1 Title

2 Pan-tropical prediction of forest structure from the largest trees

3 Short Title header

4 Pan-tropical forest structure from the largest trees

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196 Abstract

Aim. Large tropical trees form the interface between ground and airborne observations, offering a unique opportunity to capture forest properties remotely and to investigate their variations on broad scales. However, despite rapid development of metrics to characterize the forest canopy from remotely sensed data, a gap remains between aerial and field inventories. To close this gap, we propose a new pan-tropical model to predict plot-level forest structure properties and biomass from just the largest trees.

203 **Location**. Pan-tropical

204 **Time period**. Early 21st century

205 Major taxa studied. Woody plants

Method. Using a dataset of 867 plots distributed among 118 sites across the tropics, we tested the prediction of the quadratic mean diameter, basal area, Lorey's height, community wood density and aboveground biomass from the ith largest trees.

209 Result. Measuring the largest trees in tropical forests enables unbiased predictions of plot and 210 site-level forest structure. The 20 largest trees per hectare predicted quadratic mean diameter, 211 basal area, Lorey's height and community wood density and aboveground biomass with 12%, 16%, 4%, 4% and 17.7% of relative error. Most of the remaining error in biomass prediction is 212 213 driven by differences in the proportion of total biomass held in medium size trees (50-70 cm), which shows some continental dependency with American tropical forests presenting the 214 215 highest proportion of total biomass in these intermediate diameter classes relative to other continents. 216

Conclusion. Our approach provides new information on tropical forest structure and can be employed to accurately generate field estimates of tropical forest carbon stocks to support the calibration and validation of current and forthcoming space missions. It will reduce the cost of field inventories and contribute to scientific understanding of tropical forest ecosystems and response to climate change.

222 Introduction

The fundamental ecological function of large trees is well established for tropical forests. They 223 224 offer shelter to multiple organisms (Remm & Lõhmus, 2011; Lindenmayer et al., 2012), regulate forest dynamics, regeneration (Harms et al., 2000; Rutishauser et al., 2010) and total 225 biomass (Stegen et al., 2011), and are important contributors to the global carbon cycle 226 (Meakem et al., 2017). Being major components of the canopy, the largest trees may also 227 suffer more than sub-canopy and understory trees from climate change, as they are directly 228 229 exposed to variations in solar radiation, wind strength, temperature seasonality and relative air humidity (Laurance et al., 2000; Nepstad et al., 2007; Lindenmayer et al., 2012; Thomas et al., 230 2013; Bennett et al., 2015; Meakem et al., 2017). Because they are visible from the sky, large 231 trees are ideal for monitoring forest responses to climate change via remote sensing (Bennett 232 233 et al., 2015; Asner et al., 2017).

Large trees encompass a disproportionate fraction of total aboveground biomass (AGB) in 234 tropical forests (Chave et al., 2001; Lutz et al., 2018), with some variations in their relative 235 236 contribution to the total AGB among the tropical regions (Feldpausch et al., 2012). In Central 237 Africa, the largest 5% of trees in a forest sample plot, i.e. the 5% of trees with the largest 238 diameter at 130 cm, store 50% of forest plot aboveground biomass on average (Bastin et al., 2015). Consequently, the density of large trees largely explains variation in forest AGB at local 239 240 (Clark & Clark, 1996), regional (Malhi et al., 2006; Saatchi et al., 2007), and continental scales 241 (Stegen et al., 2011; Slik et al., 2013). Detailing the contribution of each single tree to the 242 diameter structure, we showed previously that plot-level AGB can be predicted from a few large trees (Bastin et al., 2015), with the measurement of the 20 largest trees per hectare being 243 sufficient to estimate plot-level biomass with less than 15% error in reference to ground 244 245 estimates. These findings suggested that a substantial gain of cost-effectiveness may be achieved by focusing forest inventories on the largest trees rather than all size classes. 246 Similarly, it suggested that remote sensing (RS) approaches could focus on the measurement 247 of the largest trees, instead of properties of the entire forest stand. 248

Several efforts are underway to close the gap between remote sensing of forest biomass and 249 field surveys (Coomes et al., 2017; Jucker et al., 2017). However, existing RS approaches 250 251 typically require ground measurement of all trees above or equal to 10 cm of diameter (D) for 252 calibration (Asner et al., 2012; Asner & Mascaro, 2014). Collecting such data in the field is costly and time-consuming, which therefore limits the spatial representativeness of available 253 plot networks. Besides, extrapolation methods of ground-based biomass estimations on RS 254 data still faces important limits. For instance, using mean canopy height extracted from active 255 256 sensors (Mascaro et al., 2011; Ho Tong Minh et al., 2016), or canopy grain derived from optical images (Proisy et al., 2007; Ploton et al., 2012, 2017; Bastin et al., 2014), the biomass is 257 predicted with an error of only 10-20% compared to ground-based estimates. However, this 258 good level of accuracy is limited to the extent of the RS scene used, which considerably 259 decrease in the upscaling step necessary for national of global maps (Xu et al., 2017). A 260 promising development to alleviate this spatial restriction lies in the 'universal approach', 261 proposed by Asner et al. (2012) and further adapted in Asner and Mascaro (2014), in which 262 263 plot-level biomass is predicted by a linear combination of ground-based and remotely-sensed 264 metrics. The 'universal approach' relies upon canopy height metrics derived from radar or LiDAR (top of canopy height, TCH), and basal area (BA, i.e. the cumulated cross-sectional 265 stems area) and community wood density (i.e. weighted by basal area, WD_{BA}) derived from 266 267 field inventories. Plot AGB is then predicted as follows (Asner et al., 2012):

268 AGB = $aTCH^{b1}BA^{b2}WD_{BA}^{b3}(1)$

While generally performing better than approaches based solely on remote sensing of tree height (Coomes *et al.*, 2017), this model relies on exhaustive ground measurements (i.e. wood density and basal area of all trees above 10 cm of diameter at 130 cm, neither of which is measured using any existing remotely sensed data).

Recent advances in remote sensing allow the identification of single trees in the canopy (Ferraz *et al.*, 2016), estimation of adult mortality rates for canopy tree species (Kellner & Hubbell,
2017), description of the forest diameter structure (Stark *et al.*, 2015), depiction of crown and
gap shapes (Coomes *et al.*, 2017), and even identification of some functional traits of canopy

species (Asner *et al.*, 2017). As routine retrieval of some canopy tree metrics is within reach, we test here the capacity of the largest trees, i.e. trees that can be potentially derived using remote sensing, to predict plot-level biomass. To this end, we adapted equation (1) as follows: $AGB = a(Dg_{LT}, H_{LT}, WD_{LT})^{b1}$ (2)

where for the ith largest trees, Dg_{LT} is the quadratic mean diameter, H_{LT} the mean height, and WD_{LT} the mean wood density among the ith largest trees.

