1 Using terrestrial laser scanning to constrain forest ecosystem

2 structure and functions in the Ecosystem Demography model (ED2.2)

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15 Abstract. Terrestrial Biosphere Models (TBMs) are invaluable tools for studying plant-atmosphere interactions at multiple 16 spatial and temporal scales, as well as how global change impacts ecosystems. Yet, TBM projections suffer from large 17 uncertainties that limit their usefulness. Forest structure drives a significant part of TBM uncertainty as it regulates key 18 processes such as the transfer of carbon, energy, and water between the land and the atmosphere, but remains challenging to 19 observe and reliably represent. The poor representation of forest structure in TBMs might actually result in simulations that 20 reproduce observed land fluxes, but that fail to capture carbon pools, forest composition, and demography. Recent advances 21 in Terrestrial Laser Scanning (TLS) offer new opportunities to capture the three-dimensional structure of the ecosystem and 22 to transfer this information to TBMs in order to increase their accuracy. In this study, we quantified the impacts of 23 prescribing initial conditions (tree size distribution), constraining key model parameters with observations, as well as 24 imposing structural observations of individual trees (namely tree height, leaf area, woody biomass, and crown area) derived 25 from TLS into the state-of-the-art Ecosystem Demography model (ED2.2) at a temperate forest site (Wytham Woods, UK). 26 We assessed the relative contribution of initial conditions, model structure, and parameters to the overall output uncertainty 27 by running ensemble simulations with multiple model configurations. We show that forest demography and ecosystem 28 functions as modelled by ED2.2 are sensitive to the imposed initial state, the model parameters, and the choice of key model 29 processes. In particular, we show that:

- parameter uncertainty drove the overall model uncertainty with a mean contribution of 63% to the overall variance
 of simulated gross primary production;
- model uncertainty on the gross primary production was reduced fourfold when both TLS and trait data were
 integrated into the model configuration;
- land fluxes and ecosystem composition could be simultaneously and accurately simulated with physically realistic
 parameters when appropriate constraints were applied to critical parameters and processes.
- 36 We conclude that integrating TLS data can inform TBMs on the most adequate model structure, constrain critical parameters,
- 37 and prescribe representative initial conditions. Our study also confirms the need for simultaneous observations of plant traits,
- 38 structure and state variables if we seek to improve the robustness of TBMs and reduce their overall uncertainties.

40 1 Introduction

Terrestrial biosphere models (TBMs) are key tools to understand the ecosystem response to anthropogenic disturbances and climate change (Medvigy and Moorcroft 2012; McGuire et al. 2001). Nowadays they are intensively used, as is or embedded in Earth system models, to study plant-atmosphere interactions and predict the future of ecosystems facing global change (e.g., Poulter et al. 2010). Yet, the usefulness of TBMs is currently limited by the large uncertainties in their projections which originate from different sources (Lin et al. 2011).

Forest structure has long been recognized as a critical component to understand forest dynamics (Hurtt et al. 2010). It influences the climatically important fluxes of carbon, energy, and water (Bonan 2008). Yet, its realistic representation is challenging and an urgent priority in the development of next-generation TBMs (Fisher et al. 2018). The representation of the forest structure within TBMs is associated with three sources of uncertainty: model structure, model initialisation, and model parameter uncertainty.

51 The model structure entails by definition all the processes included in a model, how they are implemented, and all the 52 underlying assumptions (Bonan 2019). Model structure complexity varies among TBMs and also depends on the user 53 configuration choices: different formulations of the same process can co-exist within a TBM. This complexity results from 54 the necessary compromise between an accurate representation of reality on the one hand and the computational demand and 55 observational requirements on the other (Shiklomanov et al. 2020). Model intercomparison studies have demonstrated that 56 discrepancies in the representation of key processes such as forest structure (Fisher et al. 2018) or photosynthesis (Rogers et 57 al. 2017) lead to significant uncertainties in the projections of critical variables such as the overall land carbon sequestration 58 capacity (Friedlingstein et al. 2014: Lovenduski and Bonan 2017: Friedlingstein et al. 2006).

59 The initialisation uncertainty reflects the error made when determining the initial conditions of the modelled ecosystem. 60 Several approaches exist for initialising TBMs, the most common of which is probably to start runs from near-bare ground 61 conditions, force the simulations with relevant climate-forcings, and wait for the model to reach an equilibrium state, the so-62 called potential vegetation (Antonarakis et al. 2011). Yet, such a spin-up approach does not guarantee reliable initial 63 demography, carbon pools, or ecosystem structure. Alternatively, forest inventories can be used to prescribe the initial 64 composition of the ecosystem (Medvigy et al. 2009). The derivation of the initial states of critical variables, such as the 65 aboveground biomass or the total leaf area from the plant size distribution, then relies on model default allometries which are 66 often derived from other, potentially non-representative site-specific data.

Parameter uncertainty arises among other things from the necessary simplification of the natural complexity into a coherent list of model parameters, the uncertainty in the measurements used to calibrate the model, or the methods used to upscale local measurements to scales on which TBMs operate (Zaehle et al. 2005). Previous sensitivity analyses have underlined the critical importance of parameter uncertainty for the projections of ecosystem demography and productivity (Dietze et al. 71 2014; Massoud et al. 2019; Raczka et al. 2018; Wramneby et al. 2008). In a recent comparative study, parameter uncertainty 72 was even shown to dominate the overall model uncertainty over process uncertainty (Shiklomanov et al. 2020). Among 73 model parameters, allometric coefficients scale the shape and mass of the plants or of its components with their size (Chave 74 et al. 2014). Not surprisingly, multiple TBMs were shown to be sensitive to such allometric parameters (Collalti et al. 2019; 75 Cano et al. 2020; Esprey et al. 2004). Parameter uncertainty can be reduced by constraining the range of variation of model 76 parameters through the assimilation of different sources of observations or via model optimization (LeBauer et al. 2013). In 77 the past, TBMs have often been calibrated with eddy covariance data (Fer et al. 2018; Rezende et al. 2016; Collalti et al. 78 2016). While this approach ensures that the model correctly reproduces the short timescale (diurnal/seasonal) dynamics of 79 land fluxes, it does not ensure an accurate representation of forest structure and carbon pools. This is especially true because 80 forest structure-related parameters can present a low sensitivity to those observations (LeBauer et al. 2013; Richardson et al. 81 2010), and the equifinality in TBMs (Luo et al. 2009) can lead to acceptable land fluxes with a poor representation of 82 ecosystem structure (i.e. fluxes can be reproduced from an almost infinite range of structural possibilities, some of which 83 will be much more likely than would be others).

84 Among the different sources of observations used to reduce model uncertainties, remote sensing from various platforms 85 (terrestrial, air- and space borne) has increasingly been used to monitor and understand terrestrial ecosystems (Jones and 86 Vaughan 2010). LiDAR (Light Detection And Ranging) -data in particular have been used in the past to initialise forest biomass and constrain predictions of TBMS (Thomas et al. 2008; Hurtt et al. 2019). The recent revolution in Terrestrial 87 88 Laser Scanning (TLS, also called terrestrial LiDAR) provides new opportunities for constraining TBMs, and reducing the 89 uncertainties related to the vegetation structure representation (Fischer et al. 2019). The ability of TLS to measure the 90 distance to reflecting surfaces was initially used in ecological studies to measure simple metrics like DBH and tree heights 91 (Maas et al. 2008; Hopkinson et al. 2011). Since then, TLS methods have rapidly evolved to derive more complex metrics, 92 such as the vertical profiles of the forest structure (Jupp et al. 2009; Calders et al. 2018) and whole-tree volumetric 93 assessments (Fan et al. 2020), leading to an accurate determination of forest structure across various forest types (Calders et 94 al. 2015; Tanago et al. 2018; Takoudjou et al. 2018; Ehbrecht et al. 2017; Stiers et al. 2018; Saarinen et al. 2021.). Today, the 95 ability of TLS to accurately represent the 3D structure of forests via quantitative structure modelling (OSM), see Raumonen 96 et al. (2013) and Hackenberg et al. (2015) represents a unique opportunity to improve our understanding of forest ecosystems 97 under changing climates (Calders et al. 2020). In particular, TLS snapshots of vegetation ecosystems could simultaneously 98 provide important state variables to initialise TBMs, strong constraints to some critical allometric parameters, and help 99 determine the most appropriate model structure for some key processes.

In this study, we evaluated the relative contribution of different sources of uncertainty (parameters, processes, initial conditions) to the overall uncertainty of multiple simulated outputs of a specific TBM, namely the Ecosystem Demography model version 2 (ED2.2). We also explored the benefits of constraining vegetation structure related parameters and processes using TLS on the model performance and output variability. To do so, we ran ED2.2 simulation ensembles for a temperate forest in the UK considering different initial states for the modelled ecosystem, and varying multiple model parameters and process settings with or without TLS constraints. In other words, we assessed: (i) the relative importance of the model structure, initialisation, and parameter uncertainties in the ED2.2 model representation of a temperate forest; (ii) the potential added value of TLS data for vegetation modelling. To the best of our knowledge, this study is the first attempt to constrain fuse TLS data and a TBM using TLS.

