

Reply to Editors Report

Editors Report: Thank you for providing author comments in response to the referee comments. Both referees evaluated that your manuscript has good scientific quality and fair to good scientific significance and presentation quality. They acknowledged that your study topic is timely and fits to the journal scope, and recommended to make further data-model analyses with clearer explanations. The tree-size distribution model with a fewer number of parameters would help constrain DGVMs.

I studied the referee comments, your responses, and revised Introduction and Discussion and conclude that the manuscript would be adequately improved in clarity and insightfulness. I am looking forward to receiving the revised manuscript.

Authors' Response:

Thank you for your report and agreement with the changes we proposed in our response to the referee's comments (previously submitted, please refer to them for list of major changes). Please find in this submission the updated manuscript, supplementary material and in this document the tracked changes version of the manuscript compared to that submitted in July.

The changes made are extensive, as we had proposed, and note that Most figures have changes that cannot be shown via latexdiff tracked changes and also note that the figure 10 in the previous version has been cut, again this cannot be shown in the tracked changes, so we are highlighting these changes here. Also note the previous Table 1 (the height and mass allometry coefficients used to convert trunk diameter to mass) has been moved to the supplementary material.

We have also gone through the manuscript and tried to eliminate any remaining typographic errors. We now hope these have all been found and removed. If you or the journal editorial team have any further comments or queries do not hesitate to contact us.

Changes to Figures: -

Figure 1 – Corrected longitude spelling on x axis label.

Figure 4 swapped with figure 5

New Figure 4 – Added second panel b) showing the histogram of fitted μ_1 values for both 1 and 2-parameter fits. The original panel a) was rebinned to match the number of bins in the new panel b).

New Figure 5 – Compared to its previous form as the old figure 4 the only change is the addition of the line showing equation 26 fits the trend of the points relating the two fitting parameters.

Figure 6 – Restricted the fitting lines to the range of tree masses used in the fitting process. This is to more clearly show that trees with mass below m_p were excluded from the fit.

Figure 7 – Again added a line to show the fit of equation 26 to the relationship between fitting parameters.

Figure 8 - Added second panel b) showing the histogram of fitted μ_{m1} values for both 1 and 2-parameter fits. The original panel a) was rebinned to match the number of bins in the new panel b).

Figure 9 – Added extra panel for MST biomass and changed the x-axis label from "observed biomass" to "allometric biomass".

Figure 10 - Removed

Validation of demographic equilibrium theory against tree-size distributions and biomass density in Amazonia

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Abstract. ~~Understanding the relative abundance of trees of different sizes is an important part of predicting the response of forests to changes in climate, land-use and disturbance events. Two competing theories of forest size-distributions are demographic equilibrium theory (DET), based on scaling of mortality and growth with size, and metabolic scaling theory (MST), based scaling size with metabolic rates and how trees fill space. Recently, it was shown that for US forests DET is a much better model than MST, even using the same growth scaling with size. Studies comparing DET and MST have so far focused on trunk diameter, but tree mass and the associated forest mass per unit area (biomass density) are much more relevant to climate land-use change depends on models that can simulate the time-varying distribution of different tree-sizes within a forest - so-called *forest demography* models. A necessary condition for such models to be trustworthy is that they are able to reproduce the tree-size distributions that are observed within existing forests worldwide. In a previous study, we showed that Demographic Equilibrium Theory (DET) is able to fit tree-diameter distributions for forests across North America, using a single site-specific fitting parameter (μ) which represents the ratio of the rate of mortality to growth for a tree of a reference size. We use a form of DET that assumes tree-size profiles are in a steady-state resulting from the balance between a size-independent rate of tree-mortality and tree growth-rates that vary as a power law of tree size (as measured by either trunk diameter or biomass). In this study, we extend by fitting both DET and MST to mass data for the Amazon rainforest. The conversion via allometry from trunk diameter data to mass leads to an artefact in the mass distribution, which can be corrected by excluding smaller trees. We derive equations to calculate the total forest biomass density from the mass distribution equation, for both models, and these can be used as an indicator of goodness of model fit to the data. The models were fitted to the data, test DET against ForestPlots data for 124 sites across Amazonia, fitting, using Maximum Likelihood Estimation, at the forest plot, regional and continental scale. The fits for both diameter and mass demonstrate that MST is rarely a good fit for Amazon size-distributions and that DET is much better and can estimate biomass density, at the forest plot scale, with a mean error of 6% (10% if DET allometry fixed to MST) to both directly measured trunk diameter data and also biomass estimates derived from published allometric relationships. Again, we find that DET fits the observed tree-size distributions well, with best-fit values of the exponent relating growth-rate to tree mass giving a mean of its true value, compared to 139% for MST. The median of the fitted growth-scaling power for all the 124 plots is very close $\phi = 0.71$ (0.31 for trunk diameter). This is broadly consistent exponents of $\phi = 0.75$ ($\phi = 1/3$ for trunk diameter) predicted by Metabolic Scaling Theory (MST) allometry. The fitted ϕ and μ~~

parameters also show a clear relationship that is suggestive of life-history trade-offs. When we fix to the MST allometry values, implying MST allometry is a mean scaling, around which smaller forest plots cluster. At the larger regional scale, the error in the biomass density estimate of DET reduces to 2% or less and it is less than 1% for the whole continent value of $\phi = 0.75$. we find that best-fit values of μ cluster around 0.25 for trunk diameter, which is similar to the best fit value we found for North America of 0.22. This suggests that models based on DET, such as the relatively simple Robust Ecosystem Demography model (RED), are a good basis for a next-generation dynamic global vegetation model, and that Amazonian forests remain close to demographic equilibrium on large scales, despite climate change and significant anthropogenic disturbance. an as yet unexplained preferred ratio of mortality-to-growth across forests of very different types and locations.

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10 1 Introduction

The modelling of the abundances of various tree sizes in tropical forests such as the Amazon is important in efforts to improve understanding of land-climate feedbacks and hence anthropogenic climate change. This is because the fluxes of CO_2 between land and atmosphere are sensitive to climate (Feldpausch et al., 2016; Gatti et al., 2014) and so lead to possibly significant climate-land feedbacks, such as those predicted for the Amazon (Cox et al., 2000; Brienen et al., 2015).

15 Earth System models (ESMs) are used to model climate, but currently have a large range of uncertainty in the prediction of the land carbon sink, with as much as 500 GtC uncertainty by 2100 for 1% increase in CO_2 emissions per year (Friedlingstein et al., 2014). This has an impact on the predictions uncertainty feeds through into estimates of how much emissions need to be reduced to keep global warming within a certain level.

20 These issues have led to the development of more advanced Dynamic Global Vegetation Models (DGVMs), used within ESMs, to better more effectively represent vegetation processes (Sitch et al., 2015; Fisher et al., 2018). One of the key advances has been the inclusion of tree size-distributions, which allows better representation of land-use change and recovery from disturbance. These more

25 These recent DGVMs broadly consist of two different approaches based on representing size either via individual-based to representing tree size, either based on individual-based models (Shugart et al., 2018) or using cohort-based ecosystem demography models (Moorcroft et al., 2001; Longo et al., 2019).

30 DGVMs though DGVMs also need to balance additional complexity against practical considerations of usability, and computer execution time and memory usage. Key issues in the usability of complex numerical models are the understanding of the effect of many model parameters and the dependence on initial conditions (Moore et al., 2018). One, often overlooked, solution is simple analytical models that only represent key features of a numerical model, but allow a complete view of the effect parameters have on their behaviour. These analytical models can, therefore, be used to aid understanding of more complex models and also can have practical uses when validating models against data or using equilibrium solutions to aid

model initialisation. This approach has been used with the new DGVM called the Robust Ecosystem Demography model (RED, Argles et al. (2019), in preparation). The analytical equilibrium solution, as well as being used for initialisation, can be extended to allow RED to estimate implicit disturbance rates based on average vegetation coverage and growth. To solve these issues we have been exploring simplifications to the modelling of forest demography that are both parameter sparse and have steady-state solutions that can be solved for analytically (Moore et al., 2018; Argles et al., 2019).

Moore et al. (2018) compared simple trunk diameter size-distribution models to US forest inventory data. This approach showed that even in areas like the US that are not yet completely at equilibrium, the size-distribution could be modelled well on the larger scales that are important for DGVMs. We now extend this approach by firstly comparing various analytical models to forest inventory data for the Amazon, and secondly using mass as well as trunk diameter data. These are two key extensions, as the Amazon We follow Demographic Equilibrium Theory (DET) (Muller-Landau et al., 2006b) in assuming that forests are in a steady-state with size distributions completely determined by size-dependent functions of tree growth and mortality. Previously we showed that DET was able to fit the large-scale size-distributions of forests in North America (Moore et al., 2018), even though many of these forests are net carbon sinks (and therefore not in a precise steady-state). The current study uses the simplest reasonable form of DET that assumes growth is a power law of size and mortality is constant. This has been shown to be a useful model of underlying demographic processes, with the model parameters correlating with observations (Muller-Landau et al., 2006b; Lima et al., 2016), even though individual forest plots may deviate from the simplifying assumptions. While the growth and mortality functions of a forest are often unknown, DET can provide useful indications of the patterns of the ratio of mortality-to-growth based on observed tree-size distributions alone (Moore et al., 2018)

Amazonia is one of the largest pools of land carbon on the planet (Feldpausch et al., 2012) and using mass data allows this approach to test the ability of this modelling approach to accurately predict biomass density.

As in our previous study (Moore et al., 2018), we use the Demographic Equilibrium Theory (DET) model (Muller-Landau et al., 2006b) and the Metabolic Scaling theory (MST) model (West, 1997; West et al., 2009). MST has two assumptions of interest; that of how trees fill space and of the allometric scaling of tree growth rate with tree size. MST may be vulnerable to climate change (Cox et al., 2000; Brienen et al., 2015). It is therefore vital that DGVMs are able to model this region well. We therefore extend the analysis of Moore et al. (2018) by fitting DET model to tree trunk-diameter data for this key region, and also to tree mass data derived from allometry, which is even more relevant for ESMs. As a baseline comparison we also fit the Metabolic Scaling Theory of forest demography (MSTF), which assumes that trees of varying sizes fill space in such a way that the size-distribution scales with trunk diameter D as D^{-2} and that they have power-law growth rate scaling as $D^{1/3}$. Our previous study tested only the space-filling assumption as the DET model we used also assumed the same growth scaling as MST. In this study we test two versions of DET, one with the MST growth scaling and another where the assumed power-law scaling of growth has an exponent that is also allowed to vary as a fitting parameter. (West et al., 2009).

In Section 2 below we summarise the theoretical basis for DET and also MSTF, deriving analytical formulae for total forest biomass in each case. Section 3 describes the Methods and data, and Section 4 describes the results. Finally discussion and conclusions are in Sections 5 and 6.

So this study will test which of the models is the best choice for trunk diameter distributions, for mass distributions and for also for total biomass density prediction for the Amazon.

2 Theory

2.1 Demographic Equilibrium Theory (DET)

5 The distribution of tree sizes in a forest can be understood in terms of how the growth and mortality of the trees vary with tree size (Kohyama et al., 2003; Coomes et al., 2003; Muller-Landau et al., 2006b). ~~For~~ The amount of trees in a given size class (i.e. range of tree size) ~~, then trees smaller than that range will grow into it increasing the abundance. Conversely the abundance will decrease as trees in that range grow out or die~~ depends on the number of smaller trees growing into it and the number leaving it due to growing out or dying. The balance of growth and mortality will determine whether the abundance
10 of a size class is increasing, decreasing or if it is in demographic equilibrium (Van Sickle, 1977). ~~On~~ At the scale of a whole forest ~~scale~~, there is a further balance between the rate of seedling recruitment from seeds (lower boundary condition) and the whole forest mortality. Again this balance will determine if the forest as a whole is gaining or losing both mass and/or abundance.

The governing equation for this process is variously known as the ~~one-dimensional~~ one-dimensional drift or continuity
15 equation (Van Sickle, 1977), the Kolmogorov forward or the Fokker-Planck equation with the ~~second order~~ second-order term omitted (Kohyama, 1991) ~~and the Van Foerster equation (Von Foerster, 1959):~~

$$\frac{\partial n(D,t)}{\partial t} + \frac{\partial}{\partial D} (n(D,t)g(D,t)) = -\gamma(D,t)n(D,t) \quad (1)$$

where n is the size-distribution (tree density per size class) in Trees $\text{cm}^{-1} \text{ha}^{-1}$ in terms of tree trunk diameter D in cm, trunk diameter growth rate g in cm year^{-1} , γ is the mortality rate per year and time t in years.

20 It was shown (Kohyama et al., 2003) that for an unchanging, equilibrium size-distribution, this equation can be integrated as follows :-

$$\int_{n_L}^n \frac{dn}{n} = \int_{D_L}^D \frac{1}{g(D)} \left[\frac{dg(D)}{dD} + \gamma(D) \right] dD \quad (2)$$

where n_L is the value of n at the lower boundary D_L , which for forest inventory data is the minimum sampling size (in this study 10 cm).

25 This equation can be solved to give an exact solution ~~for~~, if simplifying assumptions of size-independent mortality $\gamma(D) = \gamma$ and power law growth rate $g(D)$ are used. The growth rate $g(D)$ in cm per year is then

$$g(D) = g_1 D^\phi \quad (3)$$

where g_1 is a constant with the same value as the growth rate for a tree with trunk diameter of 1 cm. The solution (Muller-Landau et al., 2006b; Lima et al., 2016; Moore et al., 2018) for the size distribution is then the Left-Truncated Weibull Distribution (LTWD)

$$n(D) = n_L \left(\frac{D}{D_L} \right)^{-\phi} \exp \left[\frac{\mu_1}{1-\phi} \left(D_L^{1-\phi} - D^{1-\phi} \right) \right], \phi \neq 1 \quad (4)$$

- 5 where $\mu_1 = \gamma/g_1$ is the mortality to growth ratio at $D=1$ cm (Note: the units of μ_1 are $\text{cm}^{\phi-1}$ but as it is defined for the point $D = D_1 = 1$ cm can be assumed to be dimensionless if the size variable D is implicitly a ratio D/D_1 , which has the same exact numerical value as D but is dimensionless).