Using a large database of forest inventories gathered across the tropics (Figure 1), including 283 284 secondary and old growth forest plots, we test the ability of the largest trees to predict various metrics estimated at 1-ha plot level, namely the mean quadratic diameter, the basal area, the 285 Lorey's height (i.e. plot-average height weighted by basal area), the community wood density 286 (i.e. plot-average wood density weighted by basal area) and mean aboveground live biomass 287 (supplementary figure 1). By testing different numbers of largest trees as predictors, we aim to 288 propose a threshold of the minimal number of largest trees required to predict forest plot 289 290 metrics at a pan-tropical level with no bias and low uncertainty (i.e. error inferior to 20%). While 291 previous work focused on estimating biomass in Central African forests (Bastin et al., 2015), 292 the present study aims at generalizing the potential of large trees to predict these different plot 293 metrics at continental and pan-tropical scales. Taking advantage of a unique dataset gathered 294 across the tropics (867 1-ha plots), we also investigate major differences in forest structure 295 across the three main tropical regions: the Americas, Africa and Asia. We further discuss how 296 this approach can be used to guide innovative RS techniques and increase the frequency and 297 representativeness of ground data to support global calibration and validation of current and planned space missions. These include the NASA Global Ecosystem Dynamics Investigation 298 (GEDI), NASA-ISRO Synthetic Aperture Radar (NISAR), and ESA P-band radar (BIOMASS) 299 300 (Le Toan *et al.*, 2011; Dubayah *et al.*, 2014). This study is a step forward in bringing together remote sensing and field sampling techniques for guantification of terrestrial C stocks in tropical 301 302 forests.

303 Material & Methods

304 Database

305 For this study, we compiled standard forest inventories conducted in 867 1-ha plots from 118 306 sites across the three tropical regions (Figure 1), including mature and secondary forests. Each site comprises all the plots in a given geographical location, i.e. within a 10 km radius and 307 collected by a Principal Investigator and its team. These consisted of 389 plots in America (69 308 sites), 302 plots in Africa (35 sites) and 176 plots in Asia (14 sites). Data were provided by 309 310 Principal Investigators (see supplementary Table 1), and through datasets available on the following networks: TEAM (http://www.teamnetwork.org/), CTFS (http://www.forestgeo.si.edu/; 311 Condit et al., 2012) and ForestPlots (https://www.forestplots.net/) for AfriTRON (the African 312 Tropical Rainforest Observation Network; <u>www.afritron.org</u>) and RAINFOR (the Amazon forest 313 314 inventory network; http://www.rainfor.org/) networks.

We selected plots located between 23°N and 23°S, including tropical islands, with an area of 315 1-ha to ensure stable intra-sample variance in basal area (Clark & Clark, 2000). Plots in which 316 317 at least 90% of the stems were identified to species, and in which all stems with the diameter 318 at 130 cm greater than or equal to 10 cm had been measured were included. Wood density, 319 here recorded as the wood dry mass divided by its green volume, was assigned to each tree using the lowest available taxonomic level of botanical identifications (i.e. species or genus) 320 321 and the corresponding average wood density recorded in the Global Wood Density Database 322 (GWDD, Chave et al., 2009; Zanne et al., 2009). Botanical identification was harmonized 323 through the Taxonomic Names Resolution Service (http://tnrs.iplantcollaborative.org), for both plot inventories and the GWDD. For trees not identified to species or genus (~5%), we used 324 plot-average wood density. We estimated heights of all trees using Chave et al.'s (2014) pan-325 326 tropical diameter-height model which accounts for heterogeneity in the D-H relationship using an environmental proxy: 327

328 $Ln(H) = 0.893 - E + 0.760 ln(D) - 0.0340 ln(D)^{2} (3)$

Where *D* is the diameter at 130 cm and *E* is a measure of environmental stress (Chave *et al.*,
2014). For sites with tree height measurements (N=20), we developed local D-H models, using

a Michaelis-Menten function (Molto *et al.*, 2014). We used these local models to validate the predicted Lorey's height (i.e. plot average height weighted by BA) from the largest trees, of which height has been estimated with a generic H-D model (equation 3, Chave et al. 2014).

We estimated plot biomass as the sum of the biomass of live tree with diameter at 130 cm superior or equal to 10 cm, using the following pan-tropical allometric model (Réjou-Méchain *et al.*, 2017):

337 AGB=exp(-2.024-0.896E+0.920ln(WD)+2.795ln(D)-0.0461(ln(D²))) (4)

338 Plot-level metric estimation from the largest trees

The relationship between each plot metric, namely basal area (BA), the quadratic mean 339 diameter (Dg), Lorey's height (H_{BA} the mean height weighted by the basal area) and the 340 community wood density (WD_{BA} ; the mean wood density weighted by the basal area), and 341 342 those derived from largest trees was determined using an iterative procedure following Bastin et al. (2015). Trees were first ranked by decreasing diameter in each plot. An incremental 343 procedure (i.e. including a new tree at each step) was used to sum or average information of 344 345 the *i* largest trees for each plot metric. Each plot-level metric was predicted by the respective metric derived from the ith largest trees. For each increment, the ability (goodness of fit) of the 346 347 *i* largest trees to predict a given plot-metric was tested through a linear regression. To avoid overfitting, a Leave-One-Out procedure was used to develop independent site-specific models 348 (N=118). Specifically, the model to be tested at a site was developed with data from all other 349 350 sites. Errors were then estimated as the relative root mean square error (rRMSE) computed 351 between observed and predicted values (X):

352
$$rRMSE = \sqrt{\sum \frac{(Xobs - Xpred)^2}{n}} / \bar{X}$$
 (5)

The form of the regression model (i.e. linear, exponential) was selected to ensure a normal distribution of the residuals.