110 2 Material and Methods

111 2.1 Study site and data

112 **2.1.1** Study site

113 Wytham Woods is a mixed deciduous forest, predominantly broadleaved, covering approximately 40 ha. It is located 5 km 114 northwest of Oxford in southern England (Thomas et al. 2011), Owned by Oxford University, Wytham Woods has been part 115 of the UK Environmental Change Network (ECN) and of the Smithsonian Global Earth Observatory (SIGEO) network since 116 1992 and 2008, respectively, and has hosted numerous ecological studies (Savill et al. 2010). The site is classified as an 117 ancient semi-natural woodland (Hall et al. 2001), which means that the site has been continuously covered by trees through 118 recorded history (since at least 1600), occasionally managed, and experienced minimal intervention (i.e. no silvicultural 119 management) since WWII (Fenn et al. 2015). Over the 1993-2008 time period, the site was characterised by a mean annual 120 temperature of 10°C and a mean annual precipitation of 726 mm (Butt et al. 2009). The area we simulate in this study is a 121 1.4 ha forest plot nested within the 18 ha long-term monitoring site part of the ForestGEO global network of forest inventory 122 plots. This 140 m × 100 m area has a local SW-coordinate (0, 100) and local NE-coordinate (140, 200) boundary. The local 123 origin coordinate (0,0) was located with a differential GPS at Lat 51.7750579 and Lon -1.33904729.

124 2.1.2 Field inventory and Terrestrial Laser Scanning data

The studied plot was inventoried during the summer of 2016. All trees were located, measured, and identified at the species level. The plot is largely dominated by sycamore (Acer pseudoplatanus, 65.3% of the 815 inventoried trees in the 1.4 ha plot, see Table 1, Figure 1 and Supplementary Figure S1), ash (Fraxinus excelsior, 10.3% of the stems), and hazel (Corylus avellana, 8.2% of the stems). Oaks (Quercus robur) represent a limited fraction of the woody stems (4.3%) but disproportionately contribute (23.4%) to the total basal area as they mostly consist of large trees (Table 1 and Figure 1). From the inventory, tree DBH is 24.4 cm on average (DBH median is 19.8 cm), and ranges from 2.9 cm to 141.2 cm.

131 Three-dimensional forest structure data were collected using a RIEGL VZ-400 terrestrial laser scanner (RIEGL Laser 132 Measurement Systems GmbH) in leaf-on (June and July 2015) and leaf-off (December 2015 and January 2016) conditions 133 (Calders et al. 2018). The RIEGL instrument uses on-board waveform processing and records multiple return LiDAR data, 134 which improves vertical sampling (Lovell et al. 2003; Calders et al. 2014). Individual trees were extracted using treeseg 135 (Burt, et al. 2019), and their structure modelled with TreeQSM (Raumonen et al. 2013) with the leaf-off TLS point cloud. 136 Leaves were then added to the individual tree branches using both the leaf-off and -on TLS datasets with the FaNNI 137 algorithm (Åkerblom et al. 2018). Doing so, TLS allowed retrieving of individual tree height, aboveground woody biomass 138 (modelled through estimates of volume combined with species-specific wood density), and leaf area. In addition, the 139 individual tree crown area was computed from the vertical projection of the leaf-off point clouds of individual trees. For

more details, a complete description of the TLS data collection and forest stand reconstruction is available in Calders et al.(2018).

142 2.1.3 Flux tower data and species traits

Stand-scale carbon and water fluxes have been occasionally measured in Wytham Woods using the eddy covariance technique. We digitised the most recent (to our knowledge) data collection of CO_2 fluxes that was reported by Thomas et al. (2011) for the period May 2007-April 2009. To do so, we digitised the weekly mean values of ecosystem gross primary productivity (GPP), ecosystem respiration (R_{eco}), and net ecosystem productivity (NEP) from Figure 6 of the aforementioned reference using the Plot digitizer software (v.2.6.8, http://plotdigitizer.sourceforge.net/). For a more detailed description of the eddy covariance data (including the data frequency of the original data, and the data quality filtering), we refer the readers to the original publication by Thomas et al. (2011).

150 In addition, we extracted all existing records of specific leaf area (SLA) and maximum rate of carboxylation (V_{c.max}) for the 151 five most important species in Wytham woods (Acer pseudoplatanus, Corylus avellana, Crataegus monogyna, Fraxinus 152 excelsior, and Ouercus robur) from the TRY database (Kattge et al. 2020), see Table 1 (the complete list of references from 153 which the data originate is available in supplementary section 1). Individual traits were converted into ED2.2 units ($m^2 kg_c^{-1}$ 154 for SLA with a fixed leaf carbon content of 0.5 and μ mol_C m⁻² s⁻¹ for V_{c max}). V_{c max} data were also rescaled to the ED2.2 155 reference temperature (15°C) using the model default value for the temperature coefficient O10 of 2.4. Following Asner et al. (2017), we calculated the community-weighted mean (CWM) and community-weighted standard deviation (CWSD) for both 156 157 traits based on the species composition and species-level average values, using species basal area as weights:

$$CWM = \frac{\sum_{i=1}^{N} w_i \cdot x_i}{\sum_{i=1}^{N} w_i}$$
Equation (1)

158

$$CWSD = \sqrt{\frac{\frac{\sum_{i=1}^{N} w_i \cdot (x_i - CWM)^2}{(N-1)\sum_{i=1}^{N} w_i}}{N}}$$
Equation (2)

where N is the total number of species for which data was available in TRY for each trait x, x_i is the mean trait value for species i, and w_i is the species weight (here the basal area of species i).

161 Flux tower data were used as a validation dataset while the TRY data were used to constrain parameters of the TBM used in 162 this study and described just below.

163 **2.2 Model**

164 2.2.1 The terrestrial biosphere model ED2.2

ED2.2 is a terrestrial biosphere model that can simulate the vegetation dynamics of a wide range of ecosystems from boreal to tropical forests (Longo et al. 2019). It is a cohort-based, spatially implicit model that approximates the behaviour of an individual-based, spatially distributed vegetation model through a system of size- and age-structured partial differential equations (Moorcroft et al. 2001). ED2.2 integrates modules of plant growth, mortality, phenology, disturbance, hydrology, and soil biogeochemistry to predict e.g., the demography, the succession, and the dynamics of water and carbon within the simulated ecosystem.

In ED2.2, the inter- and intra-specific diversity is represented by a set of plant functional types (PFTs) that differ by their leaf physiology, phenology, growth and allocation strategies, mortality, and sensitivity to environmental conditions (D. Medvigy et al. 2009). The trees inventoried in Wytham Woods were classified as either mid- or late-successional temperate deciduous trees (see below for the reasoning of the mapping). These PFTs are cold-deciduous, i.e. leaf phenology is prognosed by the accumulation of growing degree-days (growing season) and chilling days (senescing season) (Longo et al. 2019). A comprehensive model description, including photosynthesis, allometries, radiative transfer, and phenology, is available in Longo et al. (2019).

178 The ED2.2 model is available at https://doi.org/10.5281/zenodo.3365659.

179 2.2.2 Model initialisation and forcings

In this study, the ED2.2 model was initialised using i) near-bare ground (NBG) initial conditions (i.e. seedlings only), ii) the field inventory, or iii) the TLS-reconstructed size distribution. In the latter two configurations, the 1.4 ha site was initially divided into 35 square patches of 20 x 20 m. These three types of initial conditions are referred to below as NBG, Census, and TLS respectively. Simulations were run for multiple years using the local forcing data of the corresponding years of the CRU-NCEP reanalysis dataset (Viovy 2018). Simulations were run for either five years (Census and TLS configurations) or the approximate age since the last large-scale disturbance (100 years, NBG configuration), see Table 5. Soil texture was set according to the dominant soil type (clay), based on site-level observation (Butt et al. 2009).

187 2.2.3 Allometries and model parameters

In ED2.2, the carbon made available from net assimilation is partitioned at the cohort level into the different plant pools according to DBH-dependent allometries (Longo et al. 2019). In other words, plant cohorts allocate the carbon assimilated through photosynthesis to living tissues (i.e. fine roots, sapwood, leaves, seeds), the non-structural storage pool, and the dead tissues (i.e. coarse roots, and aboveground woody biomass) depending on (i) a set of allometries and (ii) whether the plant carbon balance and environmental conditions are favourable for growth. In ED2.2, aboveground woody biomass, height, leaf

- biomass, and crown area are scaled through DBH-dependent allometries (Table 3). The ED2.2 default allometric models and
- parameters are defined according to Medvigy et al. (2009) for the leaf biomass and height, Dietze et al. (2008) for the crownarea. and Albani et al. (2006) for the aboveground woody biomass.