This solution is also applicable for other size variables such as tree dry mass m in kg :-

$$n(m) = n_L \left(\frac{m}{m_L} \right)^{-\phi_m} \exp \left[\frac{\mu_{m,1}}{1-\phi} \frac{\mu_{m1}}{1-\phi} \left(m_L^{1-\phi_m} - m^{1-\phi_m} \right) \right], \phi_m \neq 1 \quad (5)$$

- 10 where m_L , ~~m_L~~ , μ_{m1} and ϕ_m are the mass equivalents of D_L , μ_1 and ϕ .

The LTWD distribution has been shown to be a good description of tree trunk diameter distributions in a variety of tropical forests (Muller-Landau et al., 2006b; Lima et al., 2016) and in temperate forests in the US over larger scales (Moore et al., 2018). When these distributions are fitted to data then they can have both parameters ϕ and μ_1 as fitting parameters or just fit μ_1 and fix ϕ to the values used in MST allometry (Niklas and Spatz, 2004; West et al., 2009) of $\phi = 1/3$ and $\phi_m = 3/4$.

15 2.2 Total Biomass Density for DET

The total biomass density (kg of dry tree mass per hectare) of the LTWD tree mass distribution can be obtained by integrating Eq. (5) in terms of mass, between the lower boundary m_L and infinity :-

$$M_{L \rightarrow \infty} = \int_{m_L}^{\infty} m n(m) dm = n_L m_L^{\phi_m} \frac{\exp(x \mu_{m1} m_L^{1/x})}{\mu_{m1} (x \mu_{m1})^x} \Gamma(x+1, x \mu_{m,1} m_L^{1/x}) \quad (6)$$

where Γ is the upper incomplete Gamma Function, $x = 1/(1 - \phi_m)$.

- 20 As real forests do not satisfy the assumption of infinite maximum size tree, this can lead to errors in the calculated biomass density. A correction to this can be found, in terms of m_{\max} the largest tree mass in the distribution :-

$$M_{L \rightarrow m_{\max}} = \int_{m_L}^{m_{\max}} m n(m) dm = \int_{m_L}^{\infty} m n(m) dm - \int_{m_{\max}}^{\infty} m n(m) dm \quad (7)$$

In cases where m_{\max} is both large and much larger than m_L then there will be little difference between Eq. (6) and Eq. (7). m_{\max} is a somewhat arbitrary function of the sample size, due to large trees being statistically rare, meaning the infinite upper

- 25 bound solution Eq. (6) is expected to perform better be more accurate for larger sample sizes.

2.3 Metabolic Scaling Theory (MST)

Metabolic scaling theory is a theory of scaling of organisms with size, based on theories of metabolism, physics and chemistry (West, 1997; Muller-Landau et al., 2006a). This theory uses the predictions of the scaling of individuals to predict the larger scale patterns and structure of populations and communities. For forests this is in the form of using the scaling of photosynthesis of trees and the vascular structures that transport water to predict individual scaling. This is then combined with assumptions [from self-thinning](#) about how trees fill space to describe the expected forest size-distribution ([West et al., 2009](#)) ([Coomes et al., 2003](#); [West et al., 2009](#)). This leads to a power law distribution for trunk diameter \dashv

$$n(D) = n_L \left(\frac{D}{D_L} \right)^{-2} \quad (8)$$

and for mass the distribution is almost identical

$$n(m) = n_L \left(\frac{m}{m_L} \right)^{-11/8} \quad (9)$$

2.4 Total Biomass Density for MST

The MST equations also enable the calculation of biomass density (kg of dry tree mass per hectare). In this case only the finite upper bound of m_{\max} can be used as the solution goes to infinity as the upper bound goes to infinity.

$$M_{L \rightarrow \max} = \int_{m_L}^{m_{\max}} m n(m) dm = \frac{8n_L m_L^{11/8}}{5} \left[m_{\max}^{5/8} - m_L^{5/8} \right] \quad (10)$$

15 3 Methods

3.1 Forest inventory data

The tree census data used in this study is from the public access permanent sample plots of the RAINFOR (Peacock et al., 2007) network. RAINFOR provides a systematic framework for long-term monitoring of the Amazon. The RAINFOR data is stored on the ForestPlots database (<https://www.forestplots.net>). This database stores measurements (stem diameter, species ID, recruitment, growth, and mortality) of individual trees from hundreds of locations, taken using standardised techniques to allow the behaviour of tropical forests to be measured, monitored and better understood (Lopez-Gonzalez et al., 2011).

We selected 124 open access forest plots (Fig. 1) classified as mixed forest ([not monoculture](#)) and old-growth to most closely match the model assumptions of forests undisturbed by human interference and approximating to equilibrium demography. ~~All measurements below a trunk diameter of~~ [The 124 selected plots all had a consistent lower cut-off in measurements at 10 cm trunk diameter](#) ~~were excluded giving consistent left-truncation point to the distributions of~~. [Two available upper montane plots](#)

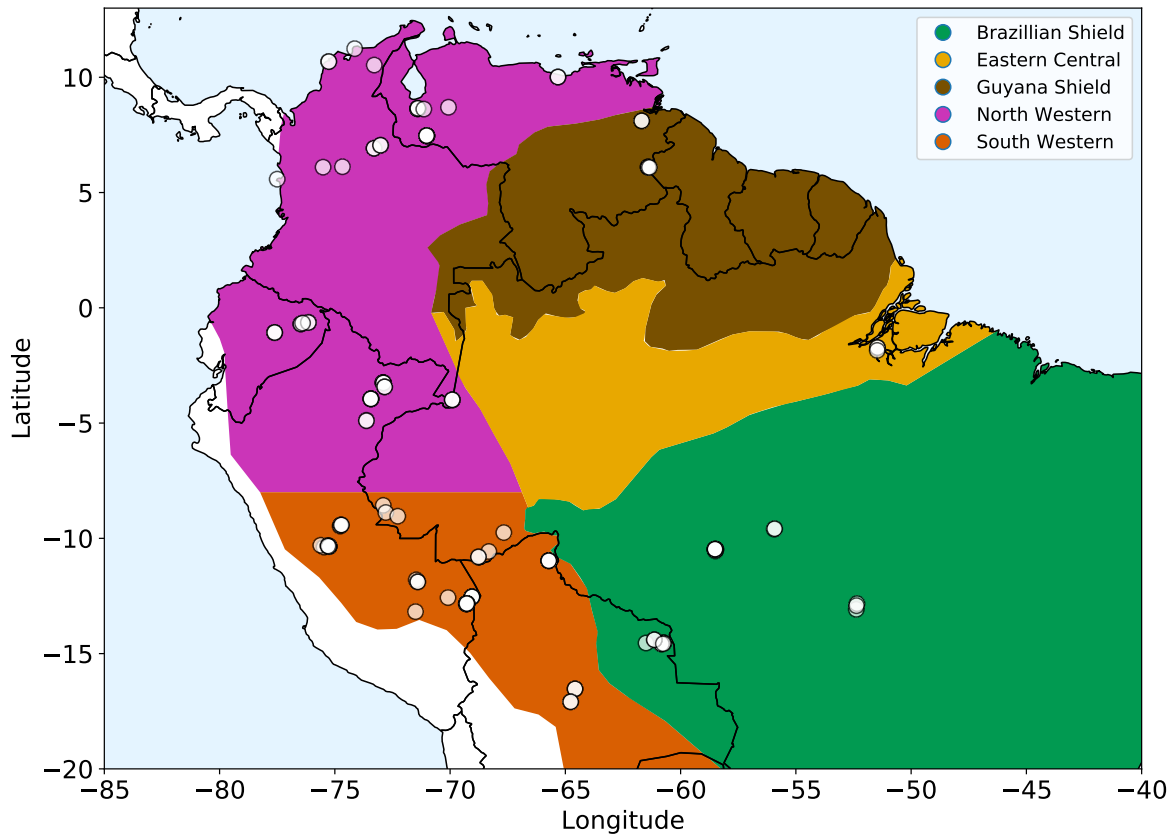


Figure 1. Amazonian Allometric Regions. Each region, shown by the coloured areas, is defined by geography, rainfall and soil substrate. White circles show location of the forest plots used. The two western regions share common allometry but are split based on rainfall seasonality for analysis purposes.

~~with very few measurements above 10 cm. Plots with a large proportion of measurements below 10 cm were not included in the 124 selected plots used, as they did not have enough measurements to allow a reliable fit.~~

3.2 Calculating Dry Tree Mass from Trunk Diameter

The open access plots of the Amazon RAINFOR dataset consists only of trunk diameter values. To estimate the tree mass, the methodology developed by Feldpausch et al. (2012) was used. In that study two functional forms (with and without height) were tested against destructively sampled mass data (trees carefully measured then cut down and weighed) to find ones which best estimated mass from trunk diameter. It was found that mass estimation accuracy doubled when including height, even if the height had in turn been estimated from trunk diameter. Out of three choices of height functional form (power law, Weibull-H and exponential), Feldpausch et al. (2012) found ~~to~~ the Weibull-H form Eq. (11) to be the best at estimating mass across multiple size classes. The height H in metres is then ~~:-~~

$$H = a_h(1 - \exp(-b_h D^{c_h})) \quad (11)$$

with the coefficients varying geographically between defined allometric regions (see Table ~~??~~ [S1 in Supplementary Material](#) and Figure 1).

The regions were defined by geography and substrate origin (Feldpausch et al., 2012). Western Amazonia (Columbia, Ecuador and Peru) being recently weathered Andean deposits, the geologically old Brazilian Shield to the south (Bolivia and Brazil), Guyana Shield on the northern side of the Amazonia Basin (Guyana, French Guyana and Venezuela) and Eastern Central Amazonia (Brazil) consisting of sedimentary substrates originating from the other regions. The western region from Feldpausch et al. (2012) was split along latitude of -8 degrees based on rainfall seasonality (Fauset et al., 2015). These two western [regions](#) still retain a common height allometry but are split for analysis.

~~Coefficients for Eq. (11) from Feldpausch et al. (2012) Region a_h b_h c_h All S.America 42.574 0.0482 0.8307 Western Regions 46.263 0.0876 0.6072 Brazilian Shield 227.35 0.0139 0.555 Guyana Shield 42.845 0.0433 0.9372 Eastern-Central 48.131 0.0375 0.8228~~

The mass function (kg [of tree dry mass](#)), when height was included as one of the parameters used was ~~:-~~

$$M = e^a (\rho_w D^2 H)^b \quad (12)$$

where the parameters are universal across all regions with values $a = -2.9205$ and $b = 0.9894$. The function was from Feldpausch et al. (2011) and the parameters were estimated in Feldpausch et al. (2012).

The wood specific gravity ρ_w was obtained from the Dryad Global Wood Density Database <https://doi.org/10.5061/dryad.234/1> (Chave et al., 2009; Zanne et al., 2009). For each tree measurement the ρ_w value used was for that species from the closest available region. Where the species data was unavailable or the species of the measurement had not been recorded then the ρ_w value of the Genus was used, based on an average of all trees in the Dryad database in that Genus. Trees without Genus data

were estimated from Family data, and any remaining measurements where the ρ_w was still unknown were set to the average ρ_w of the trees in that same forest plot with known ρ_w values.

3.3 Fitting methodology

As in our previous study (Moore et al., 2018), Maximum Likelihood Estimation (MLE) was used to find the parameters that give the best fit for both the Left-Truncated Weibull, derived from DET (DET-LTWD) and Metabolic Scaling Theory (MST) distributions. MLE is an effective method for parameter fitting of forest size distributions (Taubert et al., 2013; White et al., 2008).

Maximising the log-likelihood L results in a more numerically tractable summation of terms rather than a product of terms obtained from using the Likelihood directly. L in terms of $f(D)$ the probability distribution function (pdf) $f(D)$ is then

$$L = \sum_i \ln(f(D_i)) \quad (13)$$

where D_i is tree trunk diameter data point measurement of stem i in the dataset.

The data was fitted both by plot, by allometric region (an aggregated dataset of all plots in that region), by country (again aggregation of plots) and for all the data, from all 124 plots, grouped together as one large dataset. This allows both the study of the individual plots and the larger scale patterns across South America.

3.4 Maximum Likelihood Estimation (MLE) for Demographic Equilibrium Theory (DET)

The probability density function (pdf) $f(D)$ for the DET-LTWD, in terms of tree trunk diameter D and minimum tree size D_L is related to the number density distribution $n(D)$ (Eq. 4) :-

$$f(D) = \frac{A}{N} n(D) = \mu_1 D^{-\phi} \exp \left[\frac{\mu_1}{1-\phi} \left(D_L^{1-\phi} - D^{1-\phi} \right) \right], \text{ for } \phi \neq 1 \quad (14)$$

where N is the total number of trees in the dataset being fitted, ϕ is the growth scaling power from Eq. (3) and A the area of the plots containing the trees sampled in the dataset. This equation is equivalent to the standard form of the LTWD

$$f(D) = \frac{c}{\lambda} \left(\frac{S D}{\lambda \lambda} \right)^{c-1} \exp \left[\left(\frac{D_L}{\lambda} \right)^c - \left\{ \frac{D}{\lambda} \right\}^c \right] \quad (15)$$

where $c = 1 - \phi$ is the shape parameter and $\lambda = \left[\frac{c}{\mu_1} \right]^{1/c}$ the scale parameter.

We fit DET-LTWD twice, once with both parameters ϕ and μ_1 allowed to vary as fitting parameters and secondly with the growth scaling parameter ϕ fixed to the MST allometry values ($\phi = 1/3$ and $\phi_m = 3/4$, see Niklas and Spatz (2004) and West et al. (2009)). Fixing ϕ means we have a DET-LTWD model following just one assumption of MST (the allometry) and so acts as way of comparing the effect of the second MST assumption of space-filling when comparing DET-LTWD and MST fits.

3.4.1 One Parameter Fit

For this situation, where we are only aiming to find the parameter μ_1 and ϕ is assumed, then MLE can be solved analytically (Kizilersu et al., 2016)

$$\mu_1 = \frac{c}{(\bar{D}^c - D_L^c)} \quad (16)$$

5 where $c = 1 - \phi$. The equations are the same for tree mass, just with the symbols appropriately substituted (m for D etc).