To estimate plot basal area, we used a simple power-law constrained on the origin, as linear model resulted in non-normal residuals. Plot-level basal area (BA) was related to the basal area for the *i* largest trees (BA_i) using: 358 BA = $b_1 \Sigma BA_i^{\gamma 1}(6)$

To estimate the quadratic mean diameter, Lorey's height and the wood density of the community, we used simple linear models relating the plot-level metrics and the value of the metrics for the *i* largest trees:

$$362 \qquad D_g = a_2 + b_2 D_{gi} (7)$$

363
$$H_{BA} = a_3 + b_3 H_i$$
 (8)

364
$$WD_{BA} = a_4 + b_4 WD_i$$
 (9)

Both Lorey's height (H_{BA}) and the average height ($\overline{H_i}$) of the ith largest trees depend on the same D-H allometry, which always contains uncertainty whether we use a local, a continental or a pan-tropical model. To test the dependence of the prediction of H_{BA} from $\overline{H_i}$ on the allometric model, we used measurement from Malebo in the Democratic Republic of the Congo, where all heights were measured on the ground (see supplementary figure 2).

The quality of the predictions of plot-level metrics from the largest trees is quantified using the relative root mean square error (rRMSE) between measured and predicted values, and displayed along the cumulated number of largest trees. Model coefficients are estimated for each metric derived from the largest trees (N_{LT}) and averaged across the 118 models (see supplementary table 2).

375 Mean rRMSE is plotted as a continuous variable, while its variation is presented as a 376 continuous area between 5th and the 95th percentiles of observed rRMSE.

377 The optimal number of largest trees for plot-level biomass estimation

The optimal number of largest trees N_{LT} was determined from the prediction of each plot-level metric considered above, i.e. keeping a small number of trees while ensuring a low level of error for each structural parameter. We then predicted plot-level biomass from the N_{LT} model (equation 2). The final error was calculated by propagating the entire set of errors related to equation 4 (Réjou-Méchain *et al.*, 2017) in the N_{LT} model (i.e. error associated to each allometric model used). The model was then cross-validated across all plots (N=867).

384 Investigating residuals: what the largest trees do not explain

To understand the limits of predicting AGB through N_{LT}, we further investigated the relationship 385 386 between AGB residuals and key structural and environmental variables using linear modelling. 387 Forest structure was investigated through the total stem density (N), the quadratic mean diameter (Dg), Lorey's height (H_{BA}) and community wood density (WB_{BA}). As environmental 388 data, we used the mean annual rainfall and the mean temperature computed over the last 10 389 years at each site using the Climate Research Unit data (New et al., 1999, 2002), along with 390 391 rough information on soil types (Carré et al., 2010). Major soil types were computed from the soil classification of the Harmonized World Soil Database into IPCC (intergovernmental panel 392 on climate change) soil classes. In addition, considering observed differences in forest 393 structure across tropical continents (Feldpausch et al., 2011, 2012) and recent results on pan-394 tropical floristic affinities (Slik et al., 2015), we tested for an effect of continent (America, Africa 395 and Asia) on the AGB residuals. Differences in forest structure and AGB among continents 396 were also illustrated through the analysis of their distribution. 397

398 The importance of each variable was evaluated by calculating the type II sum of squares that 399 measures the decrease in residual sum of squares due to an added variable once all the other variables have been introduced into the model (Langsrud, 2003). Residuals were investigated 400 at both plot and site levels, the latter analyzed to test for any influence of the diameter structure, 401 which is usually unstable at the plot level due to the dominance of large trees on forest metrics 402 403 at small scales (Clark & Clark, 2000). Here we use a principal component analysis (PCA) to summarize the information held in the diameter structure by ordinating the sites along the 404 405 abundance of trees in each diameter class (from 10 to +100 cm by 10 cm bins).

407 Results

408 Plot-level metrics

409 Plot metrics averaged at the site level (867 plots, 118 sites) present important variations within and between continents. In our database, the quadratic mean diameter varies from 15 to 42 410 cm²ha⁻¹, the basal area from 2 to 58 m²ha-1, Lorey's height from 11 to 33 m and the wood 411 density weighted by the basal area from 0.48 to 0.84 gcm⁻³ (supplementary figure 1). Such 412 important differences between minimal and maximal values are observed because our 413 414 database cover sites with various forest types, from young forest colonizing savannas to old growth forest. However, most of our sites are found in mature forests, as shown by relatively 415 high average and median value of each plot metric (average aboveground biomass = 302 416 Mgha⁻¹; supplementary figure 1). In general, highest values of aboveground biomass are found 417 in Africa, driven by highest values of basal area and highest estimations of Lorey's height. 418 Highest values of wood density weighted by basal area are found in America. 419

420 Plot-level estimation from the i largest trees

421 Overall, plot metrics at 1-ha scale were well predicted by the largest trees, with qualitative agreement among global and continental models (Figure 2). When using the 20 largest trees 422 to predict basal area (BA) and quadratic mean diameter (Dg), the mean rRMSE was < 16% 423 and 12%, respectively (Figs 3a and 3b). Lorey's height (H_{BA}) and wood density weighted by 424 425 basal area (WD_{BA}) were even better predicted (Figs 3c and 3d), with mean rRMSE of 4% for 426 the 20 largest trees. The prediction of Lorey's height from the largest trees using local 427 diameter-height model (supplementary figure 2a) yielded results similar to those obtained using equation 3 of Chave et al. (2014). More importantly, it also yielded similar results to 428 prediction of Lorey's height from the largest trees using plots where all the trees were 429 430 measured on the ground (supplementary figure 2b). This suggests that our conclusions are robust to the uncertainty introduced by height-diameter allometric models. 431

432 AGB prediction from the largest trees

We selected "20" as the number of largest trees to predict plot metrics. The resulting model
predicting AGB (Mg ha⁻¹) based on the 20 largest trees is:

435 AGB = $0.0735 \times (Dg_{20}H_{20}WD_{20})^{1.1332}$ (rRMSE=0.179; R²=0.85; AIC= -260.18) (10)

436 Because the exponent was close to 1, we also developed an alternative and more operationa

437 I model with the exponent constrained to 1, given by:

438 AGB = $0.195 \times (Dg_{20}H_{20}WD_{20})$ (rRMSE=0.177; R²=0.85; AIC=-195) (11)

Ground measurements of plot AGB were predicted by our N_{LT} model with the exponent constrained to 1, with a total error of 17.9% (Figure 4), a value which encompass the error of the N_{LT} model and the error related to the allometric model chosen. The Leave-One-Out crossvalidation procedure yielded similar results (rRMSE=0.19; R²=0.81), validating the use of the model on independent sites.