196 To estimate the relative contribution of the parameter uncertainty to the variability of the model outputs, we used parameter 197 distributions from previous ED2.2 parameter uncertainty studies (Dietze et al. 2014; Shiklomanov et al. 2020; Raczka et al. 198 2018; Viskari et al. 2019). We only targeted those parameters that were shown to significantly contribute to the overall 199 parameter uncertainties in the aforementioned studies (Table 4) and set the rest to their ED2.2 default values for all 200 simulations. For SLA and V_{c.max} in particular, we defined two types of parameter distributions: either relatively wide priors 201 as in the previous sensitivity analyses listed above (Table 4) or constrained posteriors generated by the trait meta-analysis of 202 the Predictive Ecosystem Analyzer (PEcAn) run with the existing data in TRY and without random effects, see (LeBauer et 203 al. 2013; Meunier et al. 2021; Raczka et al. 2018). The meta-analysis was informed by TRY data only. Those distributions 204 are referred to below as without or with TRY-constraints, respectively. The uncertainty of the allometric coefficients was 205 determined either by the range of variation of those parameters in the ED2.2 model for hardwood tree PFTs (NBG and 206 Census configurations) or by the posterior distributions of these parameters generated when fitting the TLS data (see below).

207 2.2.4 Model configurations

To assess the importance of the model structure uncertainty, we targeted processes that were shown to induce significant variability in the model outputs in previous studies (Shiklomanov et al. 2020). In detail, we ran the model with multiple combinations of the following configurations: (i) closed canopies versus crowns of finite radii; (ii) two-stream versus multiple-scatter canopy radiative transfer models (RTMs); (iii) static versus plastic (varying with available light level) SLA and $V_{c,max}$; and (iv) a single versus two plant functional types (Table 2).

By default in ED2.2, plant canopies are represented as infinitely thin flat crowns (a.k.a. complete shading or closed canopy) that virtually occupy the entire horizontal space of the patch in which the cohort is located. In an alternative configuration, cohorts are still stacked on top of each other but have a finite radius and hence tallest plants only partially shade the underlying cohorts. In other words, the crown sub-model of ED2.2 determines the nature of the light competition between cohorts. Closed canopies have been shown to dramatically suppress competition from sub-dominant PFTs and typically result in unrealistically homogeneous patches (Fisher et al. 2015) while understorey cohorts receive more incoming diffuse and direct light if finite crowns are simulated.

The second sub-model we investigated was the choice of RTM. In both options (two-stream and multi-scatter), the full vertical radiation profile within each patch is resolved as a function of the canopy structure (e.g. leaf and wood area, clumping) and the environmental conditions (e.g. incident solar radiation, solar angle) following the approach of CLM 4.5 (Oleson et al. 2013). Both RTMs differ in the numerical resolution of the radiative transfers. By default (two-stream), the special multi-canopy solution of the two-stream approximation for vegetation canopies (Sellers 1985) is used as described in Longo et al. (2019) while the multiple-scatter is derived from first principles by Zhao and Qualls (2005) to address the longknown issues and biases of the two-stream model (Wang 2003). The multiple-scatter configuration increases diffuse light levels in the understorey as compared to the default two-stream approach (Shiklomanov et al. 2020).

The third sub-model that we evaluated is related to trait plasticity. By default (static), all cohorts of a given PFT share the same set of parameters which do not evolve over time, in contradiction with well-documented intra-specific variability of plant traits with environmental conditions (e.g. Keenan and Niinemets 2016). In the alternative configuration (plastic), cohort SLA and $V_{c,max}$ respectively decrease and increase with light availability, following empirical relationships from the tropics (Lloyd et al. 2010).

Finally, we also evaluated the impact of simulating one or multiple PFTs by either classifying all trees in the Wytham Woods inventory as belonging to the mid-successional hardwood tree PFT of ED2.2 ($N_{PFT} = 1$) or according to a classification similar to the one of Dietze and Moorcroft (2011), ($N_{PFT} = 2$), supplemented by a clustering analysis of the allometric relationships derived from the TLS data (see below).

237 2.3 Analyses

238 2.3.1 Impact of TLS data on model allometries and initial conditions

239 We first compared the model default allometries with site-specific ones constrained from the TLS data. To do so, we fitted 240 the individual plant metrics (height, crown area, aboveground woody biomass, and leaf area) versus DBH relationships 241 derived from TLS with the set of equations used in ED2.2 (Table 2). More specifically, we fitted the parameters of the four 242 allometries of ED2.2 using a Bayesian approach and the 'brms' package of R (Bürkner 2017). To account for the uncertainty 243 of the data we repeated the same analysis multiple times (N = 100) using data random sampling with replacement and 244 aggregating the resulting allometric parameter posterior distributions. To convert the leaf area obtained from TLS into leaf 245 biomass, we used the CWM of SLA. We evaluated the quality of fit of the allometric models by computing the root-mean-246 square deviations (RMSD, van Breugel et al. 2011) normalised by the observed mean and the Watanabe information 247 criterion (WAIC) for all four allometric models (height, crown area, aboveground woody biomass, leaf biomass). We fitted 248 all allometric models using multiple possible species-to-PFT classifications and only retained the classifications that 249 minimised the WAIC for the configurations $N_{PFT} = 1$ and $N_{PFT} > 1$.

- To assess the relative importance of TLS for the model initialisation, we compared the tree size distributions obtained from the field inventory and the TLS data and computed the absolute and relative differences between both DBH distributions
- 252 (ground-truthing of TLS).

253 2.3.2 Ensemble runs

For each type of initial conditions (NBG, Census, and TLS), we ran ensembles of 500 simulations with parameters randomly sampled from the parameter distributions (Table 4) and with process configuration randomly selected from the different options (Table 5). Each ensemble was equally split between runs with (250) and without (250) TRY constraints on SLA and $V_{c,max}$. The same parameter samples and process configurations were used for all three types of initial conditions, and with and without TRY restrictions on SLA and $V_{c,max}$ to allow independently evaluating the impact of the initial conditions, TRYand TLS-constraints at specific parameter values.

260 2.3.3 Sensitivity analyses and variance decomposition

261 Finally, we assessed which processes and parameters contributed the most to the overall model variance by performing a 262 sensitivity and a variance decomposition analysis following Dietze et al. (2014) and Lebauer et al. (2013). This analysis 263 allows predicting the fraction of the variance in target output variables attributable to individual parameters and processes (or 264 "partial variance"). We chose as target output variables the ecosystem GPP during the most productive month (June) or over 265 the leaf-on season (May-October), the total leaf area index (LAI) and the understorey photosynthetically active radiation 266 (PAR) in leaf-on conditions, as well as the aboveground woody biomass at the end of the simulation. For the NBG 267 configuration, we also decomposed the variance of the total stem density (which is prescribed in the other two 268 configurations). Parameters included in the variance decomposition analyses were re-classified as belonging to one of these 269 three categories: allometric parameters, TRY-constrainable parameters (SLA and $V_{c,max}$), and others. All five years of the 270 Census and TLS configurations were kept for analysis while only the last five years of the NBG runs were considered. Note 271 that the variance partitioning algorithm that we used only attributes to the parameters and processes their direct effect: 272 interactions are not accounted for in the variance decomposition.

All analyses presented in this study were performed using R 3.6 (R Core Team, 2019). Code and supporting data (including initialization and setting files) for reproducing the results presented below are publicly available in Zenodo and have the permanent DOI 10.5281/zenodo.6363617.

3 Results

278 3.1 Impact of TLS data on model allometries and initial conditions

TLS-extracted and field inventory DBHs were very well correlated ($R^2 = 0.98$, slope of the inventory vs TLS linear model = 0.998, see supplementary Figure S4). The mean (resp. median) relative difference between the TLS and field inventory DBHs was -0.2% (resp. -1.7%), see Supplementary Figure S5. The minimum and maximum absolute differences in DBH were -13.8 and 32.9 cm, respectively; the minimum and maximum relative differences were -42 and 101%, respectively (Supplementary Figure S5). The total tree basal area from the inventory was 36.8 cm² m⁻² while the total tree basal area obtained from TLS tree reconstruction was 36.2 cm² m⁻².

- Individual tree measurements from QSMs applied to the TLS point cloud could all be satisfactorily represented by the ED2.2
 allometric equations and a single PFT (Figure 2). R² of the allometric models for the individual aboveground woody biomass,
 height, crown area, and leaf biomass respectively reached 0.95, 0.83, 0.67, and 0.77. The normalised RMSD changed from
 18.3 to 16.9% (height), from 85.1 to 75.7% (crown area), from 146.1 to 95.0% (woody biomass), and from 151% to 83.5%
 (leaf biomass) when switched from ED2.2 default allometries for the mid-successional hardwood tree PFT to TLS-derived,
 site-specific ones (Table 3).
- 291 Over the DBH range in Wytham Woods, TLS-derived allometries led to systematically larger allocations to 292 above ground woody biomass (+73%) on average, up to +177% for the smallest tree) and leaf biomass (+75%) on 293 average), and smaller tree height (-1.9 m on average) as compared to ED2.2 defaults (Figure 2). Individual 294 crown areas derived from TLS measurements varied between 0.2 and 465.4 m², with a mean of 26 m². As 295 compared to the TLS-calibrated allometries, default model coefficients predicted larger crown areas for trees with 296 DBH < 64 cm (-22% on average), and smaller crown areas for trees with DBH \geq 64 cm (+17% on average), 297 see Figure 2. The latter category (DBH \geq 64 cm) comprised 30 trees (3.7% of the total) and contributed to 298 30.7% of the total basal area and 24.9% of the total leaf area.