3.4.2 Two Parameter Fit

For the two parameter case, where both ϕ and μ_1 are fitted, then we calculate the Log-Likelihood L as follows

$$L = N \left(\ln \mu_1 + \mu_1 \frac{D_L^c}{c} \right) - \frac{\mu_1}{c} \sum_i D_i^c + (c-1) \sum_i \ln D_i \quad (17)$$

10 Substituting Eq. (16) into Eq. (17) creates a function only of c and therefore ϕ . This allows minimisation of $-L$ in terms of ϕ by using Brent's bounded algorithm (Brent, 1973). Once the ~~parameters optimum~~ ϕ has been found then μ_1 can be calculated from equation 16. As equation 16 is included in the minimisation of $-L$, then it means we are in fact solving for both parameters at once and are finding the maxima of L . This algorithm was tested both with real data and data generated by computer from known LTWD distributions, by plotting the L values against ϕ and μ_1 that gives the maximum L ., to confirm the maxima was found correctly.

15 Once the parameters μ_1 and ϕ are estimated, then this allows n_L , the tree density per size class at D_L , to be obtained from these parameters and the known quantities of the total number of trees N and the plot area A , ~~which are known from the data.~~ This can be derived by integrating the equation for n (Eq. 4), to give \therefore

$$\frac{N}{A} = \int_{D_L}^{D_{\max}} n(D) dD = \frac{n_L D_L^\phi}{\mu_1} \left[1 - \exp \left[\frac{\mu_1}{c} (D_L^c - D_{\max}^c) \right] \right] \quad (18)$$

and noting that the observed number of trees is identical to the integral, we get \therefore

$$20 \quad n_L = \left(\frac{N}{A} \right) \frac{\mu_1}{D_L^\phi} \frac{1}{1 - \exp \left[\frac{\mu_1}{c} (D_L^c - D_{\max}^c) \right]} \quad (19)$$

where $c = 1 - \phi$ and D_{\max} is the largest tree size in the dataset. For this study it was found that as $D_{\max} \gg D_L$ for most cases (and that c is never much larger than μ_1), n_L ~~could~~ can be assumed to be \therefore

$$n_L \approx \left(\frac{N}{A} \right) \frac{\mu_1}{D_L^\phi} \quad (20)$$

Again, the equations are the same for tree mass, just with the symbols appropriately substituted (m for D etc).

3.5 Maximum Likelihood Estimation (MLE) for Metabolic Scaling Theory

From the equation for number density n (Eq. 8) the pdf for MST is

$$f(D) = n(D) \left(\frac{A}{N} \right) = \frac{D_L}{\left[1 - \left(\frac{D_L}{D_{\max}} \right) \right]} D^{-2} \quad (21)$$

where D_{\max} is the largest tree size in the dataset. As all the quantities are known then there are no free parameters to fit and all that needs to be done is calculate n_L , the tree density per size class at D_L :-

$$n_L = \frac{(N/A)}{D_L \left[1 - \left(\frac{D_L}{D_{\max}} \right) \right]} \quad (22)$$

Similarly the MST pdf for mass from Eq. (9) is :-

$$f(m) = n(m) \left(\frac{A}{N} \right) = \frac{3m_L^{3/8}}{8 \left[1 - \left(\frac{m_L}{m_{\max}} \right)^{3/8} \right]} m^{-11/8} \quad (23)$$

and for n_L :-

$$n_L = \frac{3(N/A)}{8m_L \left[1 - \left(\frac{m_L}{m_{\max}} \right)^{3/8} \right]} \quad (24)$$

3.6 Estimating Plot and Regional Biomass Density

To test the biomass density equations, we used the results of the MLE fits to calculate the biomass density predicted by Eq. (7) and Eq. (10). The biomass density predicted by these equations are then compared to the ~~observed~~-allometric biomass density (i.e. the sum of the mass of all trees in a dataset divided by the area of the plots). This comparison then provides a goodness of fit measure that is relevant to climate.

We chose to measure the biomass density as a function of size in terms of the total mass per unit area from trees with masses equal or greater than a given size. The main reason for this is that the forest plot data only sampled trees ~~which have with a~~ trunk diameter equal to or greater than 10 cm. Therefore it makes little sense to measure the biomass density below a given size, as would be the case with a traditional cumulative distribution function. This approach has a second benefit that the mass of a forest above a given size is a much more useful way of easily seeing the contribution of the dominant larger trees to total biomass (Bastin et al., 2018).

A correction ~~terms-term~~ is added to Eq. (7) and Eq. (10) to make sure the biomass density correctly evaluates at the upper boundary (the mass of the largest tree m_{\max}). ~~The biomass density of trees equal or greater than m_{\max} should be m_{\max}/A ,~~

where A is the total area of plots. This is because these equations only evaluate the mass up to but not including the trees with a mass equal to the largest value in the dataset. As when the second term in Eq. (7) is included the biomass density of trees equal or greater than m_{\max} . Therefore, to comply with the definition above it is necessary to add the mass of the largest trees back into the total biomass.

- 5 As the large trees are so rare this correction will be equivalent to adding just one tree of the largest mass m_{\max} is zero, so we need to add a correction term of m_{\max}/A in the dataset divided by A , the total area of plots in the dataset.

$$M_{L \rightarrow \max} = n_r m_r^{\phi_m} \frac{\exp(x \mu_{m1} m_r^{1/x})}{\mu_{m1} (x \mu_{m1})^x} \left[\Gamma(x+1, x \mu_{m1} m_L^{1/x}) - \Gamma(x+1, x \mu_{m1} m_{\max}^{1/x}) \right] + \frac{m_{\max}}{A} \quad (25)$$

This Eq. (25) is used for all biomass density estimates where the upper bound of tree size is assumed finite (based on m_{\max}), while for the cases where the simplifying assumption of infinite tree size is used then Eq. (7) is used.

10 4 Results

4.1 Mass Distribution

When the mass data was estimated from the trunk diameter measurements using the methodology of Feldpausch et al. (2012), it was noticed that the mass size-distribution (for all regions and plots) had a peak, which was not present in the trunk diameter distribution. We found this to be an artefact of the conversion from trunk diameter to mass in a distribution that was by definition truncated already in trunk diameter.

15 Fig. 2a shows the relationship between trunk diameter and tree mass for the whole dataset, illustrating that for any particular trunk diameter there is a range of tree masses. This variation to mass is in tree mass is caused by the differences in wood density between species and the variation in height allometry between regions (see Eq. 11 and Table 1 and supplementary Table S1). If instead the dataset shown in Fig. 2a is truncated in mass rather than trunk diameter, then the truncation would instead follow the horizontal dotted line and there would be data in the region between that line and the diagonal dotted line. So in effect there is “missing” data for low mass trees, which is a result of the trunk diameter observations having a minimum sampling size (truncation point) and there being a range of tree masses for trees with a given trunk diameter. This hypothesis is further confirmed by increasing the trunk diameter truncation point, as shown in Fig. 2b. As the truncation point is increased the “peak” moves to higher mass.

25 4.1.1 Eliminating the Mass Peak

When working with mass data the peak was eliminated from fitting by creating 40 bin edges (39 bins) in log-space (base e) from the smallest to largest tree in the dataset. These edges define the range of each bin and the value of each bin was selected as the midpoint in log-space. The data was then binned following these bins. Once the data was binned, the bin with the highest frequency was identified. The value of this bin was then used as the truncation point for the dataset when fitting to the dataset

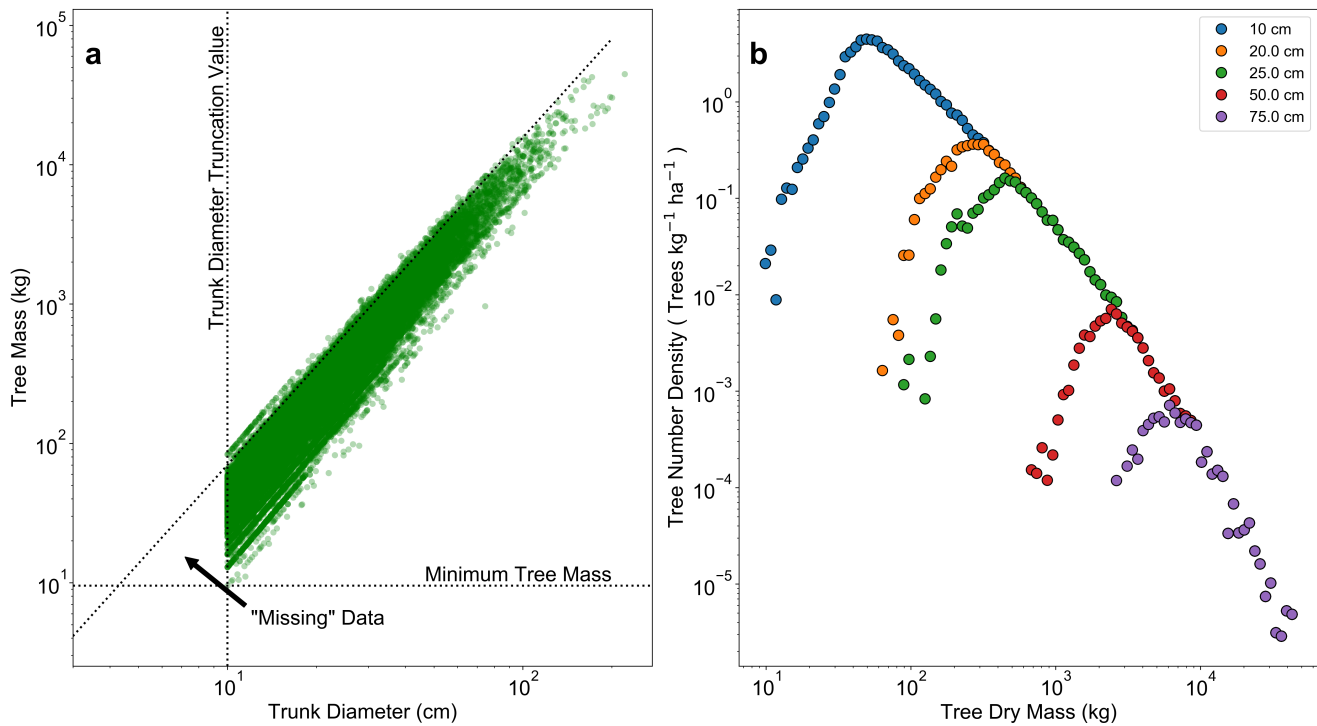


Figure 2. The effect of truncating data measured in trunk diameter and then converting to mass using allometry. In **a**), the mass for each tree [is shown](#) in terms of its trunk diameter. If the data had been truncated [to-in](#) mass there would be data in the triangle marked by the intersection of the dotted lines. This truncation effectively leads to missing data in the mass distribution, as seen in **b**). The mass distribution should constantly decrease with increasing mass but instead rises to a peak then decreases, and is due to incomplete data for the low mass end of the distribution. This peak can be seen to be an artefact of the trunk diameter truncation point. When the trunk diameter truncation point [increases-is increased](#) the mass distribution peak moves with the truncation point.

[distribution. The binning was purely used to identify the peak and for plotting the data and not used during the MLE fitting process.](#)

4.2 Trunk Diameter Results

Fitting the DET-LTWD and MST equations to the trunk diameter size distributions, showed a consistent pattern for all the geographical aggregations of plot data. In all cases, except Guyana shield, the DET-LTWD solutions (both one and two parameter versions) ~~fitted much better~~ more closely captured the curvature of the observed size-distribution than the MST solution (Fig. 5 3a and see supplementary material Fig. S1 and S2). In particular the MST model deviated from the observed data at large trunk diameters. Guyana Shield region only had four small plots, totalling 819 trees, which may explain the reason it was hard to distinguish visually the best fitting model ~~-(Supplementary Fig. S2).~~

The two parameter DET-LTWD fits gave a fitted value of the growth scaling power ϕ between 0.137 and 0.546 (Table 1) and five of the twelve regions were within 0.05 of the theoretical value of 1/3 (i.e. ϕ in range 0.28-0.38).

Table 1. Results of fitting ~~the models to~~ of the trunk diameter size-distributions for the forest plot data aggregated to regions, countries and ~~the whole continent~~ all plots combined. ~~Table~~ This table presents the fitted parameters for each model. ~~n_L is the value of the distribution at the lower truncation point of 10 cm trunk diameter.~~ μ_1 and ϕ are the model parameters from Eq. 3 fitted to the data by MLE. The one parameter DET model has $\phi = 1/3$, so only the fitted μ_1 parameter is given in the table.

Region	No.Trees	Area(ha) ha	mean D(cm) cm	DET 1 Param		DET 2 Param			MST
				μ_1	n_L $(\text{cm ha})^{-1}$	μ_1	ϕ	n_L $(\text{cm ha})^{-1}$	n_L $(\text{cm ha})^{-1}$
All S.America	63605	113.4	20.45	0.255	66.53	0.308	0.397	69.13	58.77
Brazil	12454	23.5	20.83	0.247	60.87	0.266	0.358	61.78	55.83
Bolivia	8963	16.0	20.11	0.265	68.93	0.491	0.546	78.15	59.49
Colombia	7288	13.2	19.68	0.273	69.87	0.314	0.382	71.84	58.31
Ecuador	4949	7.8	20.37	0.257	75.82	0.330	0.419	79.79	67.37
Peru	27080	44.5	20.38	0.256	72.41	0.281	0.366	73.81	63.74
Venezuela	2871	5.3	22.55	0.217	54.95	0.204	0.313	54.20	57.71
Amazonian Allometric Regions									
N.Western	22642	37.8	20.21	0.261	72.55	0.325	0.409	75.85	63.34
S.Western	24690	42.5	20.58	0.252	67.98	0.263	0.348	68.59	60.85
Brazilian Shield	13412	24.5	20.10	0.264	67.21	0.399	0.476	73.07	58.18
Guyana Shield	819	1.5	22.74	0.214	54.10	0.120	0.137	47.54	61.78
Eastern-Central	2042	4.0	22.90	0.213	50.40	0.212	0.332	50.34	53.73

10

In general the one and two parameter DET-LTWD solutions were quite similar in terms of the appearance of the fit on the distribution plots. This finding was confirmed using the Akaike ~~information criterion~~ Information Criterion (AIC) and Bayesian ~~information criterion~~ Information Criterion (BIC) (Table 2). Both the AIC and BIC are a way of determining from