444 Determining the cause of residual variations

The explanatory variables all together explain about 37% of the variance in AGB both at plot 445 446 and site levels when omitting the diameter structure, and about 63% at site level when included (Figure 5). In general, forest structure and particularly the stem density explained most of the 447 residuals (table 1; weights: 79% and 54% at plot- and site-level respectively). The stem density 448 449 was followed by a continental effect (weights: 18%, 28% and 1%, respectively for Africa, 450 America and Asia) and by the effect of H_{BA} and WD_{BA} (respective weights: 1% and 0% at the plot level, 0% and 11% at the site level, and 23% and 0% when accounting for the diameter 451 structure at the site level). Inclusion of the diameter structure provided the best explanation of 452 453 residuals, with 63% of variance explained, and a weight of 69% for the first axis of the PCA 454 (supplementary figure 3). This first axis of the PCA was related to the general abundance of 455 trees at a site, and in particular medium-sized trees (40-60cm). Among environmental variables, only rainfall was significantly related to the residuals at the site level when the 456 diameter structure was considered (2%). 457

458 Differences among continents

While diameter structure explained a large fraction of the residual variance of our global model, there was marked difference in forest structure across continents (Figure 6). Consequently, we investigated differences between continents in the distribution of residuals of the pantropical model (Figure 6a), in the relative contribution of the 20 largest trees to plot total

biomass (Figure 6b), and in the contribution to the total aboveground biomass per diameter 463 class (Figs. 6c-f). To this end, we considered the following four classes of diameter at 130 cm: 464 10 to 30 cm, 30 to 50 cm, 50 to 70 cm and above 70 cm. Results show that the prediction of 465 biomass from the 20 largest trees using the pan-tropical model tends to be slightly 466 overestimated in Africa (+ 3%) and underestimated in America (- 3%) and in Asia (-5%) (Figure 467 6a). The proportion of biomass is higher in high diameter class (over 70 cm) in Africa, in 468 469 intermediate diameter classes (between 30 and 70 cm) in America and is equally distributed among the different diameter classes in Asia (Figure 6 c-d). 470

471 Discussion

472 The largest trees, convergences and divergences between continents

473 Sampling a few largest trees per hectare generally allows an unbiased prediction of four key 474 descriptors of forest structures across the tropics. There is generally no improvement in predicting biomass, quadratic mean diameter, Lorey's height (H_{BA}) or community wood density 475 beyond the first 10-to-20 largest trees (Figure 2, Figure 3a). But when a forest plot presents 476 an abundant number of large trees (Figure 5d), increasing the number of trees sampled does 477 478 improve the model's accuracy. This is due to the fact that the higher total AGB in a plot, the 479 lower the proportion of total AGB encompassed by the largest trees. This is particularly true for BA for which rRMSE continues to decrease up to 100 largest trees (Figure 2a). In contrast, 480 Lorey's height predictions are altered when a large number of trees are included (Figure 2c), 481 i.e. when smaller, often suppressed, trees draw the average down (Farrior et al., 2016). This 482 might explain why the prediction of AGB does not mirror that of basal area (Figure 2b, Figure 483 3a), and suggest that the number of largest trees shall be set independently to each predictor 484 485 considered. Interestingly, the evolution of relative error in AGB prediction as a function of the 486 number of largest trees considered does not follow the same path between continents. For 487 instance, the error of prediction saturates more quickly in Africa and Asia than America. Investigation of residuals showed that the diameter structure (Figure 5c, supplementary Figure 488 489 3b), and in particular the number of medium size trees (Figure 5d), drives variability in AGB 490 predictions. It is therefore not surprising to see that in our dataset the site with higher levels of 491 underestimations is the one with the highest number of medium size trees, which is found in Asia in the Western Ghats of India. 492

The good performance of models based on the 20 largest trees in predicting Lorey's height and community wood density at site level was not surprising. Both metrics were indeed weighted by basal area, driven de facto by the largest trees. Their consistency across sites and continents was not expected though, which emphasize the generality of our approach.

The predictability of plot-level forest structure metrics from the largest trees implies that characteristics of smaller trees do not vary completely independently from those of the larger trees. For example, plots where the largest trees have low basal area tend to have low plotlevel basal area (Figure 3a), meaning that the total size of the smaller trees is sufficiently constrained so that it does not compensate for the small size of the largest trees. Such constraints could arise through size-frequency distributions being set by allometric scaling rules (Enquist et al., 2009), or could be due to the largest trees responding in the same way as the remaining smaller trees to environmental drivers.

Despite the general consistency of these relationships across continents, slight differences are 505 506 evident when comparing the pan-tropical model residuals across continents (Figure 6, supplementary figure 4). These differences indicate biogeographic variation in forest structure. 507 508 In America, our pan-tropical model tends to slightly underestimate basal area (mean: -5%) and overestimate Lorey's height (mean: +3%) (supplementary figure 4). This suggests that large 509 510 trees make up a smaller proportion of basal area in America and that for a given diameter we find higher trees (supplementary figure 2), the later confirming that the shape of height-511 diameter allometries varies between continents (Banin et al., 2012; Sullivan et al., 2018). In 512 513 Africa, large trees (i.e. DBH > 70 cm) are more abundant and account for a large fraction of 514 plot biomass (figure 6f). This supports previous observations that African forests are 515 characterized by fewer but larger stems (Feldpausch et al., 2012; Lewis et al., 2013), while forests in the Americas have more stems but generally have lower biomass (Sullivan et al., 516 517 2017). In Asia, the distribution of the biomass across diameter classes appears more balanced 518 (Figure 6c-f). Such differences in forest structure, even if being quite limited, suggest tropical 519 forests differ between continents in terms of dynamics, carbon cycling, response and feedback to climate and resilience to external forcings (e.g. climate change, forest degradation and 520 521 deforestation).

Interestingly, while a recent global phylogenetic classification of tropical forest groups American with African forests vs. Asian forests (Slik et al., 2018), our study of forest structure properties tends more to single out American forests, and particularly highlight the contrast in between African and American forests. Although this deserves further investigations, it might reveal a lack of close relationship between forest structure properties and phylogenic similarity,

which echoes recent results on the absence of relationship between tropical forest diversityand biomass (Sullivan et al., 2017).

529 Largest trees, a gateway to global monitoring of tropical forests

530 Revealing the predictive capacity held by the largest trees, our results constitute a major step forward to monitor forest structures and biomass stocks. The largest trees in tropical forests 531 can therefore be used to accurately predict various ground-measured properties (i.e. the 532 guadratic mean diameter, the basal area, Lorey's height and community wood density), while 533 534 previous work has predicted only biomass "estimates" (e.g. Slik et al., 2013; Bastin et al., 2015). Our approach allows us to (i) describe forest structure independently of any biomass 535 allometric model (ii) and integrates environmental-based variations in D-H relationship, known 536 to vary locally (Feldpausch et al., 2011; Kearsley et al., 2013;). It is also (iii) relatively 537 538 insensitive to differences in floristic composition and community wood density (Poorter et al., 539 2015).

