

# 1 Simulation of tree ring-widths with a model for primary production, carbon allocation 2 and growth

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## 11 Abstract

12 We present a simple, generic model of annual tree growth, called ‘T’. This model accepts input from a first-  
13 principles light-use efficiency model (the P model). The P model provides values for Gross Primary  
14 Production (GPP) per unit of absorbed photosynthetically active radiation (PAR). Absorbed PAR is estimated  
15 from the current leaf area. GPP is allocated to foliage, transport-tissue, and fine root production and  
16 respiration, in such a way as to satisfy well-understood dimensional and functional relationships. Our  
17 approach thereby integrates two modelling approaches separately developed in the global carbon-cycle and  
18 forest-science literature. The T model can represent both ontogenetic effects (impact of ageing) and the  
19 effects of environmental variations and trends (climate and CO<sub>2</sub>) on growth. Driven by local climate records,  
20 the model was applied to simulate ring widths during 1958~2006 for multiple trees of *Pinus koraiensis* from  
21 the Changbai Mountain, northeastern China. Each tree was initialised at its actual diameter at the time when  
22 local climate records started. The model produces realistic simulations of the interannual variability in ring  
23 width for different age cohorts (young, mature, old). Both the simulations and observations show a significant  
24 positive response of tree-ring width to growing-season total photosynthetically active radiation (PAR<sub>0</sub>) and  
25 the ratio of actual to potential evapotranspiration ( $\alpha$ ), and a significant negative response to mean annual  
26 temperature (MAT). The slopes of the simulated and observed relationships with PAR<sub>0</sub> and  $\alpha$  are similar; the  
27 negative response to MAT is underestimated by the model. Comparison of simulations with fixed and  
28 changing atmospheric CO<sub>2</sub> concentration shows that CO<sub>2</sub> fertilization over the past 50 years is too small to be  
29 distinguished in the ring-width data given ontogenetic trends and interannual variability in climate.

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31

## 32 1 Introduction

33 Forests cover about 30% of the land surface (Bonan, 2008) and are estimated to contain  $861 \pm 66$  PgC (Pan et  
34 al., 2011). Inventory-based estimates show that forests have been a persistent carbon sink in recent decades,  
35 with a gross uptake of  $4.0 \pm 0.5$  PgCyear<sup>-1</sup> and a net uptake of  $1.1 \pm 0.8$  PgCyear<sup>-1</sup> between 1990 and 2007 (Pan  
36 et al., 2011). This is a significant amount in comparison with the amounts of carbon released from fossil fuel  
37 burning, cement production and deforestation ( $9.5 \pm 0.8$  PgCyear<sup>-1</sup> in 2011: Ciais et al., 2013) and thus, forest  
38 growth has a substantial effect on atmospheric CO<sub>2</sub> concentration and climate (Shevliakova et al., 2013).  
39 However, there is considerable geographic variability in the trends in the carbon sink as well as the factors  
40 controlling regional trends, and uncertainty about how forest growth and carbon sequestration will be affected  
41 by climate change, and climate-driven changes in wildfire (Ciais et al., 2013; Moritz et al., 2013). The  
42 changing importance of disturbance, and its influence on forest age, is likely to have a significant impact on  
43 the ability of forests to act as carbon sinks. It is generally assumed that stand-level productivity stabilizes or  
44 declines with age (Ryan et al., 1997; Caspersen et al., 2011). However, recent analyses have shown that mass  
45 growth rate (and hence carbon accumulation) by individual trees increases continuously with tree size  
46 (Stephenson et al., 2014), pointing to a need to understand the relationship between individual and stand  
47 growth rates. Predictions of future changes in the terrestrial carbon cycle (e.g. Friedlingstein and Prentice,  
48 2010) rely on ecosystem models that explicitly represent leaf-level processes such as photosynthesis, but in  
49 most cases do not incorporate the response of individual trees. In models that do consider individual tree  
50 growth (e.g. ED: Moorcroft et al 2001, Medvigy et al. 2012; LPJ-GUESS: Smith et al., 2001; Claesson and  
51 Nycander, 2013), little attention has been paid to evaluating the realism of simulated radial growth.  
52 Incorporating the response of individual trees to climate and environmental change within such modelling  
53 frameworks should help to provide more realistic estimates of the role of forests in the global carbon cycle.