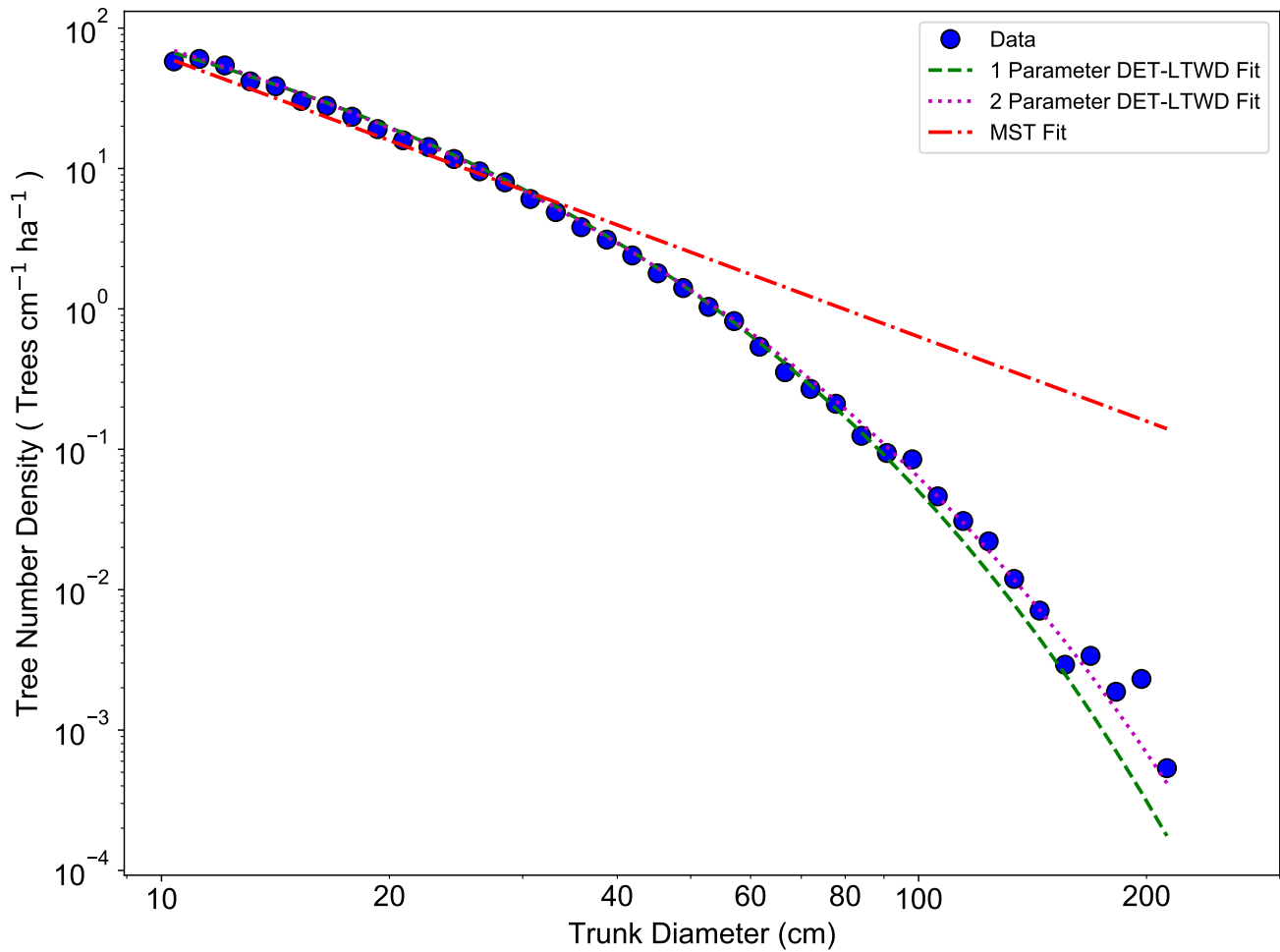


Figure 3. Fit to the trunk diameter size-distribution for all South American RAINFOR Plots as one large dataset. The blue circles show the binned data and the lines show the fitted distribution for each model.

several models which ~~represents the data better~~ has the best “goodness of fit”, with a lower value indicating a better fit. Both ~~criteria~~ criteria are calculated from the log likelihood and number of fitting parameters with a difference of 10 being the threshold where the evidence is considered to be very strongly against the higher scoring model (Kass and Raftery, 1995). BIC penalises a higher number of fitting parameters more than AIC.

- 5 It was only possible to distinguish the quality of the fits for four of the twelve geographical aggregations of forest plots. In all four cases (All S.America, Bolivia, Brazilian Shield and N.Western) the two parameter DET-LTWD fit was favoured and for the other eight it was not possible to say that the inclusion of the growth scaling power as a fitting parameter improved the fit.

Table 2. Model comparison for fits to trunk diameter size-distributions. ~~Table~~ This table shows the log Likelihood of each ~~models~~ model's fit and the corresponding AIC and BIC model comparison criterion. The best model has the lowest AIC or BIC; here the difference is shown to the best model, meaning the best model has a score of 0. Models other than the best are strongly rejected if they have a value greater than 10. ~~Best~~ Th best model and those not rejected are shown in bold.

Region	log Likelihood			Δ AIC			Δ BIC		
	MST	DET	DET	MST	DET	DET	MST	DET	DET
		1 Param	2 Param		1 Param	2 Param		1 Param	2 Param
All S.America	-218,530	-211,726	-211,699	13700.0	51.6	0.0	13600.0	42.5	0.0
Brazil	-43,146	-41,934	-41,933	2420.0	0.0	0.404	2410.0	0.0	7.83
Bolivia	-30,243	-29,433	-29,389	1710.0	87.3	0.0	1690.0	80.2	0.0
Colombia	-24,577	-23,715	-23,714	1720.0	1.57	0.0	1720.0	0.0	5.32
Ecuador	-16,889	-16,428	-16,424	927.0	5.82	0.0	915.0	0.0	0.682
Peru	-93,037	-90,049	-90,046	5980.0	3.38	0.0	5970.0	0.0	4.83
Venezuela	-10,289	-10,098	-10,098	379.0	0.0	1.67	373.0	0.0	7.63
Amazonian Allometric Regions									
N.Western	-77,148	-74,830	-74,817	4660.0	25.9	0.0	4640.0	17.8	0.0
S.Western	-85,245	-82,584	-82,583	5320.0	0.0	0.883	5310.0	0.0	9.0
Brazilian Shield	-45,391	-44,107	-44,077	2620.0	57.5	0.0	2610.0	50.0	0.0
Guyana Shield	-2,901	-2,898	-2,895	7.41	4.76	0.0	0.0	2.06	2.01
Eastern-Central	-7,370	-7,232	-7,232	274.0	0.0	2.0	268.0	0.0	7.62

4.3 Trunk Diameter Results for Individual Plots

Fitting the models to the individual forest plots (full results in supplementary material Tables S2 and S3 and Fig. S5 to S13) again resulted in the DET-LTWD models ~~fitting much better~~ generally fitting much more closely than MST. Table 3 shows the results of BIC comparison of the models for the 124 forest plots. In every case, the best model is determined by the lowest BIC value. Inferior models are only considered strongly rejected if their BIC is greater than the best model by 10 or more. This is represented by the columns in the table and shows the one parameter DET-LTWD was the ~~best model by far the most model~~ most commonly favoured by BIC score (81 plots). However, in none of those plots was it possible to strongly reject both the other models. The most common result (75 plots) was of the one parameter DET-LTWD being the best model, MST being rejected but the two parameter DET-LTWD also so closely fitting the data that it cannot be rejected. The next most common result (17 plots) was the reverse with again MST rejected but the two parameter DET-LTWD now narrowly better but not sufficient to strongly reject the one parameter DET-LTWD. The MST model was the best model for 15 plots and for 5 of those (ELD_01, ELD_02, RIO_01, RIO_02, TIP_03) the two DET-LTWD models were both strongly rejected. Four of these plots though had a very low number of trees, so it would be less likely to be able to pick a model with as much confidence from a distribution of only ~100 trees. In fact the MST model seemed more likely to have a favourable AIC or BIC score, compared to the other models, for plots with smaller sample sizes and an increasingly unfavourable score for higher sample sizes (see Supplementary Fig. S30).

Table 3. Shows the best and acceptable models for the 124 individual forest plots for trunk diameter. Models are labelled as (M) for MST, (1) for one parameter DET-LTWD and (2) for two parameter DET-LTWD. Columns refer to best fitting model (lowest BIC score). Rows refer to models that are so good a fit compared to the best that they cannot be rejected, as their BIC score is so close to the best model. For example '1M' means the MST and one parameter models are not rejected but the two parameter model is rejected based on BIC. Then the columns in this row show how many forest plots have either (1) or (M) model as best fit and the other also fitting closely.

Comparable Models	Best Model			Total
	1	2	M	
1	0	-	-	0
2	-	8	-	8
M	-	-	5	5
12	75	17	-	92
1M	0	-	2	2
2M	-	1	1	2
12M	6	2	7	15
Total	81	28	15	124

Fig. 5 shows the effect of fitting with the two parameter DET-LTWD model. There is to be a clear relationship between ϕ and μ_1 , as all results follow a curve. The black dotted vertical line shows the $\phi = 1/3$ value expected according to the MST allometry and also assumed in the one parameter DET-LTWD model.

Results of the two parameter DET-LTWD MLE fits for trunk diameter data from all 124 individual forest plots. The fitted mortality to growth ratio μ_1 is shown as a function of the fitted growth scaling power ϕ . The results from the fits to the grouped datasets of the four allometric regions are plotted as the dashed crosses of the corresponding colour. The vertical black line shows the ϕ value predicted by MST allometry.

Plotting just the ϕ results in a histogram (Fig. 4a), reveals an approximate bell-shaped distribution with a peak around close to the theoretical MST value. The median of the ϕ value for the plots is 0.34 (95% confidence interval 0.29-0.40) and the mean is 0.31 (95% confidence interval 0.26-0.36). These values are close to the theoretical value of 1/3, as suggested by the histogram. The histogram of μ_1 (Fig. 4b) shows a skewed bell-shaped distribution with a peak around 0.3 for 2-parameter DET-LTWD and a more symmetric bell curve centred around 0.25 for 1-parameter DET-LTWD. For 1-parameter DET-LTWD the the median of μ_1 for the plots is 0.25 (95% confidence interval 0.24-0.26) and the mean is 0.25 (95% confidence interval 0.24-0.26). For 2-parameter DET-LTWD the median of μ_1 for the plots is 0.27 (95% confidence interval 0.22-0.31) and the mean is 0.31 (95% confidence interval 0.26-0.35). The 1-parameter DET-LTWD mean and median μ_1 are very close to the value of 0.22 found when 1-parameter DET-LTWD was fitted to US Forest Inventory data (Moore et al., 2018) (Note in that study the fitted value of $\mu = 1.198$ was obtained for $D = 12.7$ cm, this has then be converted, by extrapolation, to the value at $D = 1$ cm to get μ_1 , this value is 0.22).

Fig. 5 shows the effect of fitting with the two parameter DET-LTWD model. There is a clear relationship between ϕ and μ_1 , as all results follow a curve.

If it is assumed that for any fixed value of ϕ there is a μ_1 value that gives the best fit for that (as can be seen in Fig. 3b) then an equation can be derived (see supplementary material Section 2) in terms of the DET theory and the known global best fit values ϕ_t and μ_{t1} (ie the values fitted to all plots together)

$$\mu_1 = \frac{1 - \phi}{\left(\frac{\exp(v_L)}{(x_t \mu_{t1})^y} \Gamma(y + 1, v_L) - D_L^{1-\phi} \right)} \quad (26)$$

where $x_t = 1/(1 - \phi_t)$, $y = \frac{1 - \phi}{1 - \phi_t}$ and $v_L = x_t \mu_{t1} D_L^{1-\phi_t}$.

Equation 26 appears to fit the general trend of the fitted values well (Fig. 5) but as can be seen in Supplementary plots S31 and S32 the curves for all plots together and individual plots do not coincide so it is unclear whether this equation explains the relationship or if it is coincidental. Whether the equation is the true description or not, the relationship between μ_1 and ϕ suggests that there is a trade-off as a high μ_1 , high ϕ tree would have a superior growth:mortality ratio at smaller sizes but an inferior growth:mortality ratio at larger sizes compared to a low μ_1 , high ϕ tree. The results here hint at a trade-off that may be taking place with the results representing the dominant strategy in each forest plot depending on local conditions affecting growth and mortality.

To test this, fitting parameters μ_1 and ϕ were compared to forest plot properties such as sample size, geographical location, and mean plot height, trunk diameter, mass, wood density and basal area. The relationships were generally weak with little correlation, suggesting a poor “signal to noise” ratio or that the metrics used above had little or no correlation to the fitting parameters.

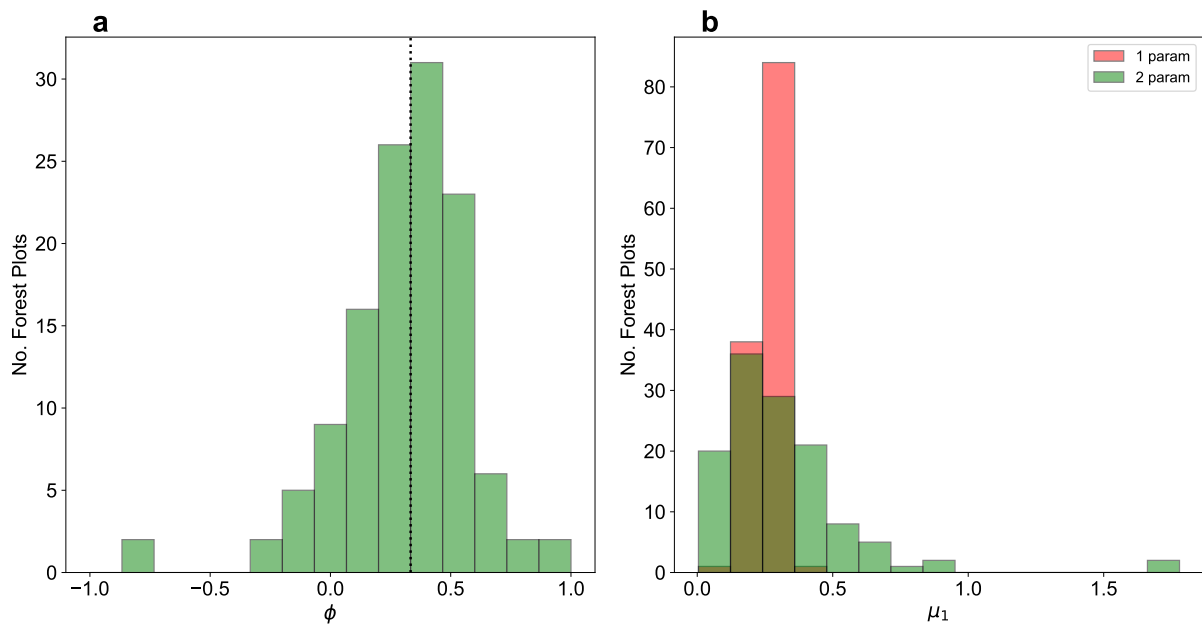


Figure 4. [a\)](#) Results for the growth scaling power ϕ when fitting the two parameter DET-LTWD via MLE for trunk diameter data from all 124 individual forest plots. The vertical black [dotted](#) line shows the value $\phi = 1/3$ predicted by MST allometry. [b\)](#) Results for fitted mortality to growth ratio μ_1 for both the 1 and 2 parameter DET-LTWD via MLE for trunk diameter data from all 124 individual forest plots.