Furthermore, the "largest trees" models were developed for each plot-level metric and for any 540 541 number of largest trees. Thus, they do not rely on any arbitrary threshold of tree diameter. Note 542 that the optimal number of largest trees to be measured (i.e. 20) was set for demonstration 543 and can vary depending on the needs and capacities of each country or project (see supplementary table 2). In the same way, local models could integrate locally-developed 544 biomass models, when available. Consequently our approach (i) can be used in young or 545 546 regenerating un-managed forests with a low "largest tree" diameter threshold and (ii) is 547 compatible with recent remote sensing approaches able to single out canopy trees and describe their crown and height metrics (Ferraz et al., 2016; Coomes et al., 2017). 548

549 Aboveground biomass model from the largest trees, a multiple opportunity

Globally, the N_{LT} model for the 20 largest trees allows plot biomass to be predicted with 17.9%
error. This result is a pan-tropical validation of results obtained in Central Africa (Bastin *et al.*,
2015). It opens new perspectives towards cost-effective methods to monitor forest structures
and carbon stocks through largest trees metrics, i.e. metrics of objects directly intercepted by
remote-sensing products.

Developing countries willing to implement Reduction of Emissions from Deforestation and 555 Forest Degradation (REDD+) activities, shall also report on their carbon emissions and develop 556 557 a national reference level (IPCC, 2006; Maniatis & Mollicone, 2010). However, most tropical 558 countries lack capacities to assume multiple, exhaustive and costly forest carbon inventories (Romijn et al., 2012). By measuring only a few large trees per hectare, our results show that it 559 is possible to obtain unbiased estimates of aboveground C stocks in a time and cost-efficient 560 manner. Assuming that 400 to 600 trees D > 10 cm are measured in a typical 1-ha sample 561 562 plot, monitoring only 20 trees is a significant improvement. Although finding the 20 largest trees in a plot of several hundred individuals requires evaluating more than 20 trees, in practice, a 563 conservative diameter threshold could be defined to ensure that the 20 largest trees are 564 sampled. An alternative approach could also be found in the development of relascope-based 565 566 approach adapted to detection of the largest trees in tropical forests. Using such approach would facilitate rapid field sampling in extensive areas to produce large scale AGB estimates. 567 Those could fulfil the needs in calibration and validation of current and forthcoming space 568 569 missions focused on aboveground biomass.

570 Our findings also point towards the potential effectiveness of using remote sensing techniques 571 to characterize canopy trees for inferring entire forest stands attributes. Remote sensing data could be used for direct measurement (e.g. tree level metrics such as height, crown width, 572 573 crown height) of the largest trees as a potential alternative to indirect development of complex 574 metrics (e.g. mean canopy height, texture) used to extrapolate forest properties. While the use 575 of single-tree approach has shown some limitations to extrapolate plot metrics (Coomes et al., 2018), we have still to investigate their potential to identify largest trees. Some further 576 refinements are needed, but most of the tools required to develop "largest trees" models are 577 578 readily available. In particular, Ferraz et al. (2016) developed an automated procedure to locate single trees based on airborne LiDAR data, to measure their height and crown area. Crown 579 area could further be linked to basal area, as the logarithm of crown area is consistently 580 correlated with a slope of 1.2-1.3 to the logarithm of tree diameter across the tropics (Blanchard 581 et al., 2016). Regarding wood density, hyperspectral signature and high resolution topography 582

583 offers a promising way to assess functional traits remotely (e.g. Asner *et al.*, 2017; Jucker *et al.*, 2018) which could potentially provide proxies of wood density. Alternative approaches 585 could focus on the development of plot-level AGB prediction by replacing the basal area of the 586 largest trees with their crown metrics. While the measurement of crown areas has yet to be 587 generalized when inventorying plots, several biomass allometric models already partition trunk 588 and crown mass (Ploton *et al.*, 2016; Coomes *et al.*, 2017; Jucker *et al.*, 2017).

The main limitation of our approach lies in the limited inference that can be made on the 589 590 understory and sub-canopy trees. We show that most of the remaining variance is explained by variations in diameter structures, and in particular among the total stem density. 591 Interestingly, stem density was generally identified as a poor predictor of plot biomass in 592 tropical forests (Slik et al., 2010; Lewis et al., 2013). However, our results show that stem 593 density explains most of the remaining variance (Table S1). This suggests that, in addition to 594 trying to understand large-scale variations in large trees and other plot metrics, which can be 595 596 directly quantified from remote sensing, we should also put more effort into understanding 597 variation in smaller trees, which mainly drives total stem density and the total floristic diversity. 598 Smaller trees are also essential to characterize forest dynamics and understand changes in 599 carbon stocks. Several options are nonetheless possible from remote sensing, considering the 600 variation in lidar point density below the canopy layer (D'Oliveira et al., 2012), the distribution 601 of leaf area density (Stark et al., 2012, 2015; Tang & Dubayah, 2017; Vincent et al., 2017) or 602 the use of multitemporal lidar data to get information on forest gap generation dynamics and 603 consequently on forest diameter structure (Kellner et al., 2009; Farrior et al., 2016).

604 Large trees in degraded forests

If large trees are a key feature of unmanaged forests, they are conspicuously absent from managed or degraded forests. Indeed, large trees are targeted by selective or illegal logging, and are the first to disappear or to suffer from incidental damages when tropical forests are exploited for timber (Sist *et al.*, 2014). The loss of largest trees drastically changes forest structures and diameter distributions, and their loss is likely to counteract the consistency in forest structures observed through this study. Understanding how, or whether, managed forests deviate from our model predictions could help characterize forest degradation, which accounts for a large fraction of carbon loss worldwide (Baccini *et al.*, 2017), acknowledging that rapid post-disturbance biomass recovery (Rutishauser *et al.*, 2015) will remain hard to capture.