54 Climate factors, such as temperature and moisture availability during the growing season, are important  
55 drivers of tree growth (Harrison et al., 2010). This forms the basis for reconstructing historical climate  
56 changes from tree-ring records of annual growth (Fritts, 2012). However, photosynthetically active radiation  
57 (PAR) is the principal driver of photosynthesis. Models for primary production that use temperature, not PAR,  
58 implicitly rely on the far-from-perfect correlation between temperature and PAR (Wang et al., 2014). PAR  
59 can change independently from temperature (through changes in cloudiness affecting PAR or atmospheric  
60 circulation changes affecting temperature) and this may help to explain why statistical relationships between  
61 tree growth and temperature at some high latitude and high elevation sites appear to breakdown in recent  
62 decades (D'Arrigo et al., 2008). CO<sub>2</sub> concentration also has an impact on tree growth, although its magnitude  
63 is still controversial: trends in tree growth have been attributed to increasing atmospheric CO<sub>2</sub> concentration  
64 in some studies (Wullschleger et al., 2002; Körner, 2006; Huang et al., 2007; Koutavas, 2013) and not others

65 (Miller, 1986; Luo et al., 2004; Reich et al., 2006). To resolve these apparent conflicts, and to understand tree  
66 growth processes better, it is necessary to analyse the response of tree growth to multiple factors acting  
67 simultaneously, including solar radiation, climate, CO<sub>2</sub>, and ontogenetic stage.

68 Modelling is needed for this purpose. Empirical models of annual tree growth and climate variables  
69 (temperature and precipitation) have been used to simulate tree radial growth (Fritts, 2012). Process-based  
70 bioclimatic models might be preferable, however, because this allows other factors to be taken into account  
71 (e.g. the direct impact of CO<sub>2</sub> concentration on photosynthesis) and for non-stationarity in the response to  
72 specific climate variables. Vaganov et al. (2006) and Rathgeber et al. (2005) have used bioclimatic variables  
73 (temperature, and soil moisture availability) chosen to reflect physiological processes to simulate radial tree  
74 growth. The MAIDEN model (Misson, 2004; Misson et al., 2004; see also MAIDENiso: Danis et al., 2012)  
75 models the phenological and meteorological controls on NPP and explicitly allocates carbon to different  
76 carbon pools (including the stem) on a daily basis using phenological stage-dependent rules. Nevertheless,  
77 MAIDEN still requires tuning of several parameters.

78 Simple equations representing functional and geometric relationships can describe carbon allocation by trees  
79 and make it possible to model individual tree growth (Yokozawa and Hara, 1995; Givnish, 1988; Falster et al.,  
80 2011; King, 2011). Such models are built on measurable relationships, such as that between stem diameter  
81 and height (Thomas, 1996; Ishii et al., 2000; Falster and Westoby, 2005), and crown area and diameter or  
82 height (Duursma et al., 2010) that arise because of functional constraints on growth. The pipe model  
83 represents the relationship between sapwood area and leaf area (Shinozaki et al., 1964; Yokozawa and Hara,  
84 1995; Mäkelä et al., 2000). The ratio of fine root mass to foliage area provides the linkage between above and  
85 below ground tissues (Falster et al., 2011). These functional relationships are expected to be stable through  
86 ontogeny, which implies that the fraction of new carbon allocated to different compartments is variable  
87 (Lloyd, 1999). In this paper, we combine the two modelling approaches previously developed in the global  
88 carbon-cycle (ecophysiology) and forest-science (geometric and carbon allocation) literature to simulate  
89 individual tree growth.

90

## 91 2 Methods

### 92 2.1 Model structure and derivation

93 We use a light-use efficiency model (the P model: Wang et al., 2014), driven by growing-season PAR, climate  
94 and ambient CO<sub>2</sub> concentration inputs, to simulate gross primary production (GPP). The simulated GPP is  
95 used as input to a species-based carbon allocation and functional geometric tree growth model (the T model)  
96 to simulate individual tree growth (Fig. 1).

97 2.1.1 The P model

98 The P model is a simple but powerful light-use efficiency and photosynthesis model, which simulates GPP per  
99 unit of absorbed PAR from latitude, elevation, temperature, precipitation and fractional cloud cover (Wang et  
100 al., 2014). The climate observations used here are monthly temperature, precipitation and fractional cloud  
101 cover, which are interpolated to a daily time step for subsequent calculations of the variables that determine  
102 annual GPP.