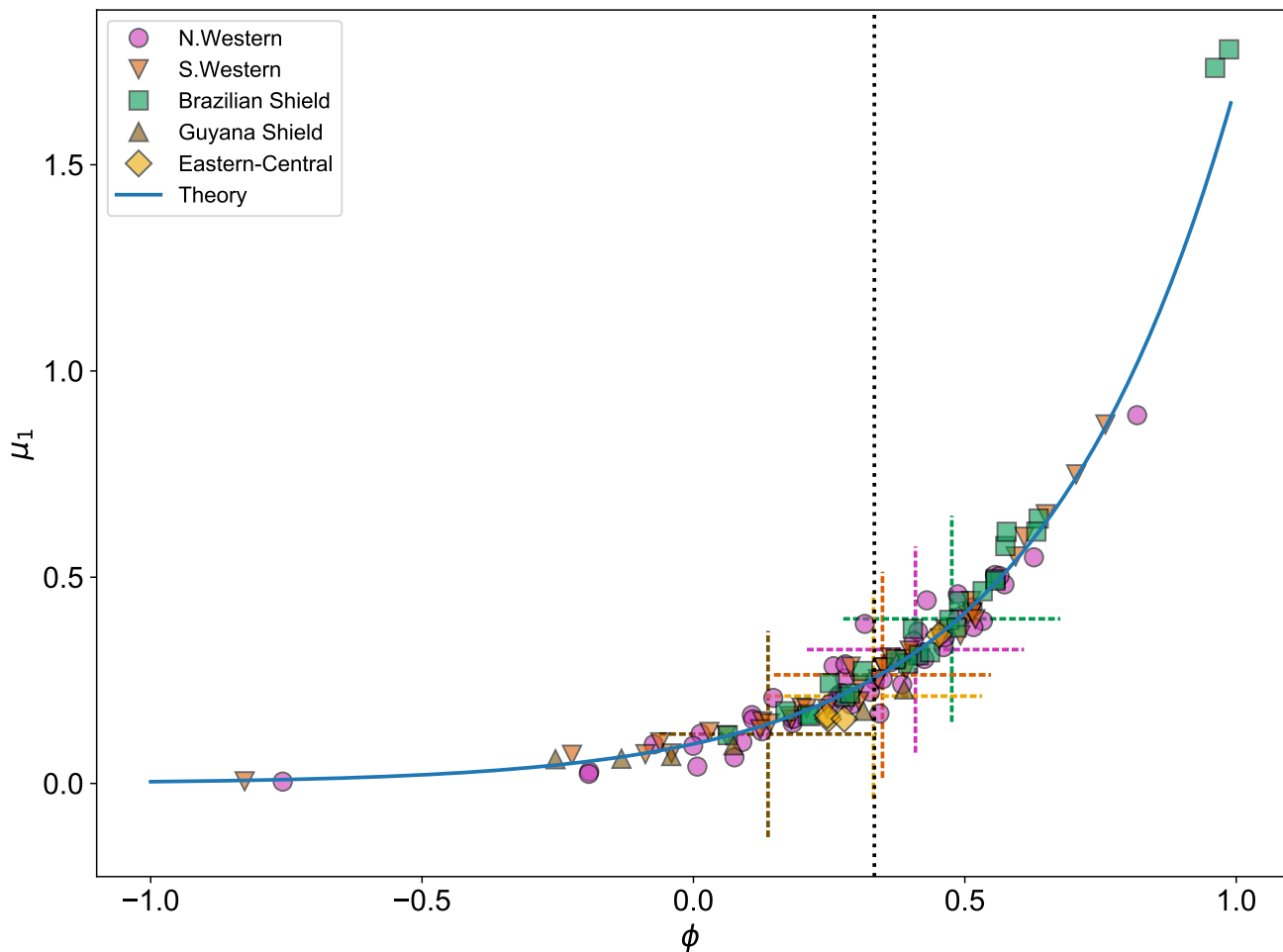


Figure 5. Results of the two parameter DET-LTWD MLE fits for trunk diameter data from all 124 individual forest plots. The fitted mortality to growth ratio μ_1 is shown as a function of the fitted growth scaling power ϕ . The results from the fits to the grouped datasets of the four allometric regions are plotted as the dashed crosses of the corresponding colour. The vertical black line shows the ϕ value predicted by MST allometry. The blue line represents the relationship derived from MLE equations for DET, showing the best fit μ_1 for a given ϕ .

4.4 Mass Results

All fitting was performed on mass data after trees smaller than m_P had been excluded. m_P was chosen based on the methodology in section 4.1.1. When fitting the DET-LTWD and MST equations to the mass size distributions, there was again a consistent pattern for all the geographical aggregations of plot data. In all cases the DET-LTWD solutions (both one and two parameter versions) fitted much better more closely than the MST solution (Fig. 6 and see supplementary material Fig. S3 and S4). Again the MST model overestimated the number of large trees.

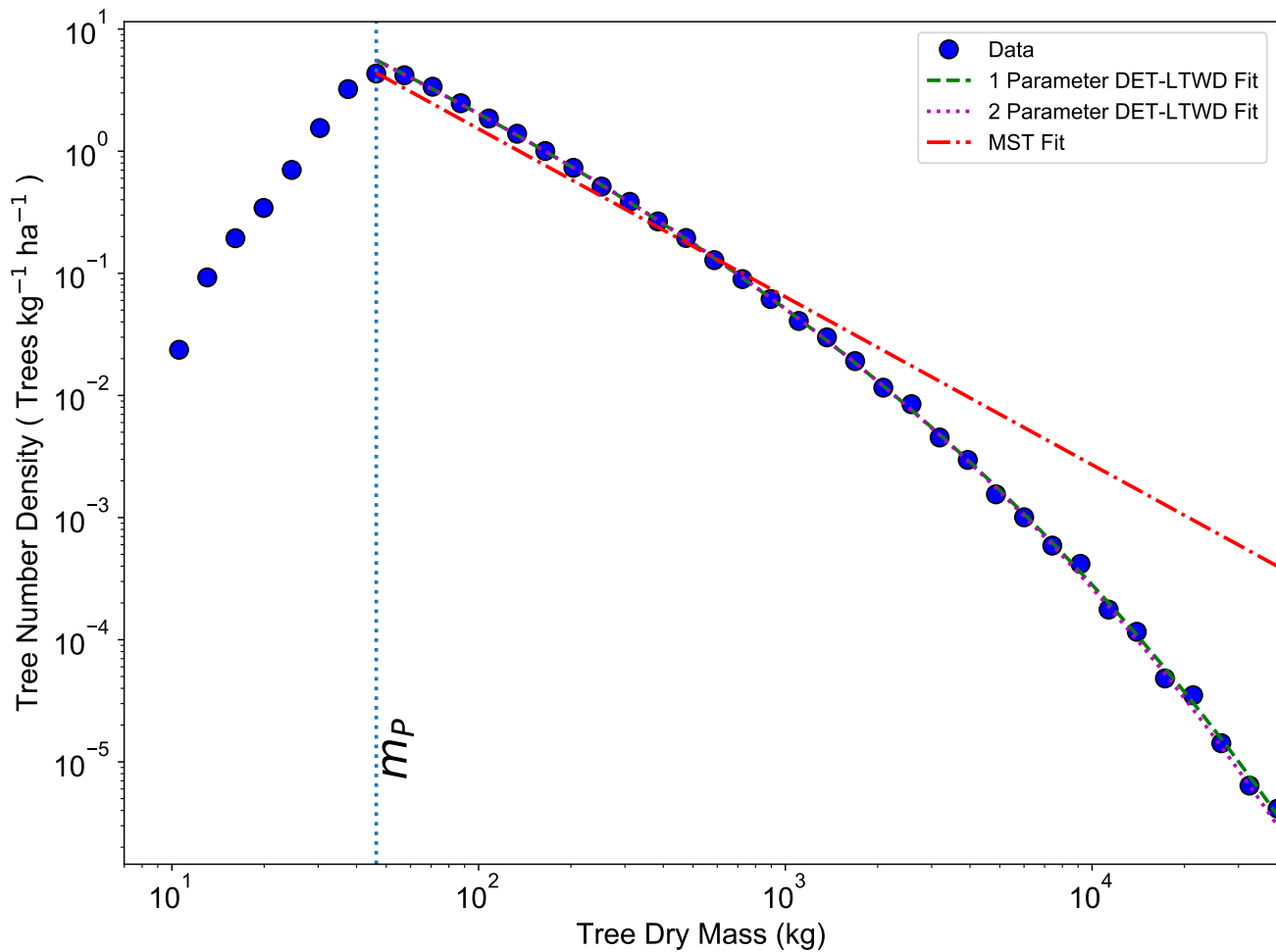


Figure 6. Fit to the mass size-distribution for all South American RAINFOR Plots as one large dataset. The blue circles show the binned data and the lines show the fitted distribution for each model. The peak in the distribution is clearly shown. The fitting is only performed on trees with mass greater than the mass of the peak.

The two parameter fits gave a fitted value of the growth scaling power ϕ_m between 0.635 and 0.794 (Table 4) which showed that the growth allometry is close to the theoretical value of 0.75 (10 of 12 regions with ϕ_m in range 0.7-0.8). The table also shows the truncation point m_P used for each dataset, and all trees with mass less than this value were excluded. The value of m_P corresponds to the peak in distribution created by the conversion from trunk diameter to mass data. The [observed allometric biomass density](#) agrees with the values found previously by Feldpausch et al. (2012), using the same [methodology biomass allometry](#). As this biomass density value is dry mass then it is a reasonable approximation (Chave et al., 2005; Martin and Thomas, 2011) to halve these values to obtain the carbon biomass density, giving a range of 10-15 kg C m⁻².

Table 4. Results of fitting the models [to of](#) the mass size-distributions for the forest plot data aggregated to regions, countries and [the whole continent all plots combined](#). Shown are the fitted parameters for each model. m_P refers to the point which all data with smaller mass was excluded to remove the allometry conversion artefact. ~~n_P is the value of the distribution at tree mass of m_P .~~ Biomass is the tree dry mass density of all trees [with dry mass](#) above m_P .

Region	No.Trees	Area ha	m_P kg	Biomass kg m ⁻²	DET 1 Param		DET 2 Param			MST
					$\mu_{m,1}$	n_P (kg ha) ⁻¹	$\mu_{m,1}$	ϕ_m	n_P (kg ha) ⁻¹	n_P (kg ha) ⁻¹
All S.America	56702	113.36	46.4	22.2	0.198	5.58	0.189	0.741	5.51	4.38
Brazil	10719	23.48	45.6	22.1	0.193	5.02	0.212	0.768	5.15	4.11
Bolivia	7892	16.00	40.6	21.7	0.199	6.12	0.225	0.773	6.33	4.92
Colombia	6167	13.21	55.5	19.0	0.216	4.95	0.188	0.724	4.79	3.45
Ecuador	4159	7.80	54.5	23.1	0.208	5.53	0.240	0.777	5.73	4.12
Peru	22414	44.50	57.3	23.5	0.204	4.93	0.194	0.741	4.87	3.59
Venezuela	2437	5.27	64.9	30.6	0.167	3.38	0.115	0.684	3.06	2.99
Amazonian Allometric Regions										
N.Western	20016	37.78	51.5	22.9	0.203	5.59	0.187	0.735	5.47	4.20
S.Western	20375	42.50	57.3	22.6	0.205	4.72	0.204	0.749	4.71	3.42
Brazilian Shield	11460	24.48	40.6	20.2	0.204	5.93	0.249	0.789	6.27	4.67
Guyana Shield	765	1.50	59.1	38.8	0.148	3.54	0.083	0.648	3.00	3.79
Eastern-Central	1773	4.00	51.8	32.7	0.157	3.61	0.147	0.737	3.54	3.53

As with the trunk diameter, fits for the two DET-LTWD solutions were, in general, quite similar in terms of the appearance on the mass distribution plots. Again the AIC and BIC fitting metrics were barely able to distinguish which DET-LTWD model best fit the data (Table 5). [Nine For nine](#) of the geographical aggregations (All S.America, Brazil, Bolivia, Colombia, Ecuador, Peru, N.Western, Guyana Shield and Eastern Central) [all could not distinguish it was not possible to distinguish between](#) the DET-LTWD fits [in with](#) either AIC or BIC. For Venezuela AIC indicated that the two parameter fit may be slightly better but BIC was not able to show any difference. The S.Western allometric region was the only one showing the one parameter fit as

being better but only for BIC. The only region to have both AIC and BIC favouring one of the fits was the Brazilian Shield region, where both AIC and BIC favoured the two parameter fit.

Table 5. Model comparison for fits to mass size-distributions. [Table-This table](#) shows the log Likelihood of each ~~models~~ ~~model's~~ fit and the corresponding AIC and BIC model comparison criterion. The best model has the lowest AIC or BIC; here the difference is shown to the best model, meaning the best model has a score of 0. Models other than the best are strongly rejected if they have a value greater than 10. Best model and those not rejected are shown in bold.

