ENSO Drives interannual variation of forest woody growth across the tropics

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ENSO drives interannual variation of forest woody growth across the tropics

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Summary

Meteorological extreme events such as El Niño events are expected to affect tropical forest net primary production (NPP) and woody growth, but there has been no large scale empirical validation of this expectation. We collected a large high temporal resolution dataset (for 1-13 years depending upon location) of more than 172,000 stem growth measurements using dendrometer bands from across 14 regions spanning Amazonia, Africa and Borneo in order to test how much month-to-month variation in stand-level woody growth of adult tree stems (NPP_{stem}) can be explained by seasonal variation and interannual meteorological anomalies. A key finding is that woody growth responds differently to meteorological variation between tropical forests with a dry season (where monthly rainfall is < 100 mm), and aseasonal wet forests lacking a consistent dry season. In seasonal tropical forests a high degree of variation in woody growth can be predicted from seasonal variation in temperature, vapour pressure deficit, in addition to anomalies of soil water deficit, and shortwave radiation. The variation of aseasonal wet forest woody growth is best predicted by the anomalies of vapor pressure deficit, water deficit, and shortwave radiation. In total, we predict the total live woody production of the global tropical forest biome to be 2.16 Pg C year^{-1}, with an interannual range 1.96-2.26 Pg C year^{-1} between 1996-2016, and with the sharpest declines during the strong El Niño events of 1997/8 and 2015/6. There is high geographical variation in hotspots of El Niño-associated impacts, with weak impacts in Africa, and strongly negative impacts in parts of SE Asia and extensive regions across central and eastern Amazonia. Overall, there is high correlation (r = -0.75) between the annual anomaly of tropical forest woody growth and the annual mean of the El Niño 3.4 index, driven mainly by strong correlations with anomalies of soil water deficit, vapor pressure deficit, and shortwave radiation.

1. Introduction

Tropical forest productivity is amongst the highest of terrestrial ecosystems [1,2], but the amount of carbon allocated to woody stems (NPP_{stem}) within tropical forests is highly variable [3–6]. We here define NPP_{stem} as the productivity of above-ground woody tissue including trunks and branches, but excluding fine woody material such as twigs, and woody coarse roots. NPP_{stem} is not the largest component of carbon allocation, typically accounting for only 20-30% of NPP and 5-10% of gross primary productivity (GPP) [7], but, because woody material is long-lived, it is a major determinant of forest biomass and carbon residence time.

In this paper we examine the seasonal and interannual variation of woody growth (NPP_{stem}) across the tropical forest biome. Meteorological variation is likely to be an important control on seasonal changes in NPP_{stem} and has only rarely been tested [8–11], but never so at a pantropical scale. Examination of NPP_{stem} variation has largely been limited to coarse temporal variation at interannual or multi-year time scales. NPP_{stem} is usually estimated by repeat census of tree diameters coupled with the use of allometric equations to translate changes into above-ground biomass. However forest census intervals typically span multiple years, and this obscures the relation of NPP_{stem} to seasonal meteorological variation and meteorological extreme events. Dendrometers enable much higher resolution tracking of tree growth (typically monthly resolution for manual dendrometers, daily for automatic dendrometers), but have not previously been employed in a consistent multi-site and multi-regional analysis. Here we present and analyse a uniquely extensive pantropical dataset of tree growth comprising more the 8,725 trees. The standardized protocol for measuring NPP_{stem} from the Global Ecosystem Monitoring network (www.gem.tropicalforests.ox.ac.uk) is unique for its use of manual dendrometers to provide high temporal resolution (~ 1-3 months), enabling examination of seasonal and interannual variation in NPP_{stem}.

At an individual level, carbon allocation to NPP_{stem} is thought to be affected by several biological processes, including photosynthetic uptake [7], its balance with respiration [12–14], tradeoffs in carbon allocation between woody parts, canopies and roots[7,15–17], source vs. sink driven biological cues[18,19], and most especially the crown exposure to light[20,21]. However when aggregated to the stand level, many of these individual-level biological drivers of growth are marginalized. After all, the amount of light and rainfall a forest receives and utilises is not so much a function of its stand structure, but of seasonality in weather and its geographic location. Here we do not specifically address the non-climatic components of spatial
variation in NPP\textsubscript{stem}, because this is an inherently more complicated question where the allocation of carbon to NPP\textsubscript{stem} is dependent upon a number of interacting factors and processes such as soil fertility, species composition, and carbon use efficiency \cite{12,20}. In this study, we purposely do not aim to explain the biological, disturbance related (e.g. catastrophic tree mortality events), or other spatially varying abiotic controls (e.g. soil fertility) upon NPP\textsubscript{stem}, but rather how month-to-month meteorological variation can explain seasonal changes in NPP\textsubscript{stem}.