299 Increasing the number of PFTs only slightly improved the goodness of fit of all four allometric models. The best species-to-300 PFT mapping according to the literature-informed minimization of the Watanabe information criterion was to classify Acer 301 pseudoplatanus as belonging to the late-successional hardwood PFT and the rest of the tree species as belonging to the mid-302 successional hardwood PFT (Table 1, Supplementary Figures S2 and S3). Using this classification, the normalised RMSD of 303 the allometric models decreased from 16.9 to 16.8% (height), 75.7 to 71.1% (crown area), 95.0 to 77.9% (aboveground 304 woody biomass), and 83.5 to 73.9% (leaf biomass). This mapping resulted in larger crown areas and larger carbon allocation 305 to woody and leaf tissues for small (DBH < 50 cm) trees of the mid-successional tree PFT and taller late-successional trees 306 across all DBHs (+1.16 m on average).

307 3.2 Ensemble runs

308 Regardless of the TRY constraints and the initial conditionsmodel configurations, the model ensembles could on average 309 reproduce both the amplitude and the seasonality of the gross ecosystem productivity, as observed by the eddy covariance 310 flux tower, with a maximum GPP in June and a leaf-off season with close-to-zero GPP in December-February (Figure 3). R² 311 of observed vs simulated monthly mean of mean of the monthly GPP was larger than 0.93 for all configurations (NBG, 312 Census, TLS) while the RMSE varied between 1.2 (NBG), 1.3 (TLS) and 1.9 (Census) umol $m^{-2} s^{-1}$, much lower than the 313 mean and standard deviation of the two years of observational data of GPP (5.5 and 4.7 μ mol m⁻² s⁻¹, respectively). Because 314 we only simulated fully deciduous tree PFTs, model ensembles underestimated GPP during winter; simulated ecosystem LAI 315 and hence ecosystem gross productivity dropped to almost zero in December-February (Supplementary Figure S6) while 316 measured ecosystem productivity was non-null during the same period (Figure 3), driven by evergreen understory plants 317 such as shrubs that were not included in our simulations.

318 The variability of the simulated GPP was critically influenced by the model configuration and the application of constraints 319 on SLA and V_{c max} (Figure 3). The standard deviation of the ensemble runs for the simulated GPP was not unexpectedly the 320 largest for the configuration with the least information on the ecosystem (the NBG configuration without TRY constraints), 321 and reached 6.33 µmol m⁻² s⁻¹ for June (Figure 3). More than 23% of the runs in that e NBG configuration without TRY constraints led to unvegetated conditions (LAI < $0.1 \text{ m}^2 \text{ m}^{-2}$, all year long, see Supplementary Figure S6) after 100 years of 322 323 simulations while about 5% of the runs simulated unrealistically dense tree covers (LAI > 10 m² m⁻² in summerwhen the tree-324 eovers reaches its maximum). Combined with the uncertainty of all other parameters, including the photosynthetic 325 onesparameters, the LAI variability explainsed the extreme variability of the simulated ecosystem's gross productivity. The 326 95% confidence interval of the simulated ecosystem GPP in June for the NBG configuration without TRY constraints (0 -327 19.8 μ mol m⁻² s⁻¹) was almost twice as large as the observed GPP at that moment (13.2 μ mol m⁻² s⁻¹).

328 Prescribing initial conditions reduced the variability of the simulated outputs: ensemble standard deviation of GPP in June 329 for the Census configuration without TRY constraints was 4.83 µmol m-2 s-1. However, for the ecosystem productivity 330 constraining SLA and Vc.max was even more critical: ensemble standard deviation of GPP in June for the Census 331 configuration with TRY constraints decreased to 1.99 µmol m-2 s-1 (see Figure 3 and also Figure 4 where the pie chart 332 radius is set proportional to the variance of the simulated ecosystem GPP). When both parameters were constrained and 333 realistic initial conditions were prescribed to the model (i.e. going from the NBG-without TRY constraints to the Census-334 with TRY constraints configuration), the variability of the simulated GPP experienced a three-fold decrease. Similarly, the 335 variability of LAI (supplementary Figure S6-7) and AGB (supplementary Figure S8) was drastically reduced, with a four-336 fold and a two-fold decrease respectively.

Given the similarities of the tree size distributions derived from the inventory and TLS (see results section 3.1), prescribing
 initial conditions had a similar impact on the variability of the outputs for the TLS and for the Census configurations.

- 339 Combined with the constraints on allometries, it led to a reduction of the ensemble standard deviation for GPP in June to 340 3.78 µmol m-2 s-1 for the TLS configuration without TRY constraints. As for the Census configuration, constraining SLA 341 and Vc.max with TRY data had a larger impact on the model uncertainty: ensemble standard deviation of GPP in June for 342 the TLS configuration with TRY constraints decreased to 1.54 umol m-2 s-1. Incrementally adding the TLS-related 343 information to the Census-with TRY constraints configuration had a positive, yet more limited effect on the reduction of the 344 model variability of GPP: ensemble standard deviation of GPP in June was reduced by 30% between the Census and TLS 345 configurations with TRY constraints. Constraining allometries with TLS had a more significant impact on LAI 346 (supplementary Figures S6-S7) and AGB (supplementary Figure S8), with a three-fold decrease of the ensemble standard 347 deviation from the Census-with TRY constraints to the TLS-with TRY constraints configurations.
- 348 All in all, Tthe predicted variability of the ecosystem LAI and GPP was on the contrary very lowest for the TLS 349 configuration with TRY constraints: 3.79 ± 0.50 m² m⁻² for the ensemble mean (\pm one standard deviation) of the ecosystem LAI (Supplementary Figure S6), $9.86 \pm 2.89 \mu mol m^{-2} s^{-1}$ for the ensemble mean (± one standard deviation) of the ecosystem 350 351 GPP (Figure 3), both during leaf-on conditions, which compared well with independent observations (Table 6). The 352 confidence interval of the simulated ecosystem GPP in June for the TLS configuration with TRY constraints was 353 significantly reduced (11.8 - 17.6 umol $m^{-2} s^{-1}$) and much closer to the confidence interval of the observations (11.5 - 14.6 354 μ mol m⁻² s⁻¹). In total, the variability of the simulated GPP experienced a four-fold decrease when parameters were 355 constrained, realistic initial conditions were prescribed, and TLS data were used to constrain the allometries (i.e. going from 356 the NBG-without TRY constraints to the TLS-with TRY constraints configuration).
- Prescribing realistic initial conditions reduced the variability of the simulated outputs (ensemble standard deviation of GPPin June for the Census configuration without TRY constraints was 4.83 μ mol m⁻² s⁻¹), just like imposing the TLS-constrained allometries (ensemble standard deviation of GPP in June for the TLS configuration without TRY constraints was 3.78 μ molm⁻² s⁻¹). However, for the ecosystem productivity constraining SLA and V_{e,max} was even more critical: ensemble standard deviation of GPP in June for the Census and TLS configurations with TRY constraints decreased to 1.99 and 1.54 μ mol m⁻²s⁻¹, respectively (Figure 3 and Figure 4 where the pie radius is proportional to the variance of ecosystem GPP).

363 3.3 Variance decomposition and sensitivity analyses

The variance of the ecosystem GPP was dominantly driven by the parameter uncertainty regardless of the configuration and the application of TRY constraints (Figure 4). Together, TRY-constrainable parameters, allometric coefficients, and the other ED2.2 parameters included in the sensitivity analysis, contributed on average to 63% of the total variance of GPP in June._-Constraining SLA and Vc,max with TRY datasets dramatically decreased the relative contribution of these two parameters to the overall variance: moving from uninformed priors to posteriors generated by the trait meta-analysis of PEcAn made the sum of their partial variances drop from a majority (57% on average for all three configurations) to a small contribution (7% on average for all three configurations), their share being mainly replaced by unconstrained parameters which increased from 6% to 50% on average across all configurations (Figure 4), especially the Quant. Eff., the Clumping
 and the Growth resp. parameters (Figure 5). The variance decomposition of the simulated ecosystem LAI and aboveground
 biomass led to very similar results, yet with a larger contribution of allometric parameters: allometric parameters contributed
 on average to 6 and 20% of the variance for LAI and AGB respectively, a larger contribution than theirs for the variance of
 GPP (3%), which illustrates the importance of TLS to constrain the ecosystem structure (Figure 5 and Supplementary
 Figures S7-S8).