Region	log Likelihood			Δ AIC			Δ BIC		
	MST	DET	DET	MST	DET	DET	MST	DET	DET
		1 Param	2 Param		1 Param	2 Param		1 Param	2 Param
All S.America	-378,596	-371,541	-371,538	14100.0	3.68	0.0	14100.0	0.0	5.38
Brazil	-71,653	-70,609	-70,607	2090.0	2.54	0.0	2080.0	0.0	4.89
Bolivia	-51,899	-51,009	-51,006	1780.0	3.58	0.0	1770.0	0.0	3.52
Colombia	-41,118	-40,122	-40,119	1990.0	2.21	0.0	1980.0	0.0	4.69
Ecuador	-27,700	-27,241	-27,240	917.0	1.24	0.0	909.0	0.0	5.26
Peru	-151,615	-148,379	-148,378	6470.0	0.0	0.004	6460.0	0.0	8.21
Venezuela	-17,382	-17,204	-17,198	364.0	10.8	0.0	352.0	4.83	0.0
Amazonian Allometric Regions									
N.Western	-134,204	-131,530	-131,528	5350.0	3.13	0.0	5340.0	0.0	4.89
S.Western	-137,602	-134,629	-134,629	5940.0	0.0	1.97	5940.0	0.0	10.1
Brazilian Shield	-74,940	-73,604	-73,592	2690.0	20.8	0.0	2680.0	13.3	0.0
Guyana Shield	-5,545	-5,547	-5,541	3.39	8.46	0.0	0.0	9.78	6.03
Eastern-Central	-12,577	-12,487	-12,487	178.0	0.0	1.58	172.0	0.0	7.2

4.5 Mass Results for Individual Plots

Fitting the models to the individual forest plots (full results in supplementary material Tables S3 and S4 and Fig. S14 to S22) again resulted in the DET-LTWD models ~~fitting much better often fitting much more closely~~ than MST. All fitting was performed on mass data after trees smaller than m_P had been excluded. m_P was chosen, for each plot, based on the methodology in section 4.1.1. Table 6 shows the results of BIC comparison of the models for the 124 forest plots. In every case, the best model is determined by the lowest BIC value. Inferior models are only considered strongly rejected if their BIC is greater than the best model by 10 or more. This is represented by the columns in the table and shows the one parameter DET-LTWD was the best model by far the most (80 plots). However, in none of those plots was it possible to strongly reject both the other models. The most common result (74 plots) was of the one parameter DET-LTWD being the best ~~model choice~~ model (according to BIC), MST being rejected but the two parameter DET-LTWD also so closely fitting the data it cannot be rejected. The next most common result (14 plots) was the reverse with again MST rejected but the two parameter DET-LTWD narrowly better but not sufficient to strongly reject the one parameter DET-LTWD. The MST model was the best model for 15 plots and for 5 of those (ELD_01, ELD_02, RIO_01, SUC_03, TIP_03) the two DET-LTWD models were both strongly rejected. Three of these plots though had very low number of trees so it would be less expected to be able to accurately pick a model from a distribution of only ~ 100 trees.

Table 6. Shows the best and acceptable models for the 124 individual forest plots for mass. Models are labelled as (M) for MST, (1) for one parameter DET-LTWD and (2) for two parameter DET-LTWD. Columns refer to best fitting model (lowest BIC score). Rows refer to models that are so good a fit compared to the best that they cannot be rejected, as their BIC score is so close to the best model. For example '1M' means the MST and one parameter models are not rejected but two parameter model is rejected based on BIC. Then the columns in this row show how many forest plots have either (1) or (M) model as best fit and the other also fitting closely.

Comparable Models	Best Model			Total
	1	2	M	
1	0	-	-	0
2	-	11	-	11
M	-	-	5	5
12	74	14	-	88
1M	0	-	2	2
2M	-	3	2	5
12M	6	1	6	13
Total	80	29	15	124

Fig. 7 shows the effect of fitting with the two parameter DET-LTWD model. There is to be a clear relationship between ϕ_m and $\mu_{m,1}$, as all results follow a curve. ~~The black dotted vertical line shows the $\phi_m = 0.75$ value expected according to the~~

MST allometry and also assumed in the one parameter DET-LTWD model. Equation 26 can be modified to apply to mass and again fits the general trend of the fitted μ_{m1} and ϕ_m well.

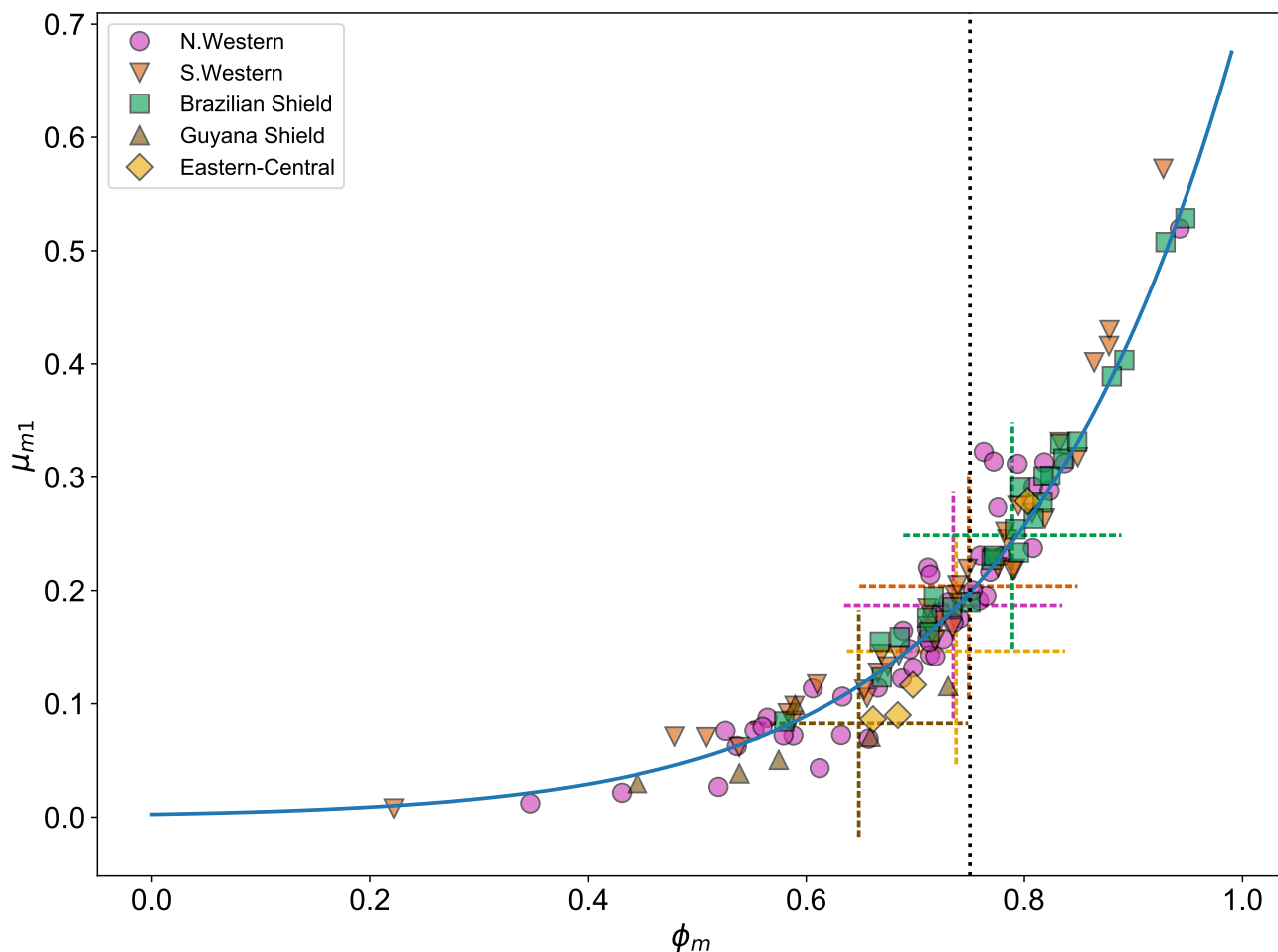


Figure 7. Results of the two parameter DET-LTWD MLE fits for mass data from all 124 individual forest plots. The fitted mortality to growth ratio μ_{m1} , for each plot, is shown as a function of the fitted growth scaling power ϕ_m . The results from the fits to the grouped datasets of the four allometric regions are plotted as the dashed crosses of corresponding colour. The vertical black line shows the ϕ_m value predicted by MST allometry. The blue line represents the relationship derived from MLE equations for DET, showing the best fit μ_{m1} for a given ϕ_m .

Plotting just the ϕ results in a histogram (Fig. 8a), reveals an approximate bell-shaped distribution with a peak around close to the theoretical MST value. The median of the ϕ_m value for the plots is 0.72 (95% confidence interval 0.71-0.75) and the mean is 0.71 (95% confidence interval 0.69-0.73). These values are close to the theoretical value of 0.75, as suggested by the histogram. The histogram of μ_{m1} (Fig. 8b) shows a bell-shaped distribution with a peak around 0.19 for both 1-parameter and

2-parameter DET-LTWD. For 1-parameter DET-LTWD the the median of μ_{m1} for the plots is 0.199 (95% confidence interval 0.196-0.205) and the mean is 0.198 (95% confidence interval 0.192-0.203). For 2-parameter DET-LTWD the median of μ_{m1} for the plots is 0.177 (95% confidence interval 0.159-0.205) and the mean is 0.194 (95% confidence interval 0.174-0.214). It is interesting that for the mass distributions all measures of central tendency cluster fairly closely to 0.19, for both 1 and

5 2-parameter fits.

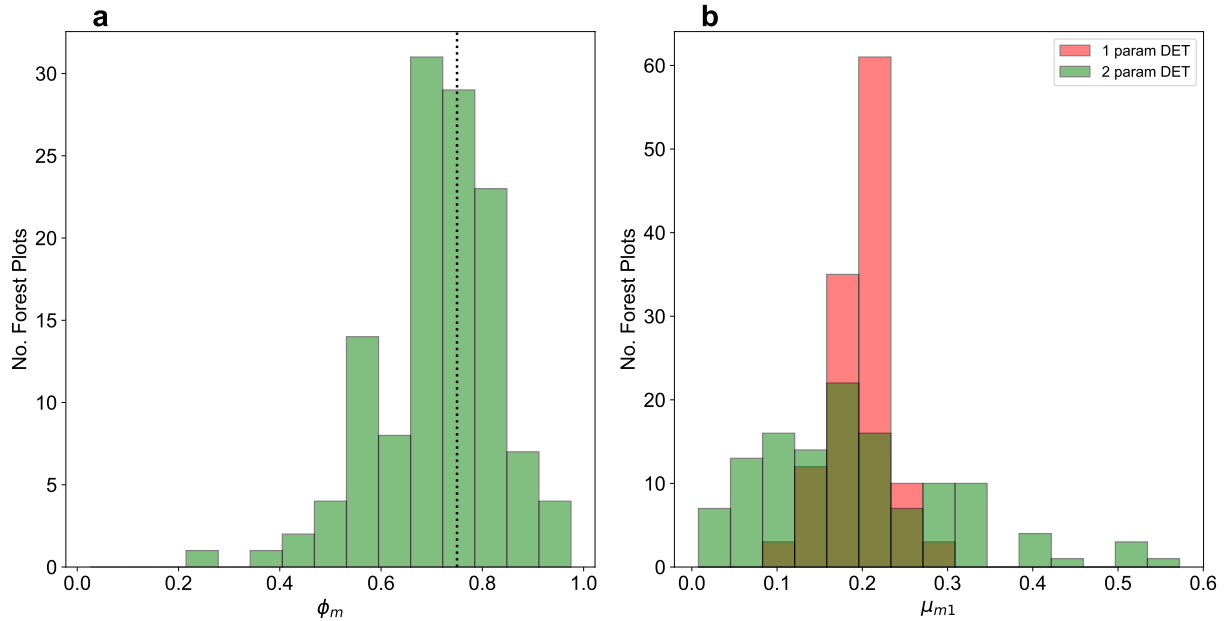


Figure 8. a) Results for the growth scaling power ϕ_m when fitting the two parameter DET-LTWD via MLE for mass data from all 124 individual forest plots. The vertical black line shows the value $\phi_m = 0.75$ predicted by MST allometry. b) Results for fitted mortality to growth ratio μ_{m1} for both the 1 and 2 parameter DET-LTWD via MLE for mass data from all 124 individual forest plots.

4.6 Biomass Results

The biomass density equations Eq. (6), Eq. (7) and Eq. (10) were tested against the ~~observed~~-allometric biomass density (summed tree mass data), as can be seen in Table 7. The biomass density equation parameters were obtained from the fits in Table 4. For the DET-LTWD solutions the biomass density was calculated for both the cases where the upper bound was infinity and the maximum tree mass in the dataset. For each of those cases, the one and two parameter DET-LTWD solutions were calculated.

The value of m_P was used for the lower bound for calculating the predicted biomass in equations Eq. (6), Eq. (7) and Eq. (10). The same values of m_P were used to truncate the data when finding the biomass density. So, comparisons between the theory and the mass obtained directly from a combination of observation and allometry were always using the same lower truncation point for each dataset but varied between datasets. The values of m_P used are given in Table 4 and the methodology used to estimate m_P is in section 4.1.1.

It is apparent that the MST biomass density equation is inferior to DET-LTWD derived biomass density equation from the DET theory. For all aggregations the biomass density was overestimated by MST, and in many cases by a considerable margin. The comparison of the different DET-LTWD biomass density equations was found to favour the two parameter fit using the finite upper bound (6 regions out of 12). Four areas had better estimates with the two parameter fit using the infinite upper bound (All S.America, Bolivia, Peru and Guyana Shield).

Interestingly, two regions (S.Western and Ecuador) had a worse fit for two parameter DET-LTWD; ~~this appears to be due to MLE fitting favouring getting the fit correct for the more numerous smaller trees rather than the larger rarer trees.~~ The S.Western region though, fits the biomass within 2% regardless of the choice of upper bound or DET model, so the very slight difference in the biomass density prediction is almost certainly not significant for this region. When the reverse cumulative biomass density, defined as biomass density of all trees above a given tree mass, is plotted for Ecuador (see supplementary material Figures S27 and S28) the error comes from the shape of the tail of the distribution, which is much flatter than theory. This could be due to it being a region with a smaller number of trees (41884159) or could be due to higher mortality for large trees in this region.