103 Potential annual *GPP* is the product of the PAR incident on vegetation canopies ( $PAR_0$ ), with the maximum  
104 quantum efficiency of photosynthesis ( $\Phi_0$ ), the fraction of absorbed PAR ( $fAPAR$ ), and the effect of  
105 photorespiration and substrate limitation at subsaturating  $[CO_2]$  represented as a function of the leaf-internal  
106  $[CO_2]$  ( $c_i$ ) and the photorespiratory compensation point ( $\Gamma^*$ ), as shown in Eq. (1).

$$107 \quad GPP = \Phi_0 (PAR_0 \times fAPAR) (c_i - \Gamma^*) / (c_i + 2\Gamma^*) \quad (1)$$

108 where  $\Phi_0$  is set to 0.48 g C/ mol photon, based on a quantum efficiency of 0.05 mol C/ mol photon and a leaf  
109 absorptance of 0.8. Daily PAR at the top of the atmosphere is calculated based on solar geometry and is  
110 subsequently modified by transmission through the atmosphere, which is dependent on elevation and cloud  
111 cover. Annual effective PAR ( $PAR_0$ ) is calculated as the annual sum of daily PAR but taking into account the  
112 low-temperature inhibition of photosynthesis and growth, using a linear ramp function to downweight PAR on  
113 days with temperatures below 10 °C. Days with temperatures below 0°C do not contribute to  $PAR_0$ . See  
114 Wang et al. (2014) for details. In this application, we first calculated potential *GPP* with  $fAPAR$  set to 1.  
115  $fAPAR$  is not an input to the model but is calculated implicitly, from the foliage cover simulated by the T  
116 model.

117 Leaf-internal  $[CO_2]$  is obtained from the ambient  $[CO_2]$  via the ‘least-cost hypothesis’ (Wright et al., 2003;  
118 Prentice et al., 2014). Wang et al. (2014) provide a continuous prediction of the  $c_i/c_a$  ratio as a function of  
119 environmental aridity, temperature and elevation based on this hypothesis:

$$120 \quad c_i/c_a = 1 / (1 + C \sqrt{(\eta \sqrt{D}) / K}) \quad (2)$$

121 where  $D$  is the cumulative water deficit over a year (proportional to an annual “effective value” of the vapour  
122 pressure deficit: VPD),  $\eta$  is the dynamic viscosity of water,  $K$  is the effective Michaelis-Menten coefficient for  
123 Rubisco-limited photosynthesis, and  $C$  is a constant. The difference between the annual actual and equilibrium  
124 evapotranspiration is used as a proxy for  $D$  (see Prentice et al., 2013).  $D$  is calculated using the daily  
125 interpolated temperature, precipitation and cloudiness data. Annual actual evapotranspiration is derived from  
126 equilibrium evapotranspiration and precipitation using a simple soil moisture accounting scheme with a daily  
127 time step, as described in Gallego-Sala et al. (2010). The temperature dependences of  $\eta$  and  $K$  follow Prentice  
128 et al. (2014). Both  $K$  and  $\eta$  change steeply with temperature:  $K$  changes from 196 ppm at 10 °C to 1094 ppm  
129 at 30 °C ;  $\eta$  decreases from 1.31 mPa s at 10 °C to only 0.798 mPa s at 30 °C.

130 The temperature dependence of  $\Gamma^*$  is described by an exponential closely approximating an Arrhenius  
131 function (Bernacchi et al., 2003):

$$132 \quad \Gamma^* = \Gamma_{25}^* \exp(0.0512\Delta T) \quad (3)$$

133 where  $\Gamma_{25}^*$  is the value of  $\Gamma^*$  at 25 °C (4.331 Pa), and  $\Delta T$  is the monthly temperature difference from 25 °C.

134 The P model has been shown to simulate well many of the global patterns of annual and maximum monthly  
135 terrestrial GPP by C<sub>3</sub> plants. The simulated seasonal cycle of GPP at different latitudes is supported by  
136 analyses of CO<sub>2</sub> flux measurements (Wang et al., 2014).

### 137 2.1.2 The T model

138 We assume that potential GPP is the first-order driver of tree growth both at stand and individual level. The T  
139 model translates potential GPP as simulated by the P model into individual tree growth, which depends on  
140 foliage cover within the canopy and the respiration of non-green tissues, carbon allocation to different tissues,  
141 and relationships between different dimensions of the tree. Although these relationships are often loosely  
142 called “allometries”, true allometries (power functions) have the undesirable mathematical property for  
143 growth modelling that, if the power is greater than one, the derivative evaluated at the start of growth is zero;  
144 if the power is between zero and one, the derivative is infinite. We have therefore avoided the use of power  
145 functions, except for geometric relationships where they are unambiguously correct.

#### 146 *Functional geometric relationship*

148 Carbon is allocated to different tissues within the constraint of the basic functional or geometric relationships  
149 between different dimensions of the tree.

150 Asymptotic height-diameter trajectories (Thomas, 1996; Ishii et al., 2000; Falster and Westoby, 2005) are  
151 modeled as:

$$152 \quad H = H_m [1 - \exp(-aD/H_m)] \quad (4)$$

153 where  $H$  is the tree height,  $D$  is the basal diameter,  $H_m$  is the (asymptotic) maximum height, and  $a$  is the initial  
154 slope of the relationship between height and diameter.

