 that fluxes decline above this threshold (Larjavaara & Muller-Landau, 2012; Sullivan et al., 2020), as is also consistent with tree-ring records indicating that tropical tree growth declines at high temperatures (e.g., 385 Vlam et al., 2014). However, these higher temperatures also tend to be associated with high PET and VPD, both of which are associated with reduced C fluxes (Figs. 4c-d, S4-S5; Slot & Winter, 2018; Zani et al., 2020). 387 Indeed, while temperature responses dominate at this global scale and within the climate space occupied 388 by forests, the effects of temperature are moderated by moisture availability (Table 1, Q2,Q3; Figs 3-4). Specifically, C fluxes are reduced under relatively dry conditions (i.e., low MAP; high VPD) and sometimes 390 under very high precipitation (Figs. 3-4). The observed positive interaction between MAT and MAP for 391 $ANPP_{stem}$ on the global scale (Fig. 3) is consistent with an analysis showing a similar interaction for ANPP392 in tropical forests, also with a cross-over point at ~20^oC (Taylor et al., 2017). However, we detect no such 393 interaction for ANPP or most other C fluxes, and we find a contrasting negative interaction for NPP (Fig. 3), suggesting that more data are required to sort out potential differences in the interactive effects of MAT395 and MAP on C fluxes in the tropics.

Forest C fluxes decline with temperature seasonality (Table 1, Q4; Fig. 4e), as is to be expected given that 397 fluxes are minimal during winters. A temperature-defined growing season length correlated strongly with 398 global-scale variation in annual C flux (Table 1, Q5; Fig. 4f; see also Churkina et al., 2005), consistent with the idea that the latitudinal gradient in carbon flux is attributable more to shorter growing seasons at high 400 latitudes than to inherently lower rates of photosynthesis or respiration by high-latitude forests (Enquist et al., 2007; Fu et al., 2019). While there is evidence that trees in high-latitude forests have adaptations to 402 maximize photosynthesis at low temperatures (Helliker & Richter, 2008; Huang, 2019), this is not sufficient to yield growing season fluxes comparable to those of tropical forests, as indicated by a number of positive 404 correlations between monthly mean flux during the growing season and growing season temperature, solar 405 radiation, and PET (Table 1, Figs. S8-S9). Thus, we reject the hypothesis that growing season length alone 406 accounts for global-scale variation in productivity-i.e., that there is no relationship between C flux per month 407 of the growing season and growing season climatic conditions (Table 1, Q5; Kerkhoff et al., 2005; Enquist et 408 al., 2007; Michaletz et al., 2014). Rather, annual C flux is shaped by both growing season length and the 409 climate of peak growing season months (Chu et al., 2016; Fu et al., 2019). Given strong co-variation between growing season length and MAT (Fig. S2; Chu et al., 2016), accurately partitioning this variation will require 411 data on intra-annual variation in C flux coupled with a higher-precision metric of growing season length than the monthly-resolution metric used here (e.g., based on leaf phenology or C exchange, sensu Fu et al., 2019). Fu et al. (2019) find that global-scale geographic variation in annual NEE is driven more strongly by growing season length than by carbon uptake rates within the growing season, whereas interannual variation in NEE and GPP at any given site appears to be driven predominantly by the maximum rate of C uptake, as opposed to growing season length (Fu et al., 2019; Zhou et al., 2016). Further analysis of interannual variation in C fluxes in relation to climate will be valuable to disentangling how seasonality shapes broad geographic patterns in forest C flux.

Our analysis clarifies how annual forest autotrophic C fluxes vary with latitude and climate on a global scale. 420 To the extent that patterns across broad scale climatic gradients can foretell ecosystem responses to climate change, our findings suggest that higher temperatures with similar moisture availability would result in a 422 generalized acceleration of forest C cycling (Figs. 2-3). This is consistent with observations of continental-423 to global-scale increases over time in GPP (Li & Xiao, 2019), ANPP_{stem} (Brienen et al., 2015; Hubau et 424 al., 2020), tree mortality (Brienen et al., 2015; McDowell et al., 2018), soil respiration (Bond-Lamberty & 425 Thomson, 2010), and heterotrophic soil respiration (Bond-Lamberty et al., 2018). However, increasing C flux 426 rates are by no means universal (e.g., Rutishauser et al., 2020; Hubau et al., 2020), likely because other factors 427 are at play, including changes to other aspects of climate, atmospheric pollution (CO₂, SO₂, NO_x), and local disturbances. Moreover, forest ecosystem responses to climatic changes outside the temperature range to 429 which forest communities are adapted and acclimatized will not necessarily parallel responses across geographic gradients in climate. Indeed, tree-ring studies from forests around the world indicate that tree growth rates 431 - along with $ANPP_{stem}$ and possibly other ecosystem C fluxes – respond negatively to temperature (e.g., Helcoski et al., 2019) **REFS**. Furthermore, in the tropics, climate change will push temperatures beyond 433 any contemporary climate, and there are some indications that this could reduce forest C flux rates (Mau 434 et al., 2018; Sullivan et al., 2020) if paralleled by VPD increases (Smith et al., 2020). Further research is 435 required to understand the extent to which forest responses to climate change will track the observed global 436 gradients, and the time scale on which they will do so. In the meantime, understanding the fundamental climatic controls on annual C cycling in Earth's forests sets a firmer foundation for understanding forest C 438 cycle responses to accelerating climate change.

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