Seasonal differences in NPP\textsubscript{stem} (or xylogenesis) are likely to be concentrated towards the transition between the dry to the wet seasons because xylogenesis is inhibited when cell turgor is low \cite{18}, and trees recovering from extreme drought stress may improve their hydraulic conductivity by replacing xylem that have cavitated over the dry season \cite{22}. This pattern may be stronger in highly seasonal forests that experience annual drought stress, whereas differences in the temporal allocation of carbon to woody growth may be non-existent in aseasonal forests where few droughts occur to impair stem hydraulic conductivity. The extent to which a seasonal increase in woody stem growth reflects an increase in overall productivity, or simply a shift in carbon allocation among roots, wood, the canopy, and non-structural carbohydrate storage pools remains uncertain. In lowland Amazonia, allocation shifts were found to be more important than overall changes in carbon assimilation in explaining interannual variability in canopy, wood, and fine root growth rates \cite{16,17}.

Here, we utilize the anomalous drought conditions produced by El Niño events to examine how much spatial and temporal variation in in NPP\textsubscript{stem} can be explained by purely meteorological variation. El Niño events tend to increase temperatures and atmospheric water vapour deficit (VPD) across the tropics, and cause strong declines in precipitation in some regions, most notably Amazonia and insular SE Asia \cite{23}. These meteorological factors are likely to affect NPP\textsubscript{stem} through underlying ecophysiological mechanisms. We focus on relating temperature, VPD, cloudiness, and precipitation metrics to NPP\textsubscript{stem}. First, negative precipitation anomalies and soil water deficits are likely to impede growth by increasing soil-root hydraulic resistance \cite{24} and reducing stem conductance through cavitation \cite{25}. Precipitation deficits from drought can eventually lead to declines in NPP \cite{26}; but see \cite{11}. Relating precipitation to forest growth can be challenging because monthly precipitation can be decomposed into numerous metrics with greater ecophysiological relevance, but here we focus on four aspects: a one dimensional Thornthwaite-Mather water balance model from a high resolution climate product \cite{27}, climatic water deficit (CWD) which is a simpler proxy for sub annually varying soil water deficit, the maximum climatic water deficit (MCWD) which represents that maximum CWD for the preceding 12-months \cite{28}, and lagged differences in monthly precipitation which can serve as a proxy for the transition between dry and wet seasons. Second, temperature, even in the tropics, can control or act as a cue for much of the seasonality of growth and carbon allocation \cite{29,30}, yet reductions in photosynthesis occur when trees are exposed to temperatures beyond their optimum for photosynthesis \cite{31–33}. A recent comparison of an evergreen and semi-deciduous forest in Panama found that the community temperature optimum closely mirrored the mean maximum daytime temperature \cite{33}. Thus positive temperature anomalies during drought events may push leaves over their optimum temperature for photosynthesis, increase respiration costs \cite{34}, and by extension reduce the amount of plant expendable carbon that can be allocated to NPP\textsubscript{stem}. Alternatively, higher temperatures may push forest canopies into or beyond their optimal temperature range and either leading to an increase or saturation of gross primary productivity \cite{35}. Third, high temperatures with invariant or reduced atmospheric humidity lead to high VPD, which can induce stomata to close \cite{36–38} even when soil moisture is non-limiting \cite{39}. Of course stomatal conductance does not work independent of leaf energy balance, so positive VPD anomalies may result in a reduction of leaf conductance which may induce higher leaf surface temperatures and VPDs, and perhaps further reduce photosynthesis. Finally, shortwave radiation is highly correlated with photosynthetic assimilation of CO\textsubscript{2}. El Niño events can reduce cloudiness in the same regions where it reduces precipitation, which results in increased shortwave irradiance. A positive shortwave anomaly could increase photosynthesis in tropical regions with weak dry seasons, such as northwest Amazonia, and Borneo \cite{30}, although prior evidence suggests an increase in carbon assimilation may not necessarily manifest in higher NPP\textsubscript{stem} \cite{5,7,40}.