377 On average, processes only accounted for 124% of the overall variance of GPP with a maximum (resp. minimum) for the 378 TLS configuration with TRY constraints with 20% (resp. for the NBG without TRY constraints with 5%). Constraining SLA-379 and V_{e max} with TRY datasets dramatically decreased the relative contribution of these two parameters to the overall variance: 380 moving from uninformed priors to posteriors generated by the trait meta-analysis of PEcAn made the sum of their partial 381 variances drop from a majority (57% on average for all three configurations) to a small contribution (7% on average for all 382 three configurations), their share being mainly replaced by unconstrained parameters which increased from 6% to 50% on 383 average across all configurations (Figure 4), especially the Quant, Eff., the Clumping and the Growth resp. parameters-384 (Figure 5). Process uncertainty was dominated by the type of crown model (5%) and the radiative transfer model (4%). Trait 385 plasticity only contributed marginally to the overall variance (< 1% on average).

386 The variance decomposition of the simulated ecosystem LAI and aboveground biomass led to very similar results, yet with a 387 larger contribution of allometric parameterses (. With an average contribution of 6 and 20% for LAI and AGB respectively (. 388 to be compared with anthe mean average contribution of of 3% for GPP), allometric parameters had) and hHence a stronger-389 impact of TLS constraints on the variance of thesethose output variables, which reinforced the impact of TLS constraints on the ecosystem structure (Figure 5 and Supplementary Figures S7-S8). Processes (especially the choice of the RTM) played a 390 391 stronger role for the available light in the understorey (on average 40% of the total variance), especially in runs with 392 prescribed initial conditions (on average 56% of the total variance, see Supplementary Figure S9). Due to compensatory 393 effects (Supplementary Figure S2), the number of simulated PFTs had a limited impact on all of the considered model 394 outputs: NPFT only contributed to 3% of the variance of ecosystem GPP, 2% of the variance of LAI and PAR, and 1% of the 395 variance of AGB.

396 3.4 Ecosystem <u>structure</u> and functions

397 Despite similar seasonal cycles of ecosystem productivity (Figure 3), ensemble means exhibited very contrasted ecosystem 398 structure (Figures 6-7). None of the unprescribed simulations (NBG configuration) could capture the size distribution 399 observed through the inventory (Figure 6). Small-size stem (especially DBH < 50 cm) densities were underestimated while 400 large tree (DBH > 100 cm) densities were overestimated in the vegetated simulations (LAI > 0.1 m² m⁻²) of the NBG 401 configuration with or without TRY constraints. Switching from closed canopy to finite crowns systematically increased the 402 density of small (DBH < 50 cm) trees, by 73% on average; just like constraining SLA and V_{c.max} with TRY data. While the 403 ecosystem LAI of the NBG configuration with closed canopies compared well with independent observations from the 404 literature $(3.83 \pm 1.94 \text{ versus the range of } 3.6 - 4.1 \text{ m}^2 \text{ m}^{-2} \text{ observed in Wytham Woods, Table 6})$, the vertical arrangement of 405 the leaves significantly differed from what was observed by TLS and imposed in the TLS configuration (Figure 7), as a 406 result of the differences in tree size distribution (Figure 6).

407 Despite lower total leaf areas, the infinitely wide crown configuration (closed canopies, Table 6) made the forest more 408 opaque to the incoming solar radiation than the finite crowns. Across all configurations, the PAR available in the understory 409 decreased by 15% throughout the year while the ecosystem LAI decreased by 18% when closed canopies were simulated 410 (Table 6). For near bare-ground configurations, the LAI of the potential vegetation simulated was 23% lower with infinite 411 crowns, and 16% less PAR reached the understorey.

412 As the soil received more radiation when finite crowns were simulated, it was warmer and as a result, heterotrophic (and 413 ecosystem respiration, see Table 6) increased (+ 25% on average) when switching from infinite to finite crowns. Forest 414 carbon stocks also diverged between configurations: driven by higher allocations to leaf and aboveground woody biomass 415 (Figure 2), aboveground carbon storage was larger (+74% on average) in TLS-derived runs than when default allometries 416 were applied (Table 6). Aboveground woody biomass from configurations starting from near bare-ground conditions was 417 systematically underestimated compared to the TLS estimates (11.4 kg_C m⁻² on average for the NBG configuration versus 418 24.5 kg_c m⁻² on average for the TLS configuration). However, the larger allocation to woody biomass induced by the use of 419 TLS-derived allometries mostly did not impact any other model outputs (Figure 5) as that carbon pool is inert and does not 420 influence a lot of processes downstream (e.g. more woody biomass does not translate into exacerbated light interception). 421 Leaf biomass allometry derived from TLS both reduced the simulated LAI and ecosystem GPP to more realistic values and 422 constrained its variability (Figures 3, Table 6, and Supplementary Figure S6).

423 None of the simulation/configurations could accurately represent all features of Wytham woods. The model simulations 424 starting from near bare-ground conditions failed to capture the vertical distribution of leaves (Figure 6) and the tree size 425 distribution (Figure 7); the model simulations prescribed with the inventory overestimated the ecosystem GPP (Table 6); and 426 the model simulations from the three configurations all overestimated the net ecosystem productivity (NEP), due to an 427 overestimation of GPP (Census) and/or an underestimation of the ecosystem respiration (Census, NBG, and TLS), see Table 428 6. Model simulations underestimated R_{eco} on average by -17% leading to unrealistic NEP predictions, which illustrates the 429 need for constraining or optimising autotrophic and heterotrophic respiration parameters along with the photosynthetic and 430 allometric parameters to align those with observational data.

431 4 Discussion

432 4.1 The relative weight of the different sources of uncertainty

433 The different model configurations tested in this study led to contrasting predictions of vegetation states. Depending on the 434 chosen model outputs, the relative weights of the sources of uncertainty considerably varied. Near bare-ground simulations 435 generated potential vegetations that significantly differed in their demography from observations (Figure 4) while 436 prescribing initial tree size distribution was not a guarantee for accurately reproducing observed land fluxes (Figure 3, Table 437 6). The finite crown area representation also had a substantial impact on the model outputs. In particular, limiting the crown 438 radius to finite values promoted smaller plants in the understorey (Figure 6), increased the simulated LAI (Table 6) and 439 profoundly modified the vertical distribution of light in the canopy (Figure 8 and Table 6). Carbon pools also considerably 440 diverged between model configurations, especially when TLS-derived allometries were taken into account (Table 6).

However, in general, it was the parameter uncertainty that dominated the overall model uncertainty (Figure 3, Supplementary Figure S7 and S8), just like it was previously observed for ED2.2 simulations of temperate forests (Shiklomanov et al. 2020). The parameters that dominated the variance depended on the use of TRY and/or TLS constraints. When observations were available, uncertainty was transferred to other unconstrained parameters while the overall variance was reduced, like in previous similar studies (Meunier et al. 2021), which supports the process of progressively integrating observations of most sensitive parameters until the model variance is reduced to satisfactorily levels in an efficient data-model fusion loop (Dietze et al. 2014).

Although parameter uncertainty was larger in magnitude than process uncertainty, crown size representation and the choice of RTMs appear to drive a significant part of the model process uncertainty and should be paid more attention to in future analyses. Especially, because the implementation and the sensitivity of the radiative transfer processes are currently overlooked in ED2.2 like other vegetation models (Fisher et al. 2018; Viskari et al. 2019).

452 4.2 The added value of TLS for vegetation modelling

453 The quantitative information that remote sensing generates at unprecedented spatial and temporal scales can serve the 454 purpose to reduce uncertainties in TBM projections. It has already been shown that airborne laser scanning (ALS) combined 455 with an individual-based forest model could offer new insights into the contribution of plant size to ecosystem functioning 456 (Fischer et al. 2019). Similarly, ALS and synthetic-aperture Radar have successfully been applied to prescribe the initial 457 structure and composition of tropical forests (Antonarakis et al. 2011; Antonarakis et al. Moorcroft 2014; Longo et al. 2020), 458 and LiDAR data have been coupled to allometric models to estimate carbon stocks and fluxes at large scale (Hurtt et al. 2019; 459 Thomas et al. 2008). Yet, our study is the first attempt to inform a TBM withfuse TLS data-and TBMs. As compared to ALS, 460 TLS offers a few significant advantages, as well as some drawbacks, that are important to remember. Airborne techniques 461 allow for wall-to-wall coverage characterising 3D forest structure at the regional scale, whereas TLS offers far more detailed

462 information but only at the local (up to a few ha) scale. However, TLS is <u>can</u> capable of estimating the volume of individual 463 trees directly, instead of relying on allometries that require calibration and thus field measurements. In addition, it can 464 accurately capture the entire size distribution (DBH and height) of the sample plot while smaller trees can easily be missed 465 with airborne surveys (Wang et al. 2016) leading to incorrect demography, especially in dense forests.

466 Because TLS data are complementary to the datasets that are frequently used for model calibration (e.g. eddy covariance 467 data), they can contribute in a collective effort towards realistic representations of ecosystems in TBMs. TLS has the 468 potential to fill important parameter and process gaps and in doing so, to help reduce the uncertainties in vegetation model 469 simulations. The steep increase in the amount of available forest TLS data over the past decade (Calders et al. 2020) makes 470 its coupling with TBMs even more timely. As demonstrated in this study, TLS observation can ensure a more adequate 471 model structure, constrain model allometric parameters and prescribe representative initial conditions. Yet, only a 472 combination of constraints on both allometries (using TLS data) and photosynthetic parameters (thanks to TRY data) could 473 satisfactorily reduce the model uncertainties to its lowest level, which supports the integration of multiple data sources into 474 TBMs for more realistic simulations (Peylin et al. 2016). Such a combination fusion of a TBM and multiple data streams 475 allowed us to accurately simulate both ecosystem productivity and ecosystem community composition with physically 476 realistic parameters, which was previously highlighted as a challenge for dynamic vegetation models (Shiklomanov et al. 477 2020; Fisher et al. 2010).