Table 7. Model Biomass Comparison. Table shows the percentage difference between each model of the biomass density predicted by the parameters obtained from fitting the mass distribution using MLE, and the **observed-allometric** mass in the dataset. This comparison is only for data where the tree mass is greater than the peak in the mass distribution m_P . Bold indicates the model that is the closest fit to the **observed-allometric** value.

	m_P	Observed-allometric	% Difference to allometric Biomass Density				
	Biomass		LTWD m_P to ∞		LTWD m_P to m_{\max}		MST
	kg	kg m ⁻²	DET 1 Param	DET 2 Param	DET 1 Param	DET 2 Param	
All S.America	46.4	22.2	1.62%	-0.09%	0.67%	-0.77%	389.9%
Brazil	45.6	22.1	-1.67%	2.21%	-3.79%	-1.34%	253.5%
Bolivia	40.6	21.7	-4.62%	0.30%	-4.50%	-0.68%	355.7%
Colombia	55.5	19.0	2.01%	-1.69%	2.88%	-0.41%	439.6%
Ecuador	54.5	23.1	1.97%	7.10%	-0.54%	2.10%	226.5%
Peru	57.3	23.5	1.17%	-0.31%	0.67%	-0.58%	407.4%
Venezuela	64.9	30.6	16.09%	2.70%	8.45%	1.62%	170.6%
Amazonian Allometric Regions							
N.Western	51.5	22.9	5.13%	2.43%	4.37%	2.12%	394.7%
S.Western	57.3	22.6	-1.12%	-1.32%	-1.51%	-1.68%	402.4%
Brazilian Shield	40.6	20.2	-7.04%	1.33%	-7.10%	-0.72%	364.4%
Guyana Shield	59.1	38.8	29.40%	4.58%	-8.53%	-8.25%	28.7%
Eastern-Central	51.8	32.7	8.76%	5.30%	1.72%	0.25%	143.7%

4.7 Biomass Results for Individual Plots

To look deeper at the relationship between model choice and predicted biomass density the analysis was repeated for the individual forest plots. In Fig. 9, the results of the biomass density predicted by the models is shown as a function of the actual ~~observed~~-allometric biomass density. It can be observed that correcting for the largest tree size in each plot is much better than assuming an infinite maximum tree size and that the one parameter model performs less well for finite maximum tree size case.

This finding is supported by looking at the relative root mean squared error (root mean squared error divided by ~~observed~~-allometric biomass density) for each model, as shown in Table 8.

Table 8. The relative root mean squared error (RMSE) of the biomass density prediction of the 124 forest plots using the parameters fitted via MLE to the mass size-distribution. The table compares the results from the different DET-LTWD models and the MST model. The range column indicates the integration limits of the biomass density calculation. The DET-LTWD model assumes no maximum size and by default integrates out to infinity. This can be corrected in terms of the largest tree mass m_{\max} in the dataset.

Model	Range	Relative RMSE
1 Parameter DET-LTWD	m_P to ∞	0.236
2 Parameter DET-LTWD	m_P to ∞	0.295
1 Parameter DET-LTWD	m_P to m_{\max}	0.098
2 Parameter DET-LTWD	m_P to m_{\max}	0.069
MST	m_P to m_{\max}	1.387

For the small individual forest plots, finite maximum tree size has a larger effect on accuracy than using the two parameter DET-LTWD over the one parameter version.

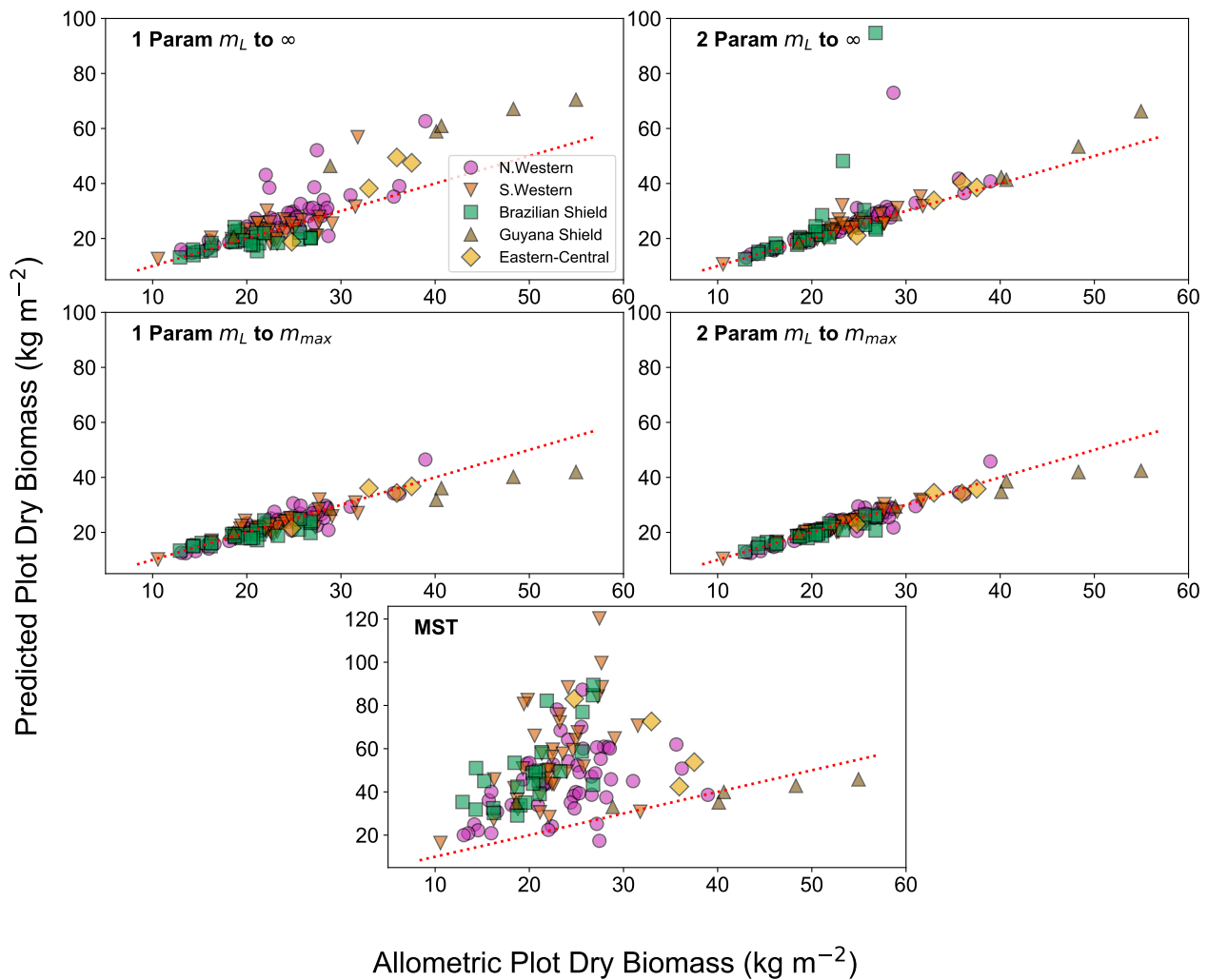


Figure 9. Comparison of the biomass density prediction based of-on the size-distribution fits to the mass data, to the observed-allometric biomass density in each of the 124 forest plots. Results are plotted for both the one and two parameter fits and for both the assumption of infinite and finite maximum tree size. The finite tree size case is limited to the largest tree mass m_{max} in each forest plot. The red dotted line shows illustrates the line of a perfect one to-to-one relationship (i.e. theory matching the data perfectly).

5 Discussion

In this paper we ~~investigated which of three demographic models (MST and two forms of DET) could be best used to model the size-distributions and total biomass density of the Amazon forest. This analysis was done for~~ show that the Left-Truncated Weibull (LTWD), which is consistent with the Demographic Equilibrium Theory (DET) when the mortality is size independent and the growth is a power-law of tree size, fits the observed tree-size distributions for 124 forest plots across Amazonia. Our fitting was undertaken with either two free parameters or with one free parameter and the growth scaling power ϕ constrained to that specified in Metabolic Scaling Theory (1/3 for trunk diameter and 3/4 for mass, see West et al. 2009; Niklas and Spatz 2004). We also compared the performance of DET-LTWD to that of the Metabolic Scaling Theory for forest demography (MSTF, West et al. 2009). Our analyses were carried out for both trunk diameter measurements and for those same measurements converted allometrically (Feldpausch et al., 2012) to mass. This conversion trunk diameter converted allometrically to mass (Feldpausch et al., 2012).

We found that this conversion of trunk diameter to mass introduces a peak in the mass distribution that is purely an artefact of the conversion. The peak is due to the variation in mass of trees of a given trunk diameter, due to height and wood density variation leading to some small mass trees being in effect “missing” from the mass distribution. If the diameter to mass relationship was purely one-to-one, then the artefact peak would not occur. This has implications for anyone using mass size-distributions converted from trunk diameter data. Our solution ~~is to only to fit to~~ was to fit only to trees with mass greater than the mass distribution peak.

~~Each of the three models of size-distribution could predict total biomass density by the integration of the analytical form of mass-distribution. One interesting implication of the resulting equations is that the mortality and growth only ever appear in the form of the ratio μ_T and never independently. This suggests that the ratio of mortality to growth determines the equilibrium state of a forest, and that the actual magnitude of the individual mortality and growth terms only determines the transient effects away from a steady state.~~

~~Of the three models, MST is rarely a good fit at plot, regional or continental level for either trunk diameter or mass distributions and significantly overestimates total biomass density. The MST was only the undisputed best model in 4% of forest plots and only for 20% was the model a good enough fit not to be rejected. For the aggregated datasets (continent, regions and countries) only the Guyana region was the MST model not rejected, and this region only has a small number of trees, which leads to higher uncertainty in the fit.~~ The model fitting shows that Amazon size-distributions are generally better fit by the DET-LTWD based models than MSTF. The 2 and 1-parameter DET-LTWD fits were often not significantly different enough from each other for comparison by AIC or BIC (which balance the quality of the fit against the number of unknown parameters) to choose which is the best description of the size-distributions. The few plots and regions (including all plots combined) where one model was found to have a significantly better AIC or BIC score all favoured the 2-parameter model.

~~The DET model assumed power-law growth with size and size-independent mortality, which gives a left truncated Weibull distribution (LTWD). best-fit growth-scaling exponent ϕ varied between plots and regions, but the mean value of ϕ across all 124 plots fell close to the values predicted by MST. For the 1-parameter DET-LTWD was tested both when its growth~~

allometry was constrained to the MST growth allometry (West et al., 2009; Niklas and Spatz, 2004) and when the growth allometry parameter was allowed to vary as a fitting parameter (one and two parameter fitting respectively) best-fit values of μ_1 for trunk diameter cluster tightly around 0.25 (and around $\mu_{m1} = 0.19$ for mass). This comparison allows the MST allometry to be tested independently of the space-filling assumptions of MST. If the MST allometry is true, then the two DET-LTWD models should be in agreement as the unconstrained version of the model can still select the MST allometry if it is the best fit. The BIC criterion for model selection showed that the MST-constrained DET-LTWD was in contention for 89% of forest plots, for five of the six countries and for three of the five regions, and for mass distribution of the whole continent. The unconstrained DET-LTWD was only clearly the best model for the trunk diameter distribution of the whole continent, Bolivia and the Brazilian Shield region.

For the unconstrained DET-LTWD model, the growth scaling power is close to the mean value of $\mu_1 = 0.22$ that we found for North American forests (Moore et al., 2018), hinting at a preferred value of the ratio of mortality to growth across different regions and forest types.

The clustering of ϕ did vary between plots and regions but the median/mean values and the continental result all fell very close to the values predicted by the MST allometry. For the aggregated datasets, the fitted ϕ values for the mass distributions were closer to MST than for the trunk diameter distributions. This finding suggests that the MST allometry value predicted by MST allometry (Niklas and Spatz, 2004; West et al., 2009) suggest two possibilities. Either that the clustering represents an underlying reality for the scaling of trees "basin of attraction" that is modified by local conditions. This idea was first raised by Price et al. (2007) and later disputed by Coomes and Allen (2009). It was further suggested (Coomes et al., 2011) (Price et al., 2007) or that plots do not meet the model assumptions of growth, mortality and equilibrium and this in turn somehow leads to this clustering. We cannot say for certain why the plots cluster close to the MST values but it does lead to intriguing future avenues of study.

It was suggested (Coomes and Allen, 2009; Coomes et al., 2011) that light competition should modify the growth function MST scaling of growth with size. This would mean that for trunk diameter the growth scaling power would vary with size and be greater than the predicted MST value of 1/3. For our regional fits the fitted power, while close to the theoretical was slightly larger than the MST value of 1/3, it was larger in most cases but for the individual forest plots, the value was very close to MST with no clear bias. So our results cannot be taken as conclusive evidence of light competition modifying the growth scaling but neither are they completely inconsistent with it.

We find the fitted 2-parameter DET-LTWD ϕ values for both mass and trunk diameter also have a well defined relationship to the fitted mortality : growth ratio μ_1 . This relationship does not appear to be a fitting artefact, as if artificial data is generated with known μ_1 and ϕ values off the observed curve the fitting process ϕ correctly fits it to the generated values, not the curve seen in this study. This relationship suggests an interesting but as yet unknown property of the Amazon forests but may represent life-history trade-offs (Uriarte et al., 2012). Trees have different strategies such as live-fast die-young pioneer species versus grow-slow live-long canopy species. This is one possible explanation of the relationship between μ_1 and ϕ , as when both are high the early growth at small size will be slower but keep increasing, while when ϕ and μ_1 are both low the early growth will be higher but more quickly level off. Interestingly no plots had low phi, with high μ_1 , which would correspond to

uncompetitive low growth at all sizes. As these results are at the plot level rather than per tree basis, it would suggest that each site has a dominance of one life-history strategy. As there is no correlation of μ_1 or ϕ with plot metrics such as height or wood density, this hypothesis remains unconfirmed.

When considering how well the models predicted total biomass density from the fitted size-distribution, the biggest source of error at the plot scale is the model assumption of infinite maximum tree size. However, this can be easily corrected for and allows the MST-constrained DET-LTWD to estimate biomass density with relative root-mean-square error of 10% over the 124 forest plots and unconstrained DET within 6%. Conversely, the MST model consistently overestimated the biomass density, often by a considerable margin. The regional scale, where there were more trees, showed much better prediction of the biomass density and the unconstrained DET-LTWD with finite upper bound had the smallest error in biomass density.