Specifically we address the following questions:
(1) How much variation in tropical NPP\textsubscript{stem} can be explained by meteorological variation?
(2) What meteorological drivers most affect NPP\textsubscript{stem} during El Niño associated drought events?
(3) What is the total annual woody production of the tropical forest biome, how much does it decline during El Niño events, and which regions contribute most strongly to these declines?

2. Methods
2.1 Scaling from individuals to forest stand
We employed the standard protocols of the Global Ecosystems Monitoring (GEM) network, described at gem.tropicalforests.ox.ac.uk). Simply, constructed manual dendrometer bands were installed on trees and measured at intervals typically ranging from 1-3 months across 14 geographic regions encompassing a large rainfall gradient ranging from highly seasonal dry tropical forests to aseasonal wet tropical forests (Fig. 1 & SM Fig. 1), encompassing 50 individual plots. In total, 8,725 trees were attached with dendrometers, and more than 187,000 readings were taken over 65 plot-years of data. The duration of measurement and number of observations varied across plots (See Table 1). Dendrometers were installed on a subset of adult trees (≥10 cm DBH). The sample coverage and size distribution of trees with dendrometer bands varied across plots, and rarely matched the corresponding size distribution from the full plot census of all adult trees. A nonlinear height allometry was derived for each site, and used to update tree height with every dendrometer measurement (detailed in SM section 1). The biomass was estimated for each tree using allometric equation from Chave et al. (2014) with wood density derived from the Global Wood Density Database and for each species or regional-genus mean. The mean individual growth rate in Mg C was calculated using a dry-biomass carbon content of 47.8%. This growth rate was multiplied by the number of individuals (≥10 cm DBH) in each plot when the number of trees with dendrometers was > 50% of the number of trees in the plot. We also applied the mean growth rate to all trees in the plot when 30-50% of the trees had dendrometer bands and the median DBH of trees with dendrometer bands matched the median DBH of all trees in the plot to within 5%. When measurements did not meet these criteria, but still had at least 60 individuals with dendrometer measurements - size, wood density, and estimated height were used to construct non-linear generalized additive models to predict growth for each date, which were then used to predict total carbon accumulation for each tree in the plot that did not have a dendrometer. The resulting NPP\textsubscript{stem} observation is the scaled forest-level woody growth (in carbon units Mg C month\textsuperscript{-1} ha\textsuperscript{-1}) estimated by summing the observed growth rates from trees with dendrometer bands, and the sum of tree level growth predictions over trees in the plot lacking dendrometer bands. The effects of stochastic tree mortality events are large upon month-to-month changes in forest biomass. Our goal was to isolate the climatic signal upon only live woody tree growth so we removed the demographic responses of carbon entering the plot from tree recruitment, and carbon leaving the plot from tree mortality. To do so, the regression growth models of each date were applied to a single fixed date census corresponding to each forest plot. Finally it is worth noting that the error from scaling the individual growth to plot-level NPP\textsubscript{stem} are not propagated throughout subsequent analyses on the plot-level estimates of NPP\textsubscript{stem}.

2.2 Deriving meteorological predictors
Temperature and VPD data time series for each site were derived from a gridded climate product (TerraClimate) [27]. The TerraClimate product is a statistically downscaled (~4 km) merge between the CRU TSv4.01 empirical climate interpolation [44] and the JRA-55 climate reanalysis product [45]. Meteorological time series from TerraClimate were compared with downscaled site-level meteorological predictions from local automatic weather stations and the ERA-Interim climate reanalysis product (detailed in SM section 2) [46]. The monthly meteorological estimates from TerraClimate corresponded well with the downscaled site level meteorological records for most sites (SM Section 2; SM Figs. 2 & 3) with the exception of shortwave radiation at the Borneo sites. Surface level shortwave radiation over wet tropical forest regions is not well estimated by most climate reanalysis products, so we calculated the 3-month moving mean cloud fraction using the satellite derived NOAA CDR PATMOS-X v5.3 cloud properties product [47] and the 3-month moving surface level shortwave radiation estimates from the Clouds and the Earth's Radiant Energy Budget product [48].