478 In the future, TLS could inform vegetation models even more. The TLS community is indeed actively working on the

479 derivation of additional tree- or stand-scale parameters from lidar raw data and 3D point clouds. Those parameters include 480

leaf angle distributions (Boni Vicari et al. 2019), clumping (Zhao et al. 2012), and reflectance (Calders et al. 2017), which

481 have been shown to significantly contribute to the overall model uncertainty (Meunier et al. 2021; Shiklomanov et al. 2020;

482 Viskari et al. 2019). Yet, theoretical, technological, and technical challenges specific to each parameter still need to be raised

483 before one can constrain these sensitive traits with TLS in a study similar to this one.

484 4.3 Model equifinality

485 Some runs from all three configurations (prescribed or not with initial size distributions) could reproduce the seasonal cycle 486 of GPP observed by the flux tower (Figure 3). However, those 'optimal' simulations were very different from the forest 487 structure point of view (Table 6, Figures 6-7). This situation illustrates the low identifiability of numerous TBM parameters 488 and the need for multiple simultaneous constraints and observations. While aboveground carbon storage is critical to 489 estimate forest sink strength and the overall carbon storage capacity of the ecosystem (Keeling and Phillips 2007), it has a 490 limited impact on simulated land fluxes (GPP in particular, see Figure 5) that are often used to calibrate TBMs. The 491 parameters controlling land fluxes, namely those controlling ecosystem LAI (Williams and Torn 2015; Wei et al. 2013) and 492 those related to photosynthesis (Figure 5), are also confounded, echoing observed trade-offs of the Leaf Economic Spectrum 493 (Wright et al. 2004: Peaucelle et al. 2019). TLS has the potential to discriminate equifinal model simulations with similar

494 land fluxes but contrasting structure. On-site trait measurements (Figure 3) could further help avoid those risks of495 equifinality (Babst et al. 2020; Peaucelle et al. 2019).

496 4.4 Study limitations

497 Our findings come with several important limitations. First, the eddy covariance flux data (2007-2009) preceded the 498 observation of the forest structure (TLS and field inventory occurred over the 2015-2016 period) by almost a decade. The 499 forest composition and demography might have changed in the meanwhile, which reduces the confidence of the validation 500 with eddy covariance data (Figure 3). This is even more true as one realises that the validation dataset is rather limited in size 501 and information content (very low year-to-year variability in observed fluxestwo very similar seasonal cycles of GPP). Yet, 502 in this study we were more interested in the variance decomposition for different model configurations (Figures 3-4) than the 503 actual goodness of fit of every single configuration. In addition, in the absence of locally observed meteorological drivers, 504 we had to force the model simulations with regional datasets that cannot serve the purpose of capturing the day-to-day 505 variability or the diel cycle, which forced us to only compare the modelled and observed seasonal GPP cycle. Furthermore, 506 GPP is not directly observed but rather a derived (modelled) quantity as opposed to the net ecosystem exchange of carbon 507 and the latent heat flux of water that are directly measured. We could not access water flux raw data nor were they reported 508 in publications that we knew of. GPP uncertainties were also not quantified in the original publication of Thomas et al. 509 (2011). While NEP values were reported, validating the model simulations with those values would have biassed our 510 analyses as we could not constrain respiration parameters with data. Mismatches between different data sources and/or the 511 low availability of good-quality data are recurrent issues in vegetation modelling exercises. Despite multiple initiatives to 512 standardise high quality data such as Fluxnet (Baldocchi et al. 2001), we emphasise here the need for concomitant 513 observations in experimental and observational plots.

514 Second, the comparison between the potential vegetations as simulated by ED2.2 and the field inventory data are also 515 imperfect as Wytham Woods is a managed forest that has been frequently coppiced and pollarded. Disturbance history 516 experienced by the ecosystem is mostly unknown, preventing us from reproducing the current forest demography by the 517 model.

518 Third, the trait meta-analysis was run with random effects turned off, which can generate too narrow parameter posterior
 519 distributions (Raczka et al. 2018), and hence underestimate the contribution of the TRY-constrained parameters (see e.g.
 520 Figure 4). A similar analysis including random effects should be repeated to evaluate such an underestimation.

521 Finally, the ecosystem growth form complexity was neglected in this study. We only simulated tree PFTs while shrubs and 522 grass species also coexist in Wytham Woods. Integrating this ecological complexity would not have brought additional 523 information or robustness regarding the objectives of our study on the variance decomposition while increasing the

- 524 dimensionality and complexity of the problem. Future research should investigate whether the main findings highlighted in
- 525 this study hold with other PFTs, across other sites and biomes, or even in other vegetation models (Dokoohaki et al. 2021).

527 **5** Conclusion

528 Vegetation models are important tools to predict the fate of ecosystems in a changing climate but are often used as black-box 529 tools due to their complexity. They have been designed to realistically represent the ecosystem that they simulate, but often 530 fail to do so primarily because of considerable parameter uncertainties as well as process and initialisation errors. Even for 531 the state-of-the-art process-based terrestrial biosphere models, not all parameters can be constrained with data: some cannot 532 be observed in the field, require calibration, or the appropriate observational trait data may be missing. In addition, model 533 initialisation and the choice of model structure necessarily lead to additional uncertainties. We demonstrate in this study that 534 TLS has the potential to provide initial condition estimates and to constrain some critical vegetation model parameters 535 (allometries) and processes (crown representation). Combined with trait-based constraints on a few key parameters, TLS was 536 able to define a model configuration that could reproduce both the ecosystem productivity and the plant community 537 composition of the simulated site with physically realistic parameters, as well as considerably reduce model uncertainties.

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915 Tables

917 available in the TRY database for each of the five dominant species in Wytham woods, and their local prevalence (in terms of 918 individual density and basal area). Missing traits were unavailable in TRY. The table also summarises the abundance of those five

919 dominant species in the 1.4 ha plot in terms of absolute and relative density and basal area, as well as the PFT mapping when more

920 than one PFT were simulated (N_{PFT} > 1). The community weighted means (CWM) and standard deviations (CWSD) were obtained

921 using the basal areas as weights.

Ap = Acer pseudoplatanus, Ca = Corylus avellana, Cm = Crataegus monogyna, Fe = Fraxinus excelsior, and Qr = Quercus robur.
 The colours of the different species in the first row of the Table are consistent with Figures 1 and 2.

Trait	Ар	Са	Cm	Fe	Qr	Others	CWM (± CWSD)
SLA (m² kg _c -¹)	-	34.7 (± 36.1)	62.8 (± 65.5)	3 0	22.9 (± 23.9)	-	25.1 (± 1.5)
V _{c,max} (µmol m ⁻² s ⁻¹)	31.9 (± 16.1)	-	-	39.7 (± 18.0)	31.1 (± 18.8)	-	32.6 (± 0.9)
PFT (if N _{PFT} > 1)	LH ⁽¹⁾	MH ⁽¹⁾	мн	МН	МН	мн	
State variable				teril.	2. 1	1.	Total
Density (-)	532	67	24	84	35	73	815
Relative density (%)	65.3	8.2	2.9	10.3	4.3	9.0	100
Basal area (m²)	31.59	0.48	0.24	5.96	11.87	0.57	50.71
Relative basal area (%)	62.3	0.9	0.5	11.8	23.4	1.1	100

924

⁽¹⁾MH = Mid successional Hardwood trees, LH = Late successional Hardwood trees

Table 2: List of varying processes included in the model ensembles in order to evaluate the model structural uncertainty as well as
 their different possible configurations. Adapted from Shiklomanov et al. 2020.

Process	Description
Crown model	Choice of the crown representation in the canopy radiation model and in the turbulence scheme
Closed	Crowns are evenly spread throughout the patch area and cohorts are stacked on the top of each other
Finite	Cohorts have a finite radius and are stacked on the top of each other (Dietze et al. 2008)
Radiative transfer model (RTM)	Choice of the canopy radiation model
Two-stream	Two-stream approximation (Oleson et al. 2013; Sellers 1985)
Multi-scatter	Multiple-scatter approximation (Zhao and Qualls 2005)
Trait plasticity	Choice of including plant trait variation with the local environment
False	SLA and $V_{c,max}$ are constant
True	SLA and $V_{c,max}$ respectively increases and decreases with shading
Plant functional diversity (N _{PFT})	Number of PFTs included in the simulation
1	All plant species are classified as mid-successional temperate deciduous trees
2	Plant species are mapped into two PFTs according to Table 1 classification

929 Table 3: List of allometries modified in this study, ED2.2 default and TLS-derived allometric coefficients (for one or multiple 930 simulated PFTs). The corresponding curves are plotted in Figure 2.