In terms of what this means, again MSTF was rarely a good fit at plot, regional or all plots level for either trunk diameter or mass distributions, and significantly overestimated total biomass density, so we reject the MST-MSTF model as a good model of forest size-distributions. This rejection is consistent with the recent study by Zhou and Lin (2018) that showed there was a fundamental flaw in the derivation of MST. They had noticed that the MST-the MSTF model failed to account for the effect of size-dependent growth rate on how fast a tree transitions through a given size class. This observation explains that the assumptions of MST-MSTF of the size distribution scaling D^{-2} is inconsistent with the assumption of individual tree resource use scaling as D^2 . Here, we have confirmed the D^{-2} (and $m^{-11/8}$) size-distribution model should be rejected for South American tropical forests. Furthermore, for most plots also we can reject a general power law distribution also, as the distributions observed are rarely linear when plotted in log-log space.

For large scale modelling, such as DGVMs, the extra parameter of the unconstrained DET-LTWD performs better, particularly for biomass density. Despite this, the constrained version still does quite well, and for some regions can actually be better. So, the DET-LTWD model constrained by MST allometry has the benefits of simplicity but at a small cost in accuracy. At the smaller scale of individual forest plots, the variation in fitted growth scaling allometry is greater. At these smaller scales, the extra allometry parameter in fitting is much more important but even then still allows biomass density modelling with 10% mean error. So, including the extra fitting parameter is a trade-off between simplicity and accuracy and model choice will depend on the needs of the particular application. There was a strong correlation between sample size and how likely MSTF was to be considered either the best or a acceptable model, with small sample sizes favouring MSTF. This suggests that sample sizes may lead to difficulty identifying the best model or even wrongly choose the best model, most likely as rarer large trees are more likely to be absent from a small sample. Meaning, where practical, larger forest plots of at least a 1000 stems are desirable when analysing size-distributions.

5.1 Closed Form of DET Solutions for Applications in DGVMs

As Demographic Equilibrium Theory (DET) provides such a good description of the mass-distributions observed in Amazonia, this suggests that dynamic forest demography (as given by Eq. 1) could form the basis of a next-generation Dynamic Global Vegetation Model (DGVM) to be used with an Earth System Model (ESM). For such DGVM applications, the DET model solutions need to be extended by deriving a closed form that includes seedling recruitment. We do this by partitioning a fixed

fraction α of All three models of size distribution were used to predict total biomass density by the integration of the analytical form of their respective mass distributions. One interesting implication of the resulting equations for DET is that mortality and growth only ever appear in the form of the net assimilate P (net primary productivity remaining after accounting for litterfall) of each tree to seedling production, with the remainder of P going into the growth of the tree itself ratio μ_1 and never independently. The ratio of mortality to growth therefore determines the equilibrium state of a forest, while the absolute magnitudes of the individual mortality and growth terms determine the transient effects away from a steady state.

This allows a solution to be derived (see Appendix) for the total equilibrium fractional coverage ν (fraction of ground area covered by tree crowns) that is only a function of three parameters, under the assumption that there is negligible overlap of When considering how well the models predicted total biomass density from the fitted size-distribution, the crowns in the canopy (Fig. ??).

$$\nu = 1 - \left(\frac{1 - \alpha}{\alpha} \right) \frac{\mu_s (x\mu_s)^{x-1}}{\exp(x\mu_s) \Gamma(x, x\mu_s)}$$

where $\mu_s = \mu_1 m_s^{(1-\phi)}$ is the mortality to growth ratio at the seedling size m_s .

Further solutions can then be derived (see Appendix) for total forest properties such as biomass density M (Fig. ??) and in a way that eliminates dependent terms such as n_s (number density per size class at seedling size). To do this we assume the crown area of biggest source of error at the plot scale is the model assumption of infinite maximum tree size. However, this can be corrected for and allows the 1-parameter DET-LTWD to estimate biomass density with relative root mean square error of 10% over the 124 forest plots and 2-parameter DET-LTWD within 6%. Conversely, the MST model consistently overestimated the biomass density, often by a considerable margin. The regional scale, which has larger sample size, showed much better prediction of the biomass density and the trees in the forest scales as a power law with scaling power ϕ_a and seedlings have crown area a_s , mass m_s and growth g_s . 2-parameter DET-LTWD with finite upper bound had the smallest error in biomass density. This suggests the DET-LTWD model is a useful model of biomass for large-scale applications such as being used to initialise a DGVM based on the continuity equation equation (Argles et al., 2019) or as a climate relevant measure of goodness of fit.

One of our priorities for further work is to investigate whether the commonality found in the values of μ_1 and the relationship between μ_1 and ϕ is indicative of some form of optimality operating at the forest scale.

$$M = \frac{m_s (x\mu_s)^{x(\phi_a-1)} \Gamma(x+1, x\mu_s)}{a_s \Gamma(\phi_a x + 1, x\mu_s)} \nu$$

This means we have simple equilibrium solutions that are a function of known parameters of each plant functional type (PFT) and of parameters that can be provided by the land surface scheme of an ESM. Equilibrium solutions can also be obtained for a discrete set of size classes, allowing the demographic profile to be initialised in an equilibrium state consistent with observed mass distributions and forest area coverage. Once such a model is initialised it can be run dynamically to simulate transient

states based on inputs of additional mortality due to disturbance processes and time-varying net assimilate from a driving land-surface scheme. (Argles et al., 2019). We are optimistic that this will allow more realistic simulation of the response of forests to changes in climate and atmospheric carbon dioxide, including possible forest die-back due to increasing temperatures, drought and forest fires.

- 5 Shows the dependence of the total forest fractional coverage ν and total biomass density M on the mortality to growth ratio of seedlings μ_s for the closed form solutions where the rate of seedling production is a fixed fraction α of tree productivity. These plots assume $\phi_m = 0.75$, and seedlings with mass m_s of 10g and that a 1 kg tree would have a crown area of 0.5 m^2 .

6 Conclusions

This study demonstrates that demographic equilibrium theory (DET) is able to fit measured tree size-distributions in Amazonian forests. ~~DET was found to outperform Metabolic Sealing theory (MST, West et al. (2009)), even when using just a single fitting parameter (μ_1 —the ratio of mortality to growth), and assuming the MST allometric relation between tree growth-rate~~ The fitted growth scaling parameter ϕ was clustered for both trunk diameter (0.31 ± 0.02) and tree-mass ($g \propto m^{3/4}$, Niklas and Spatz (2004)). ~~When the exponent relating growth-rate to tree-mass~~ mass diameter (0.71 ± 0.01) distributions close to the values predicted by Metabolic Sealing Theory (MST). The small bias seen could be indicative of deviations from MST allometry due to light competition. The fitted mortality: growth ratio parameter μ_1 was clearly related to the fitted ϕ was also allowed to vary by site, DET produced an even better fit to the observed tree-mass distribution, and also an intriguing (but as yet unexplained) parameter suggesting a possible life-history trade-off in the forest plots. If the DET ϕ is constrained to the MST value then the fit is often as good as the 2-parameter fit and with one less fitting parameter is preferred by the Bayesian Information Criterion and μ_1 clusters with a value (0.25 for trunk diameter) close to that of 0.22 previously reported for US forests. We therefore find evidence that the 1-parameter DET is useful in modelling forests on the global scale, particularly for applications where parameter sparsity is important (Argles et al., 2019). Further support for such applications comes from the model's ability to replicate forest biomass density over large scales, when compared to the data. The relationship between μ_1 and ϕ and a common value of μ_1 across sites. Furthermore, equations derived from DET, predicting the biomass density in terms of ϕ and μ_1 are a very climate relevant measure of goodness of fit for these models. The success of DET in explaining tree-size distributions across both North America (Moore et al., 2018) and South America (this study) indicates that relatively simple and robust ecosystem demography is a good basis for a next-generation dynamic global vegetation model (Argles et al. (2019), in preparation). between the US and Amazon may indicate some optimality principle is in play.

Code availability. Code is available on reasonable request to the corresponding author.

Appendix A

~~The closed form model (see Section ??) makes the following assumptions:—~~

- ~~– All trees produce seeds and there is no minimum reproductive size threshold.~~
- 25 ~~– All seeds have mass m_s .~~
- ~~– Seeds reaching the ground either die or are recruited as a sapling, no seed pool is modelled.~~
- ~~– All trees have a crown area that follows a power law with tree mass, and that the crown area is a big leaf and totally opaque with no light allowed through.~~
- ~~– Trees have the minimum overlap in crowns possible (perfect plasticity assumption).~~

- The proportion of seeds recruited ω is determined by shading. This proportion $\omega = 1 - v$, where v is the fractional coverage of crowns of all trees with mass greater than the seed mass m_s .
- A proportion of productivity α is assumed to go into seed production.

The total growth per hectare G , for all trees larger than m_s , is

$$5 \quad \underline{G = (1 - \alpha)P}$$

where P is the Net Productivity after losses from respiration and litter are accounted for. The amount of mass going into seeds P_S is then defined as

$$\underline{P_S = \alpha P}$$

Therefore

$$10 \quad \underline{P_S = \frac{\alpha}{1 - \alpha} G}$$

The DET solution describes the size distribution, for tree mass growth rate $g(m) = g_s(m/m_s)^\phi$

$$\underline{n(m) = n_s \left(\frac{m}{m_s}\right)^{-\phi} \exp\left\{\frac{\mu_s}{1 - \phi} \left[1 - \left(\frac{m}{m_s}\right)^{1-\phi}\right]\right\}, \phi \neq 1}$$

The mortality growth ratio μ_s is a function of α as the individual tree growth g_s is a fraction $(1 - \alpha)$ of the tree productivity $p(m) = p_s(m/m_s)^\phi$.

$$15 \quad \underline{\mu_s = \frac{\gamma m_s}{g_s} = \frac{\gamma m_s^{(1-\phi)}}{g_1} = \mu_1 m_s^{(1-\phi)} = \frac{\gamma m_s}{p_s(1 - \alpha)}}$$

The equation for the total mass growth from trees with mass from m_s to ∞ is

$$\underline{G = \int_{m_s}^{\infty} n(m)g(m)dm = g_s N \frac{\exp(x\mu_s)}{(x\mu_s)^{x-1}} \Gamma(x, x\mu_s)}$$

While the tree / stem density N is

$$N = \int_{m_s}^{\infty} n(m) dm = \frac{n_s m_s}{\mu_s} = \frac{n_s g_s}{\gamma}$$

The boundary condition defines the seed recruitment rate $n_s g_s$ (in units saplings $\text{yr}^{-1} \text{ha}^{-1}$)

$$n_s g_s = \frac{P_S}{m_s} (1 - v_s) = \frac{\alpha}{1 - \alpha} \frac{G}{m_s} (1 - v)$$

- 5 By substituting equation ?? into equation ?? and then rearranging in terms of v_s , and eliminating N using equation ??, a solution for v can be obtained that only depends on known variables $x(\phi)$, α and μ_s .

$$v = 1 - \left(\frac{1 - \alpha}{\alpha} \right) \frac{\mu_s (x \mu_s)^{x-1}}{\exp(x \mu_s) \Gamma(x, x \mu_s)}$$

v is also defined by integrating the crown area weighted by $n(m)$ between m_s and ∞

$$v = \int_{m_s}^{\infty} a(m) n(m) dm = \frac{N a_s \exp(x \mu_s)}{(x \mu_s)^{\theta_a x}} \Gamma(\theta_a x + 1, x \mu_s)$$

where the individual tree crown area $a(m) = a_s m^{\phi_a}$.

- 10 Rearranging equation ?? gives another equation for N

$$N = \frac{1}{a_s} \frac{(x \mu_s)^{\theta_a x}}{\exp(x \mu_s)} \frac{1}{\Gamma(\phi_a x + 1, x \mu_s)} \nu$$

This result can then be combined with the equations for G (equation ??) and M (equation ??) to obtain results independent of N or n_s .

Firstly, the equation for total mass M is

$$15 \quad M = m_s N \frac{\exp(x \mu_s)}{(x \mu_s)^x} \Gamma(x + 1, x \mu_s)$$

The N independent versions are then

$$M = \frac{m_s (x \mu_s)^{x(\theta_a - 1)}}{a_s} \frac{\Gamma(x + 1, x \mu_s)}{\Gamma(\phi_a x + 1, x \mu_s)} \nu$$

$$G = \frac{g_s (x\mu_s)^{x(\theta_a-1)+1}}{a_s} \frac{\Gamma(x, x\mu_s)}{\Gamma(\phi_a x + 1, x\mu_s)} \nu$$

For the MST allometry then $x = 4$ and $\theta_a = 1/2$

$$v = 1 - \left(\frac{1-\alpha}{\alpha} \right) \frac{\mu_s (4\mu_s)^3}{\exp(4\mu_s) \Gamma(4, 4\mu_s)}$$

$$N = \frac{1}{a_s} \frac{(4\mu_s)^2}{\exp(4\mu_s)} \frac{1}{\Gamma(3, 4\mu_s)} \nu$$

$$5 \quad M = \frac{m_s}{a_s} \frac{1}{(4\mu_s)^2} \frac{\Gamma(5, 4\mu_s)}{\Gamma(3, 4\mu_s)} \nu$$

$$G = \frac{g_s}{a_s} \frac{1}{(4\mu_s)} \frac{\Gamma(4, 4\mu_s)}{\Gamma(3, 4\mu_s)} \nu$$

For these particular MST allometry values the Gamma functions can be represented by a finite series as when a is an integer in $\Gamma(a, z)$ then

$$\Gamma(a, z) = (a-1)! e^{-z} \sum_{k=0}^{a-1} \frac{z^k}{k!}$$

$$10 \quad v = 1 - \left(\frac{1-\alpha}{\alpha} \right) \frac{\mu_s}{\left(1 + \frac{3}{4\mu_s} + \frac{3}{8\mu_s^2} + \frac{3}{32\mu_s^3} \right)}$$

$$N = \frac{\nu}{a_s} \left(\frac{1}{1 + \frac{1}{2\mu_s} + \frac{1}{8\mu_s^2}} \right)$$

$$G = \frac{\nu g_s}{a_s} \left(\frac{1 + \frac{3}{4\mu_s} + \frac{3}{8\mu_s^2} + \frac{3}{32\mu_s^3}}{1 + \frac{1}{2\mu_s} + \frac{1}{8\mu_s^2}} \right)$$

$$M = \frac{\nu m_s}{a_s} \left(\frac{1 + \frac{1}{\mu_s} + \frac{3}{4\mu_s^2} + \frac{3}{8\mu_s^3} + \frac{3}{32\mu_s^4}}{1 + \frac{1}{2\mu_s} + \frac{1}{8\mu_s^2}} \right)$$

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Also see the supplementary material Table S1 for a more detailed list of which plot each PI worked on.

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