2.3 Estimating the effects of meteorological drivers upon NPP\textsubscript{stem}
We calculated the long-term monthly means (•) of monthly diurnal min/mean/max were calculated for air temperature, VPD, and shortwave radiation. We also calculated metrics of precipitation (monthly), water deficit (CWD and MCWD), a metric of the wet-dry season transition (detailed in SM Section 2). The monthly anomalies of each meteorological variable were calculated, and divided by their location specific interannual monthly standard deviation. The resulting anomaly terms are expressed in units of standard deviation (σ) from their long-term monthly mean. It is important to note that both the • and σ terms vary by month and the corresponding forest plot’s location. For example, a 1°C increase above the mean temperature in the month of August would be less than one unit σ at the Kenya site in the (highly seasonal) Bolivian Amazon, whereas it would be more than three units σ across all of the (relatively aseasonal) Borneo sites. Therefore both the • and σ terms have an inherent spatial context.

We fit generalized linear mixed models (GLMMs) and Generalized Additive Models (GAMs) to examine how NPP\textsubscript{stem} is affected by seasonal meteorological variables and their corresponding anomalies. Several of the meteorological covariates used in the model comparison process were highly correlated, so we
3.2 Overall explanatory power of the meteorologically driven model

http://mc.manuscriptcentral.com/issue-ptrsb
Our meteorologically driven final statistical models explained approximately 52% (35% excluding random effects) and 41% (20% excluding random effects) of NPP\textsubscript{stem} seasonal variation for tropical seasonal forests and aseasonal wet forests, respectively. The range in the amount of variation explained (R\textsuperscript{2}) was large across sites (Table 1), but the predictive distribution of the models generally covered the observed range of NPP\textsubscript{stem} (Fig. 2). The R\textsuperscript{2} of aseasonal wet forest sites improved the most when allowing random effects (i.e. variation in plot-specific mean values of NPP\textsubscript{stem}) which is due to the general lack of seasonal variation in NPP\textsubscript{stem}. Despite the improved performance, the plot specific intercept (random effect) acts as a categorical variable that cannot be applied for up-scaling the model across the tropics so we present conditional model predictions without random effects (Fig. 2c-h). A higher degree of predictive ability was found for sites with strongly pronounced dry seasons (e.g. the Kenia plots in Bolivia and the Santarém region plots in eastern Amazonia; Fig. 3a,c) while the R\textsuperscript{2} was poorest for the more aseasonal sites (e.g. in Borneo) where there was less seasonal variation in woody growth to explain (e.g. MLA, SAF; Table 1; Fig. 3f,g & SM Fig. 5). Despite this apparent increase in explained variation with increasing precipitation seasonality, this may be because the aseasonal wet forest model was estimated using far fewer observations (N = 110) than the seasonal forest model (N = 674).

3.3 Tropical NPP\textsubscript{stem} and its response to El Niño events
Overall, our pantropical scaling estimates that the mean total annual above-ground woody production of the tropical forest biome is 2.16 Pg C yr\textsuperscript{-1}, and this varied interannually in the range 1.96-2.26 Pg C (i.e. 12 %) between years 1996-2016. Global minima occur during El Niño events, with Amazonia and insular Southeast Asia being the most impacted (Figs. 4&5). The spatial anomalies of NPP\textsubscript{stem} are not consistent across El Niño events (Fig. 4). For example different parts of Amazonia were most strongly affected by the El Niño events in 1997/1998 and 2015/2016. Conversely the pronounced negative impact seems spatially consistent across eastern Borneo, whereas equatorial Africa may have been moderately negatively affected by the 1997/1998 El Niño but less so during the 2015/2016 event (with an important caveat that climatological products for this data-poor region are particularly unreliable).

The detrended long-term prediction of the anomaly in NPP\textsubscript{stem} is highly correlated with the moving annual average of the El Niño 3.4 Index (r = -0.7; Fig. 5). Hence interannual variation of the total woody growth of the tropical forest biome can be at least partially predicted from the El Niño 3.4 Index. The interannual anomaly of NPP\textsubscript{stem} is most highly correlated with the annual anomalies of VPD (r = -0.59), but also correlates with water deficit (r = -0.51), temperature (r = -0.49) and shortwave radiation (r = -0.38). This finding is consistent with inversion modelling results that show that the carbon cycle of the terrestrial tropics is strongly correlated with tropical land surface temperatures; however, our analysis suggests that the local mechanistic drivers are more linked to water deficits, VPD and shortwave radiation than to temperature (Fig. 2a,b).