615 Conclusion – towards improved estimates of tropical forest biomass

The acquisition, accessibility and processing capabilities of very high spatial, spectral and 616 temporal resolution remote sensing data has increased exponentially in recent years (Bastin 617 618 et al., 2017). However, to develop accurate global maps, we will have to obtain a greater number of field plots and develop new ways to use remote sensing data. Our results provide 619 620 a step forward for both by (i) drastically decreasing the number of individual tree measurements required to get an accurate, yet less precise, estimate of plot biomass and (ii) opening the way 621 622 to direct measurement of plot metrics measured from remote sensing to estimate plot biomass. As highlighted by Clark and Kellner (2012), new biomass allometric models relating plot-level 623 biomass measured from destructive sampling and plot-level metric measured from remote-624 625 sensing products should be developed, as an alternative to current tree-level allometric 626 models. Such an effort will largely lower operational costs and uncertainties surrounding 627 terrestrial C estimates, and consequently, will help developing countries in the development of national forest inventories and aid the scientific community in better understanding the effect 628 of climate change on forest ecosystems. 629

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649 manuscript.

650 **Contributions**

J.F.Bastin and E.Rutishauser conceptualized the study, gathered the data, performed the analysis and wrote the manuscript. All the co-authors contributed by sharing data and reviewing the main text. A.R.Marshall, J.Poulsen and J.Kellner revised the English.

654 **Conflict of interest**

The authors declare there is no conflict of interest associated to this study.

656 Figures

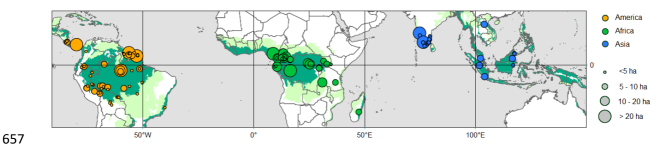
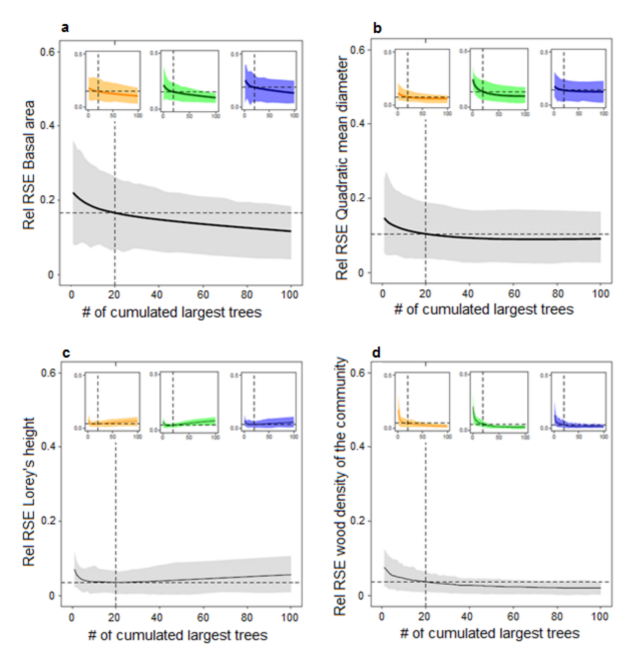
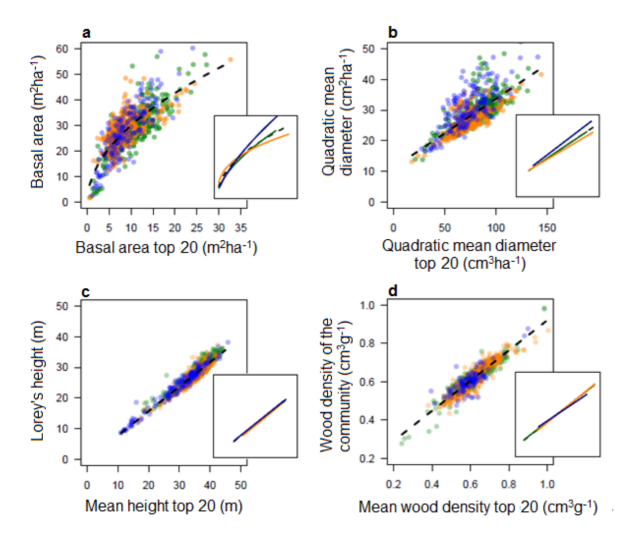


Figure 1. Geographic distribution of the plot database. We used 867 plots of 1 hectare from 118 sites. Dots are colored according to floristic affinities (Slik et al. 2015), with America, Africa and Asia respectively in orange, green and blue. They are also sized according the total area surveyed in each site. In the background, moist forests are displayed in dark green and dry forest in light green.



663

664 Figure 2. Quality of the prediction of plot metrics from largest trees. Variation of the 665 relative Root Mean Square Error (rRMSE) of the prediction of plot metric from i largest trees versus the cumulative number of largest trees for (a) basal area, (b) quadratic mean diameter, 666 (c) Lorey's height and (d) wood density weighted by the basal area. Results are displayed at 667 the pan-tropical level (main plot in grey) and at the continental level (subplots; orange = 668 America; green = Africa; blue = Asia). The solid line and shading shows the mean rRMSE and 669 the $\mathbf{5}^{th}$ and the $\mathbf{95}^{th}$ percentiles. Dashed lines represent the mean rRMSE observed for each 670 671 model, when considering the 20 largest trees.



672

673 Figure 3. Prediction of plot metrics (y-axis) from the 20 largest trees (x-axis). Results are shown for (a) basal area, (b) quadratic mean diameter, (c) Lorey's Height and (d) wood density 674 weighted by the basal area. Each dot corresponds to a single plot, colored in orange, green 675 and blue for America, Africa and Asia respectively. Both pan-tropical (black dashed lines) and 676 677 continental (coloured lines) regression models are displayed. These results show that substantial part of remaining variance, i.e. not explained by largest trees, is found when 678 predicting the basal area and the quadratic mean diameter, with slight but significant 679 differences between continents. 680

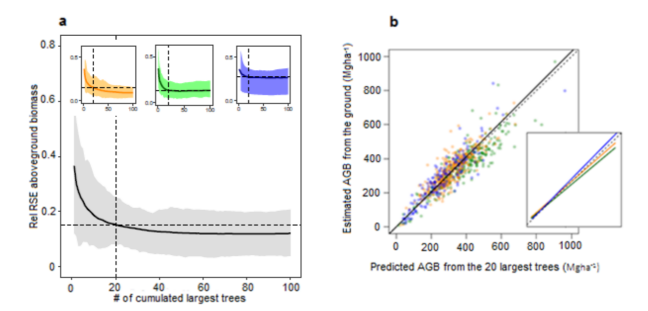




Figure 4. Prediction of AGB from plot metrics of the 20 largest trees. Results are shown for the 867 plots, among the three continents colored orange, green and blue for America, Africa and Asia respectively. The regression line of the model is shown as a continuous black line while the dashed black line shows a 1:1 relationship. The figure shows an unbiased prediction of AGB across the 867 plots, with slight but significant differences between the 3 continents.