					TLS		
Allometry	Equation ⁽¹⁾	$\begin{array}{c c c c c c c c c c c c c c c c c c c $	2 default	N _{PFT} = 1	N _{PFT} = 2		
			MH ⁽²⁾	LH ⁽²⁾	мн	мн	LH
		h _{ref}	1.3	1.3	-3.2	-3.2	-2.8
Height, <i>h</i> (m)	$h = h_{ref} + h_1 \cdot [1 - exp(DBH \cdot h_2)]$	h_1	25.2	23.4	26.2	25.4	26.4
		h_2	-0.05	-0.054	-0.074	Npr MH -3.2 25.4 -0.074 0.67 2.13 1.4 0.95 0.095	-0.07
		B_{d1}	0.16	0.24	0.37	0.67	0.23
Aboveground woody biomass, B _d (kg)	$B_d = B_{d1} \cdot DBH^{B_{d2}}$	B_{d2}	2.46	2.25	2.29	2.13	2.42
0		$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	2.49	0.6	1.4	0.3	
Crown area, CA (m ²) $CA = CA_1 \cdot DBH^{CA_2}$	$CA = CA_1 \cdot DBH^{CA_2}$	CA_2	0.81	0.81	1.15	0.95	1.33
		B_{l1}	25.2 23.4 -0.05 -0.054 0.16 0.24 2.46 2.25 2.49 2.49 0.81 0.81 0.048 0.017	0.065	0.095	0.015	
Leaf biomass, <i>B</i> _i (kg)	$B_l = B_{l1} \cdot DBH^{B_{l2}}$	B_{l2}	1.46	1.73	1.48	1.22	1.69

⁽¹⁾DBH = Diameter at Breast Height (cm)

⁽²⁾MH = Mid successional Hardwood trees, LH = Late successional Hardwood trees

932 Table 4: Description of the ED2.2 parameters varied in this stuy, their unit, and the definition of their prior used to evaluate the 933 model parameter uncertainty. "Source code name" is the name of the parameter as it appears in the ED2.2 source code. When 934 trait plasticity is enabled, both SLA and V_{c,max} may change over time and for different cohorts of the same PFT.

			Prior			
Parameter name	Description	Unit	Function ⁽¹⁾	a ⁽²⁾	b ⁽²⁾	Source code name
Water cond.	Soil-plant hydraulic conductance	m² (kg _{c,root})-1 yr -1	Inorm	-10.8	3.5	water_conductance
Growth resp.	Fraction of assimilation lost to growth respiration	Unitess (0-1)	beta	4.06	7.2	growth_resp_factor
Mort. C bal.	C balance ratio at which mortality rapidly increases	Unitless	gamma	1.47	0.058	mort2
V _{c,max}	Maximum rate of CO ₂ carboxylation at 15°C (baseline)	µmol _c m² s¹	weibull	1.7	80	Vm0
Leaf resp.	Leaf dark respiration at 15°C	µmol _c m ⁻² s ⁻¹	gamma	1.5	0.4	Rd0
Root:leaf	Ratio of fine root to leaf biomass	Unitless	Inorm	0.21	0.6	q
SLA	Specific leaf area (baseline)	m² (kg _{c,leat}) ⁻¹	gamma	5.13	0.23	SLA
Clumping	Canopy clumping factor	Unitless (0-1)	beta	3	1.5	clumping_factor
Quant. eff.	Fraction of absorbed light used for CO_2 fixation	mol CO ₂ (mol photon) ⁻¹	weibull	3.32	0.08	quantum_efficiency
Refl. (VIS)	Leaf reflectance in the visible range (400-700 nm)	Unitless (0-1)	beta	10.1	157	leaf_reflect_vis
Refl. (NIR)	Leaf reflectance in the NIR ⁽³⁾ range (700-2500 nm)	Unitless (0-1)	beta	35	56	leaf_reflect_nir
Stomatal slope	Slope between leaf assimilation and stomatal conductance (Leuning)	Unitless	Inorm	2.3	1	stomatal_slope
Min. height	Minimum height for plant reproduction	m	gamma	1.5	0.2	repro_min_h

⁽¹⁾Inorm = log-normal distribution

(3)NIR = near-infrared

⁽²⁾The values a and b define the parameters of the prior distributions (LeBauer et al., 2013).

936 Table 5: Summary of the model configurations used in this study and the underlying model settings.

			Configuration name	
		NBG	Census	TLS
	Initial conditions	Near-bare ground	Inventory	TLS
	Allometric parameters	Unconstrained	Unconstrained	TLS-constrained
	Run length (years)	100	5	5
	Crown model	Closed or finite	Closed or finite	Finite
Settings	RTM		Two-stream or multi-scatter	
	Trait plasticity		True or false	
	N _{PFT}		1 or 2	
	Ensemble size		500	

Table 6: Summary of most important states and fluxes-variables in all three model configurations and how they compare with

939 observational datasets, including flux tower data of ecosystem respiration and net ecosystem productivity. Those numbers take 940 into account the full five years of simulation for the prescribed model configurations (Census and TLS), and the last five years of 941 simulation for the near bare-ground conditions (NBG), and the two years of eddy covariance observational data. For the 942 observations of LAI in the leaf-on season, we provide a range of variation.

LAI = Leaf Area Index, AGB = Aboveground Biomass, GPP = Gross Primary Production, NEP = Net Ecosystem Productivity,
 PAR = Photosynthetically Active Radiation

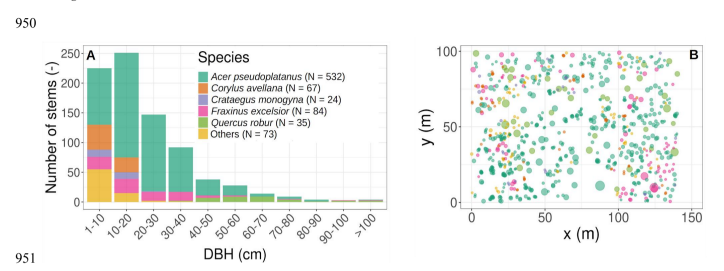
		Configuration					
		N	BG	Cer	nsus	TLS	Observations
	Units	Closed canopies	Finite crowns	Closed canopies	Finite crowns	Finite crowns	
LAI ⁽¹⁾	m² m ⁻²	3.83 ± 1.94	4.72 ± 3.67	4.71 ± 1.28	5.75 ± 2.74	3.79 ± 0.50	3.6 - 4.1 ⁽³⁾
AGB	kg _c m ⁻²	11.9 ± 7.4	10.8 ± 6.8	16.4±5.3	17.1 ± 4.7	24.5 ± 2.5	π.
GPP ⁽¹⁾	µmol m ⁻² s ⁻¹	9.55 ± 4.34	9.81 ± 4.70	10.90 ± 2.91	11.80 ± 2.95	9.86 ± 2.89	9.8 ± 3.4 ⁽²⁾
Ecosystem respiration	µmol m ⁻² s ⁻¹	4.51 ± 2.04	4.64 ± 2.24	4.56 ± 1.16	4.78 ± 1.15	3.98 ± 1.17	5.3 ± 2.2 ⁽²⁾
NEP	µmol m ⁻² s ⁻¹	1.53 ± 0.86	1.63 ± 0.89	2.32 ± 1.05	2.68 ± 0.42	2.26 ± 1.02	0.3 ± 2.9 ⁽²⁾
PAR reaching the ground ⁽¹⁾	µmol m ⁻² s ⁻¹	78.6 ± 93.2	90.9 ± 95.4	44.8 ± 34.7	58.2 ± 35.3	98.2 ± 36.0	π.

⁽¹⁾Leaf-on only (May to October)

⁽²⁾Reference: Thomas et al. (2011) and Fenn et al. (2015).