4. Discussion
4.1 How much variation in tropical NPP\textsubscript{stem} can be explained by meteorological variation?
Using our statistical models, as much as 55% of monthly woody growth can be predicted for seasonal tropical forests, and 45% for aseasonal wet forests. This amount of explained variation on high temporal resolution changes in NPP\textsubscript{stem} is not so dissimilar from the variation in forest biomass changed explained over much longer periods of time by considerably more sophisticated forest simulation models (e.g. [54,55]). However the GLMMs presented here should not be viewed as authoritative, but rather as an initial attempt to understand and separate the effect of the long-term mean of month-to-month meteorological seasonality from interannual meteorological variation upon tropical forest woody growth. These statistical models are simplistic representations of complex biological responses. Tropical forests have to mitigate several forms of ecophysiological stress from meteorological variation and in many cases the underlying ecophysiological mechanisms of tropical forests response to drought are still not well understood [56]. So it is noteworthy that the models presented here do have predictive ability across all sites, and that this predictive ability is greater across the vast majority of tropical forest regions with rainfall seasonality (Figs. 1, 2 & 3; Table 1).

There are many opportunities to improve the model. The data used to fit the model are imbalanced across sites (Table 1), with notable data limitations for the aseasonal wet tropics. By extension the uncertainty and poorer predictive performance in the aseasonal wet forest regions is likely due to data deficiency, which will in many cases improve over time. The meteorological variables used in this study are often highly correlated, which precludes the incorporation of all relevant variables into a linear predictor because standard statistical methods cannot identify effects that are highly collinear. The environmental drivers used to model here also fail to capture temporal directionality. For example, the water deficit anomaly makes no distinction whether a soil is on a trend towards drying or wetting. The representation of temperature in the model also makes no distinction between short temporal pulses, versus longer
4.2 What meteorological drivers most affect NPP during El Niño associated drought events?

We can only make cautiously qualified statements about the most important meteorological drivers affecting growth because this question is hindered by both uncertainty in the true meteorological conditions, and by insufficient data at both ends of the extremes of a meteorological variable (e.g., where observations are needed during both anomalously wet and anomalously dry conditions). The effects of VPD are consistent and large across both the seasonal and aseasonal wet tropics, but in different ways. In the seasonal forest model, the effect of VPD only has explanatory power in the seasonal component, while the interannual anomaly does not appear to be important. Conversely in the aseasonal wet tropics, VPD has no effect upon the seasonal component (as variation is low in the aseasonal tropics; SM Fig. 6), but has a large effect in the interannual anomaly term (Fig. 2b & 2h). The impediment of VPD upon NPP is consistent with stomatal conductance models where VPD incurs a non-linear stomatal limitation which restricts CO₂ assimilation rates [36,38]. The inability of the seasonal forest model to isolate a consistent VPD anomaly effect could be due to the fact that the monthly range of VPD is far larger in seasonal forest sites (SM Fig. 6), and that the dry season anomalies would have to be very large in absolute units of kPa to significantly impact stomatal conductance, because the VPD reduction on stomata closure may have largely already been exerted (a visual diagram is shown in SM Fig. 7).

Both the seasonal forest and aseasonal wet forest models indicate that the effect of VPD (either seasonal or anomaly) is especially compounded with anomalies in short wave radiation. Although the effect of a short wave anomaly effect seems important across tropical forests, it appears to reduce NPP far more in seasonal forests than it does for aseasonal wet forests. Some caution is warranted with respect to ranking of the effects of the VPD, water deficit, and shortwave anomalies because these are correlated, and their relative importance could change with prediction error from the gridded climate products. Also despite not presenting an effect of temperature anomalies, the long-term increase in air temperature is increasing VPD and may also be pushing tree communities above their normal acclimated optimum temperatures for photosynthesis [31–33]. In combination, an El Niño event that reduces rainfall and increases VPD, temperature and shortwave radiation will likely work in conjunction to limit transpiration, increase leaf temperatures, and by extension reduce photosynthesis [33]. It is noteworthy that there is little evidence that positive shortwave anomalies increase NPP, as would perhaps been expected in aseasonal forests [60,61].