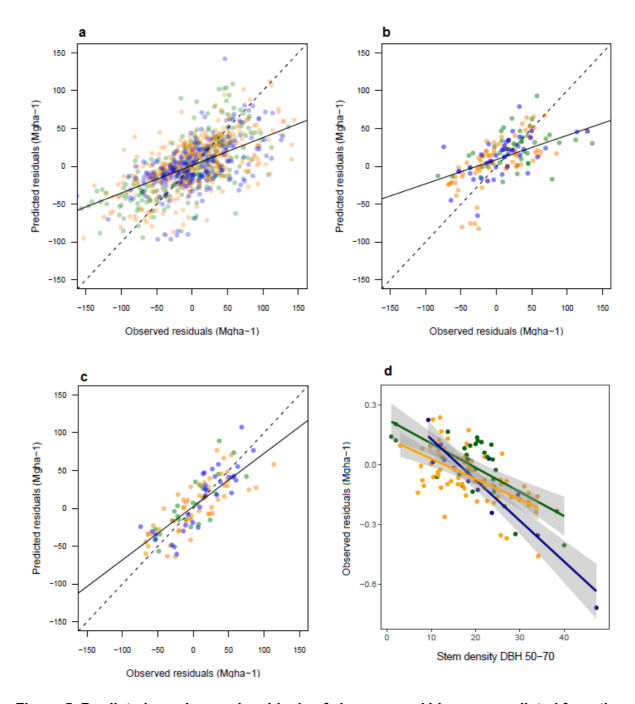
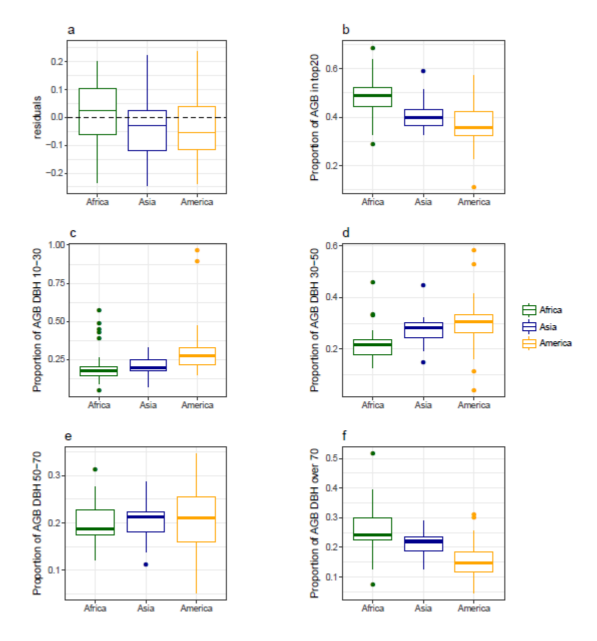


Figure 5. Predicted vs. observed residuals of aboveground biomass predicted from the **20 largest trees.** Residuals are explored at three different levels: (a) plot, (b) site [without considering the diameter structure as an explanatory variable], (c) site [considering the diameter structure] and (d) along the stem density of medium size trees. America, Africa and Asia are colored in orange, green and blue respectively. The figures show a good prediction of residuals in (a) and (b), driven by stem density, and a less biased prediction in (c), driven by

- the diameter structure. Variance of observed residuals are also well explained by the stem
- density of medium size trees (d), which mainly drive the first axis of the PCA.



697

Figure 6. Comparison across continents of aboveground biomass prediction per site and their 698 contribution to different share of the diameter structure. Africa, Asia and America, are colored 699 700 in green, blue and orange, respectively. The distribution of the residuals of pan-tropical aboveground biomass prediction from the 20 largest trees (a) shows predictions are slightly 701 overestimated in Africa (+3%), and slightly underestimated in Asia (-3%) and America (-5%). 702 The proportion of aboveground biomass in the 20 largest trees (b) is highest in Africa (48%), 703 704 followed by Asia (40%) and America (35%). The decomposition across four diameter classes 705 (c-f, i.e. from 10 to 30, 30 to 50, 50 to 70 and beyond 70 cm) of their relative share of the total 706 biomass shows that most of the biomass is found in the large trees in Africa, and in the small

- to medium trees in America. Asia presenting a more balanced distribution of biomass across
- the diameter structure.

709 Tables

Table 1. Weight of each variable retained for the explanation of AGB residuals. Weights are calculated as a type II sum of squares, which measures the decreased residual sum of squares due to an added variable once all the other variables have been introduced into the model. Results are shown for the exploration of residuals at the plot and at the site level, with and without consideration of the diameter structure. Weights are dominated by structural variables, and in particular the stem density and the diameter structure. Height, wood density and continent have also a non-negligible influence on residuals.

Level of residual	Parameter	Weight	717
Plot			
	Stem density*	79	
	Continent*	18	
	Lorey's height*	1	
	Major soil types	1	
	Temperature	1	
	Wood density weighted by the basal area	0	
	Rainfall	0	
Site without			
diametric structure			
	Stem density*	54	
	Continent*	28	
	Wood density weighted by the basal area*	11	
	Rainfall	3	
	Major soil types	3	
	Temperature	2	
	Lorey's height	0	
Site with diametric			
structure			
	PCA axis 1*	69	
	Lorey's height*	23	
	Rainfall*	3	
	Major soil types	3	
	Continent	1	
	Temperature	1	
	Wood density weighted by the basal area	0	
	PCA axis 2	0	