⁽³⁾Reference: Roberts et al. (1999)

		Configuration					2
	Units	NBG		Census		TLS	Observations
		Closed canopies	Finite crowns	Closed canopies	Finite crowns	Finite crowns	
AGB	kg _c m ⁻²	11.9 ± 7.4	10.8 ± 6.8	16.4 ± 5.3	17.1 ± 4.7	24.5 ± 2.5	-
Leaf-on only perio	od (May to Octo	ober)					1
LAI	m² m-²	3.83 ± 1.94	4.72 ± 3.67	4.71 ± 1.28	5.75 ± 2.74	3.79 ± 0.50	3.6 - 4.1 ⁽²⁾
PAR reaching the ground	µmol m ⁻² s ⁻¹	78.6 ± 93.2	90.9 ± 95.4	44.8 ± 34.7	58.2 ± 35.3	98.2 ± 36.0	-
GPP	µmol m ⁻² s ⁻¹	9.55 ± 4.34	9.81 ± 4.70	10.94 ± 2.91	11.83 ± 2.95	9.86 ± 2.89	9.8 ± 3.4 ⁽¹⁾
Ecosystem respiration	µmol m ⁻² s ⁻¹	6.92 ± 3.13	7.03 ± 3.43	7.03 ± 1.82	7.32 ± 1.80	6.07 ± 1.81	7.2 ± 1.3 ⁽¹⁾
NEP	µmol m ⁻² s ⁻¹	2.63 ± 1.46	2.78 ± 1.49	3.91 ± 1.74	4.51 ± 1.92	3.79 ± 1.67	2.6 ± 2.5 ⁽¹⁾
All year round				1			I
GPP	µmol m ⁻² s ⁻¹	6.04 ± 2.77	6.26 ± 3.02	6.88 ± 1.84	7.46 ± 1.87	6.24 ± 1.85	5.5 ± 4.7 ⁽¹⁾
Ecosystem respiration	μmol m ⁻² s ⁻¹	4.51 ± 2.04	4.64 ± 2.24	4.56 ± 1.16	4.78 ± 1.15	3.98 ± 1.17	5.3 ± 2.1 ⁽¹⁾
NEP	µmol m ⁻² s ⁻¹	1.53 ± 0.86	1.63 ± 0.89	2.32 ± 1.05	2.68 ± 0.42	2.26 ± 1.02	0.3 ± 2.9 ⁽¹⁾

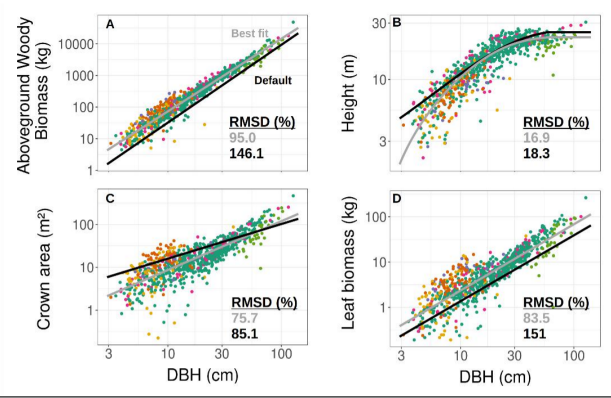


952 Figure 1: Initial conditions in terms of tree size distribution and species composition (A), horizontal position, basal area (the size of the circles in panel B is proportional to the individual basal area), and species composition (B). The species colour legend applies to

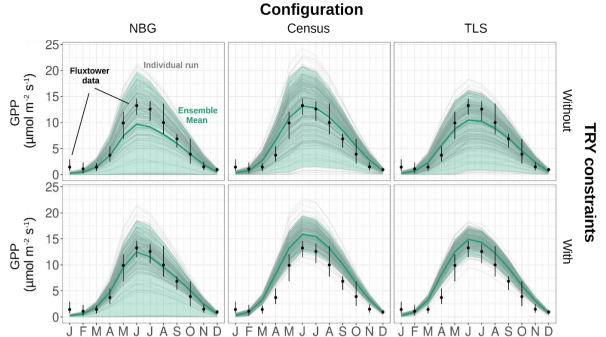
953 954 both panels and is kept the same for Figure 2 and Table 1. In the simulations, all trees were classified into either a single or 955 multiple plant functional types according to the species-PFT of Table 1.

949

Figures

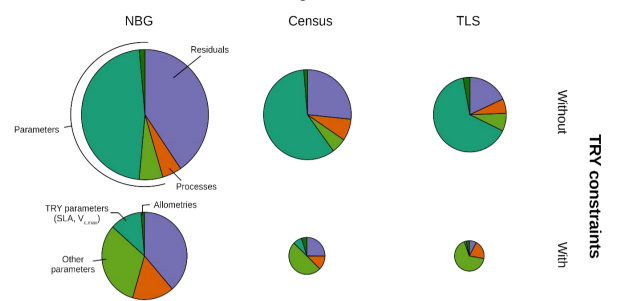


956 957 Figure 2: TLS-derived (grey, considering all tree species belonging to a single PFT) and model default (black, mid successional 958 hardwood trees in ED2) allometries for the aboveground woody biomass (A), tree height (B), crown area (C), and leaf biomass (D). 959 960 The data to which the TLS allometries were fitted (coloured points corresponding to the tree species detailed in Figure 1) were obtained using TLS. Coefficients used to plot the best fit and default allometries can be found in Table 3.

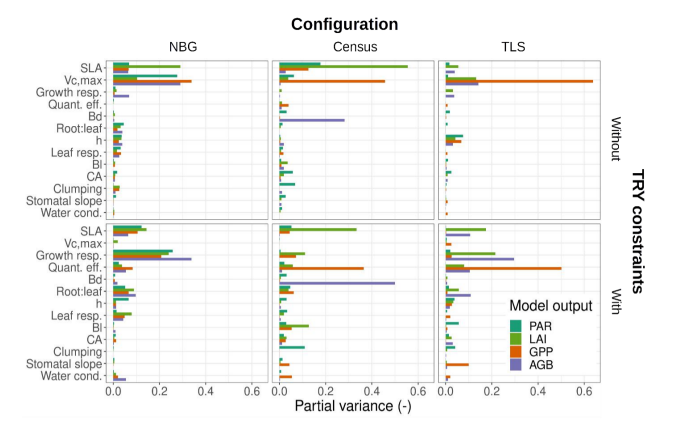


961 962 Figure 3: Seasonal cycle of the ecosystem GPP, as observed by eddy-covariance data (black dots) or as simulated by ED2.2 for 963 multiple model configurations (columns) and with or without TRY constraints on SLA and V_{c,max} (rows). The green thick lines are 964 the ensemble means while the shaded envelopes encompass 95% of the ensemble members. The individual ensemble members are 965 also plotted as thin grey lines. The vertical error bars for the flux tower data represent the 95% confidence interval of the monthly 966 GPP. The settings of the model configurations are detailed in Table 5.

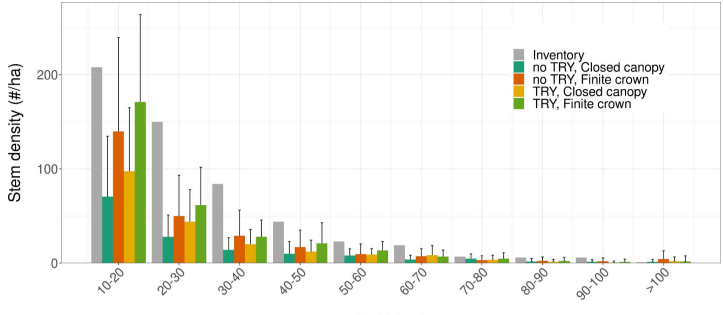
Configuration



967 968 Figure 4: Decomposition of the simulated GPP variance into process (orange), parameter (green), and residual (mauve) 969 uncertainty for multiple model configurations (columns) and with or without TRY constraints on SLA and V_{c,max} (rows). The 970 parameter uncertainty was further decomposed into the contribution of the allometric, TRY-constrainable (SLA and V_{c.max}), and 971 other parameters (shades of green). The radii of the pie charts are proportional to the total variance of the ecosystem GPP in each 972 configuration for the month of June (maximum GPP). The settings of the model configurations are detailed in Table 5.



975Figure 5: Contribution of individual or allometric parameters (Bd, Bl, CA and height include all parameters for the respective
allometries, see Table 2) to the predicted uncertainty in ED2.2 of multiple state variables (PAR = photosynthetically active
radiation reaching the ground, LAI = leaf-on ecosystem leaf area index, AGB = final ecosystem aboveground biomass, GPP = leaf-
on ecosystem gross primary production) for multiple model configuration (columns) and with or without TRY constraints on SLA
and $V_{c,max}$ (rows). Only those parameters that contributed at least once to 5% or more of the total variance were included in the
panels. Parameter description and distributions are given in Table 4. The settings of the model configurations are detailed in Table
9819815.



982 983 984

DBH (cm)

Figure 6: Tree size distribution for multiple model configurations starting from near bare-ground conditions after 100 years of simulations (coloured bars), and how they compare to the field inventory (grey). The histograms and the vertical error bars 985 986 represent the means ± one standard deviation of the ensemble member runs. Only runs that generated vegetation were kept for plotting this figure.

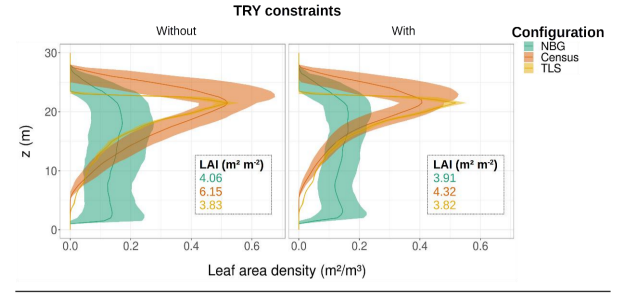


Figure 7: Ecosystem average of the leaf area density vertical distribution for the month of June for different model configurations (colourd lines and envelopes) without (left) and with (right) TRY constraints on SLA and $V_{C,max}$. The envelopes encompass the mean \pm one standard deviation of the 500 ensemble member runs. Only runs that generated vegetation were kept for plotting the NBG envelopes. The settings of the model configurations are detailed in Table 5.