The effect of soil water deficit is negative upon woody growth, but this effect is less identifiable in the aseasonal wet tropics where soil water deficit seldom deviates from zero. CWD and MCWD have been highly effective metrics of water deficit in previous studies [11,62], but here we found TerraClimate’s water deficit estimates to offer greater predictive ability than (M)CWD. The Thornthwaite-Mather water balance model used to produce the water deficit estimates in the TerraClimate product may be more effective than our calculation of (M)CWD because its calculation of water deficit includes information on soil water holding capacity and infiltration, and calculates a runoff term. However all metrics of water deficit are likely hindered by both uncertainty in rainfall estimates, and the current state of high uncertainty around how tropical forest vary their rates of evapotranspiration both seasonally and interannually [63].

4.3 How much do El Niño events suppress tropical woody growth and what can this tell us about how tropical forests are likely to respond to climate change?

The pantropical model predicts pronounced declines in global tropical forest NPP over two strong El Niño events (8.3% in 1997/1998, and 9% in 2015/2016). The impacts were largest in the Americas (Fig. 5) highlighting the importance of Amazonia in dominating the global signal because it accounts for around half of total tropical forest area and is adjacent to the eastern Pacific warm anomaly during El Niño events. Insular SE Asia also has a substantial influence on the global anomaly, but Africa appears to have a negligible role as El Niño signals are weaker and less consistent there. The meteorological teleconnections caused by El Niño events are not spatially consistent across events [64]. Similar to other findings that have correlated tropical air temperatures and El Niño indices to atmospheric CO₂ growth rates [65,66], we have...
demonstrated that the variability of total woody production of the tropics can be well-predicted from the ENSO 3.4 index. We should note that our study period does not include a major stratospheric aerosol volcanic eruption, the last major one of which being that of Mt. Pinatubo in 1991, and some models suggest that such eruptions alter vegetation productivity through increasing diffuse light [67] (not tested as meteorological predictor in our analysis) which could weaken the correlation with ENSO. While \( \text{NPP}_{\text{stem}} \) is not necessarily a good proxy for overall gross primary productivity or net ecosystem exchange, as there are likely to be concurrent shifts in plant respiration and carbon allocation [7], a depression in \( \text{NPP}_{\text{stem}} \) still probably indicates ecophysiological stress imposed upon the ecosystem [11].

Our analysis is driven by growth responses to seasonal variation and interannual anomalies, whereas growth responses to short term variation in VPD and temperature may not be the same as long-term growth responses to secular shifts in these meteorological variables. It is possible that ecosystems acclimate to longer term shifts (either through within-individual acclimation within limits, or on longer timescales through turnover in community dominance). Our analysis also does not consider changes in demography, so shifts in either recruitment or mortality could either act to counterbalance or exacerbate the magnitude of our predictions. Finally additional environmental variables come into play, in particular the secular increase in atmospheric \( \text{CO}_2 \) which may boost productivity and increase water use efficiency. Nevertheless, our analysis does highlight the potentially important role of increasing temperatures and VPD. Changes in atmospheric water demand may be more important than changes in seasonal water supply in driving ecosystem water stress in the aseasonal wet tropics, and deserve more analytical attention. It is worth noting that the peak temperatures and VPDs experienced during the 2015/6 El Niño were higher than for the 1997/8 El Niño (SM Fig. 8), because of the long-term warming trend between these events. The baseline upon which each anomaly sits is consistently shifting towards a hotter, higher VPD atmosphere, pushing ecosystems into new climate space.

Moving forward, the predictions here need to be challenged so we encourage collection and development of similar seasonally monitored dendrometer band datasets that can be applied to the same stem-to-stand scaling techniques used here. It should also be possible to draw on a wide set of dendrometer data collected by unconnected studies (some in the grey literature) to improve the span of the dataset. Because these predictions deal with a specific component of ecosystem carbon, few empirical measures are available to test our model predictions. Ecosystem models still struggle to simulate realistic ecophysiological impacts from drought [68], while they also have vastly different approaches to carbon allocation that may produce unrealistic predictions [3,69–71]. Earth System Models typically represent the entirety of the tropical forest biome with a very few plant functional types. Our analysis highlights a key difference between seasonal and aseasonal wet forests in the underlying meteorological drivers that suppress woody growth during drought events. This message is consistent with Guan et al., (2015) [72] who highlighted different phenological and photosynthetic responses between tropical forests receiving more or less than 2000 mm yr\(^{-1}\) in precipitation, suggesting an important functional ecotone in the tropical forest biomes. The "empirical upscacling" spatiotemporal products developed from applying ensembles of machine learning models to global FluxNet data [73] have served as a benchmark of sorts to ecosystem models in recent years. However comparison to our \( \text{NPP}_{\text{stem}} \) predictions may not be straightforward because \( \text{NPP}_{\text{stem}} \) is a poor proxy for both GPP and total NPP in the wet tropics [3,7,16], and there are very few eddy covariance time series in the tropics outside of Brazil. Thus we reiterate the need for more collection of seasonally monitored tropical forest \( \text{NPP}_{\text{stem}} \) data, because the causal attribution of what drives variability in carbon allocation is still an emerging science. A logical next step is also to expand this analysis to other components of NPP and respiration, and thereby to total NPP and carbon balance. This will be the focus of our forthcoming analyses.