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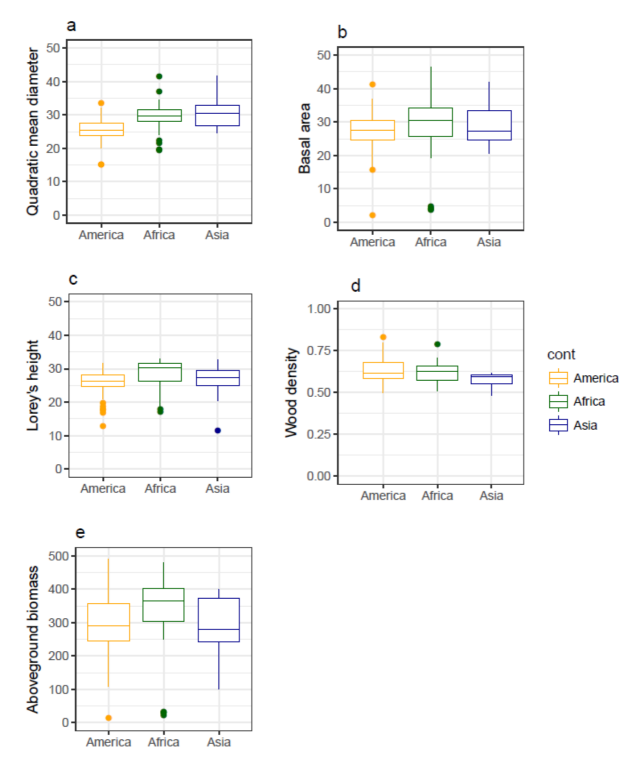
1110 Data accessibility

- 1111 Data for plots in the CTFS network are available through the online portal at
- 1112 <u>http://www.forestgeo.si.edu;</u> in the Forestplot network at <u>https://www.forestplots.net/</u> and in
- 1113 the TEAM network at <u>http://www.teamnetwork.org/</u>.

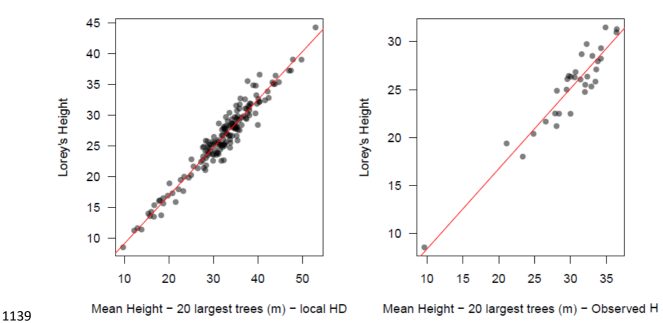
1114 Biosketches

Jean-Francois Bastin is a Post-Doctoral Fellow of the Crowther Lab at the Institute of Integrative Biology, Department of Environmental Systems Science, ETH-Zurich. He is an ecologist and a geographer using remote sensing to study the effect of global change on terrestrial ecosystems.

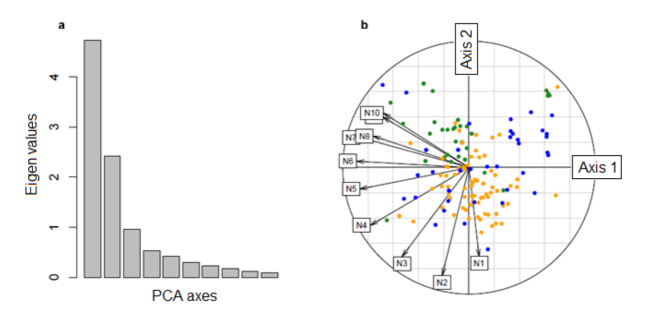
- 1119 Ervan Rutishauser is a Post-doctoral Fellow at Smithsonian Tropical Research Institute, 1120 broadly interested in understanding environmental resilience to natural or human-induced 1121 disturbances. He aims at providing rigorous and practical evidences to trigger a shift towards 1122 better ressources management and conservation.
- 1123 **Supplementary information.**
- 1124 Supplementary table 1. Plot, Site and Pls
- 1125 Supplementary table 2. Coefficients of plot level structure prediction from the *ith*
- 1126 largest trees.
- 1127 Supplementary figure 1. Cross-continent comparison of plot-metrics distribution
- 1128 averaged at the site level.
- 1129 Supplementary figure 2. Lorey's Height prediction from the 20 largest trees.
- 1130 Supplementary figure 3. PCA on the diameter structure and corresponding mean
- distribution for high contributions of axis 1 and axis 2.
- 1132 Supplementary figure 4. Cross-continent comparison of the relative residuals from the
- 1133 prediction of plot-metrics from the 20 largest trees.



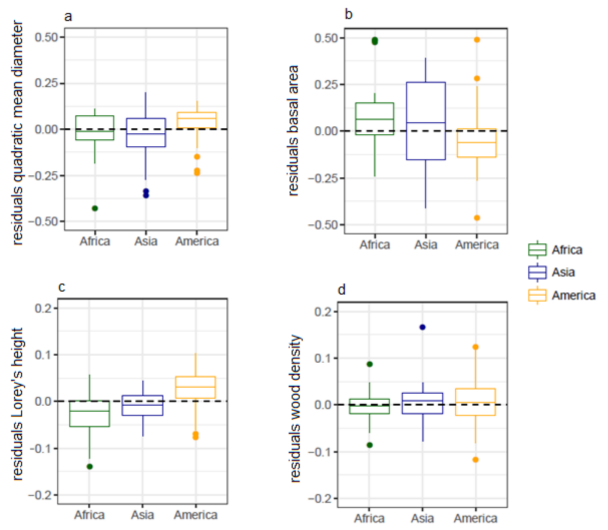
1134 1135 Supplementary figure 1. Cross-continent comparison of plot-metrics distribution averaged at the site level. Figures illustrates respectively the distribution of the values for the 1136 1137 quadratic mean diameter (a), basal area (b), Lorey's height (c), wood density (d) and 1138 aboveground biomass (e).



1140 **Supplementary figure 2. Lorey's Height prediction from the 20 largest trees.** Figures 1141 show the results using (i) local D-H allometries for 20 sites (left subfigure) and (ii) using plots 1142 where height is measured on all trees in Malebo site in the Democratic Republic of the Congo 1143 (right subfigure).



Supplementary figure 3. PCA on the diameter structure and corresponding mean 1145 1146 distribution for high contributions of axis 1 and axis 2. (A) Illustration of top and low 1147 percentile observed for each axis, with diameter distributions represented as the relative 1148 difference with the average observed distribution.(B) Biplot with contribution to the PCA of all 1149 the diameter classes, with the respective position of each site in the space defined by axis1 1150 and 2. Axis 1 is driven by differences in global abundance of trees and axis 2 is driven by a 1151 difference of balance between abundance of small vs. large trees. Colors represent continent, with Africa, America and Asia respectively in green, orange and blue. 1152



Supplementary figure 4. Cross-continent comparison of the relative residuals from the prediction of plot-metrics from the 20 largest trees. The relative residuals are generally low (<10%). Systematic small differences can however be found in America, where the quadratic mean diameter and Lorey's height tend to be slightly overestimated and the basal area slightly underestimated.</p>