### Additional Information

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Data Accessibility
Stand-level NPP used in this study will be uploaded as supplemental material.

Authors’ Contributions

Competing Interests

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References


42. Zanne AE *et al.* 2009 Data from: Towards a worldwide wood economics spectrum. (doi:10.5061/dryad.234)


72. Guan K et al. 2015 Photosynthetic seasonality of global tropical forests constrained by hydroclimate. Nature Geoscience 8, 284–289. (doi:10.1038/ngeo2382)
Figure and table captions

Figure 1. The location of the Global Ecosystem Monitoring sites used in this study, overlaid on a map of mean annual precipitation.

Figure 2. (panels a & b) Coefficient plots for the Seasonal Forest NPP\textsubscript{stem} and Aseasonal Wet Forest NPP\textsubscript{stem} models with 50% and 90% credible intervals for the meteorologically driven statistical model. Abbreviations are as follows: SWmean, is the long term monthly mean of shortwave radiation, Tmean\textsubscript{μ}, is the long term monthly mean of temperature, VPDmean\textsubscript{μ}, is the long term monthly mean of vapor pressure deficit, VPDmean\textsubscript{anom}\textsubscript{3-mo}, is the moving 3-month mean moving anomaly of vapor pressure deficit, SWanom\textsubscript{3-mo} is the 3-month moving mean anomaly of shortwave radiation, Wet anom. and Dry anom. are the excessively wet and excessively dry parts of the water deficit anomaly. (panels c - h) The effect of the model terms are expressed on hypothetical conditional plots with median posterior prediction and 50 and 99% posterior predictive intervals in shaded colors. Apart from the model term that is varied along the x-axis, all other model terms in the conditional plots are set to the mean from the season or aseasonal forest data sets. All panels on the left correspond to the seasonal forest model, while panels on the right correspond to the aseasonal wet forest model.

Figure 3. Site level observations (open circles) and predictions (solid circles) with corresponding 50 and 99% prediction intervals of monthly NPP\textsubscript{stem} for individual plots located near (A) Kenia, Bolivia, (B) Tambopata, Perú, (C) Santarém, Brazil, (C) Tambopata, Perú, (D) Kogyae, Ghana, (E) Bobiri, Ghana, (F & G) regions in the east of Sabah, Malaysian Borneo, and (H) Jenaro Herrera, Perú.

Figure 4. The detrended Pantropical spatial anomalies of NPP\textsubscript{stem} during the El Niño events of 1997-1998 and 2015-2016, expressed Mg C ha\textsuperscript{-1} month\textsuperscript{-1}.

Figure 5. (Top) The 12-month detrended and running mean anomaly (expressed in Pg C yr\textsuperscript{-1}) of annual NPP\textsubscript{stem} (black) across the tropical regions and the Pannotropics. The vertical colored bars represent corresponding El Niño 3.4 index through time.

Table 1. Climatic characteristics of Global Ecosystem Monitoring regions used in this study. We divide the forest biomes as follows: WTF - wet tropical forest (>2200 mm), MTF - moist tropical forest (1800-2200), SDTF- semi-deciduous tropical forest (1400-1800 mm), and DTF - dry tropical forest (<1400 mm). Precipitation seasonality was calculated according to Feng et al., (2013), where a higher value indicates a more temporally concentrated distribution of annual rainfall.
## Tables

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Figures

Figure 1.
Figure 2.
Figure 3.
Figure 4.
Figure 5.