155 The model also requires the derivative of this relationship:

$$156 \quad dH/dD = a \exp(-aD/H_m) = a (1 - H/H_m). \quad (5)$$

157 The form of the stem is assumed to be paraboloid (Jonson, 1910; Larsen, 1963). It can be shown (assuming  
158 the pipe model) that this form is uniquely consistent with a uniform vertical distribution of foliage area during  
159 early growth, i.e. in the absence of heartwood. Here, the total stem mass ( $W_s$ ) is expressed as a function of  $D$   
160 and  $H$ :

161  $W_s = (\pi/8) \rho_s D^2 H$  (6)

162 where  $\rho_s$  is the density of the wood, and  $(\pi/8) D^2 H$  is the volume of a paraboloid stem.

163 The relationship of diameter increment to stem increment is then given by:

164  $dW_s/dt = (\pi/8) \rho_s [D^2 (dH/dD) + 2 DH] dD/dt$  (7)

165 The projected crown area ( $A_c$ ) is estimated from  $D$  and  $H$  using an empirical relationship:

166  $A_c = (\pi c/4a) DH$  (8)

167 where  $c$  is the initial ratio of crown area to stem cross-sectional area. This relationship was chosen as an  
 168 intermediate between previously published expressions that relate  $A_c$  either to  $D^2$  or  $H$ . It is consistent with  
 169 reported allometric coefficients typically between 1 and 2 for the relationship between  $A_c$  and  $D$ .

170 Crown fraction ( $f_c$ ) is also derived from  $H$  and  $D$ . As we assumed the stem to be paraboloid, the stem cross  
 171 sectional area at height  $z$  is :

172  $A_s(z) = A_s (1-z/H)$  (9)

173 where  $A_s$  is the basal area:  $A_s = (\pi/4)D^2$ . We find the height ( $z^*$ ) at which the ratio of foliage area ( $A_f$ ) to stem  
 174 area at height  $z^*$  ( $A_s(z^*)$ ) is the same as the initial ratio of crown area to stem cross-sectional area ( $c$ ). We  
 175 obtain crown area ( $A_c$ ) from:

176  $A_c = c A_s(z^*) = c A_s (1-z^*/H)$ . (10)

177 Combining this with Eq. (8), we obtain  $(\pi c/4a) DH = c A_s (1-z^*/H)$ , which reduces to:

178  $f_c = (1-z^*/H) = H/aD$ . (11)

179 The initial slope ( $a$ ) is in principle dependent both on species growth form and on ambient conditions,  
 180 including light availability. Here it is determined directly from observations.

### 181 *Carbon allocation*

182 Actual GPP ( $P$ ) is obtained from potential GPP ( $P_0$ ) using Beer's law (Jarvis and Leverenz, 1983):

183  $P = P_0 A_c (1 - \exp(-kL))$  (12)

184 where  $k$  is the extinction coefficient for PAR, and  $L$  is the leaf area index within the crown.

185 Net primary production (NPP) is derived from annual GPP, corrected for foliage respiration (which is set at  
 186 10% of total GPP, an approximation based on the theory developed by Prentice et al., 2014 and Wang et al.,

187 2014), by further deducting growth respiration and the maintenance respiration of sapwood and fine roots.  
 188 Growth respiration is assumed to be proportional to NPP, following:

$$189 \quad P_{net} = y(P - R_m) = y(P - W_s r_s - \zeta \sigma W_f r_r) \quad (13)$$

190 where  $P_{net}$  is NPP,  $R_m$  is the maintenance respiration of stem and fine roots, and  $y$  is the ‘yield factor’  
 191 accounting for growth respiration. Total maintenance respiration of non-green parts comprises fine-root  
 192 respiration ( $\zeta \sigma W_f r_r$ , where  $\zeta$  is the ratio of fine-root mass to foliage area,  $\sigma$  is the specific leaf area,  $W_f$  is the  
 193 mass of carbon in foliage ( $(1/\sigma) L A_c$ ), and  $r_r$  is the specific respiration rate of fine roots), and stem (sapwood)  
 194 respiration ( $W_s r_s$ , where  $W_s$  is the mass of carbon in sapwood, and  $r_s$  is the specific respiration rate of  
 195 sapwood).  $W_s$  can be estimated from  $A_c$  through the pipe model:

$$196 \quad W_s = L A_c v_H \rho_s H_f \quad (14)$$

197 where  $v_H$  is the Huber value (ratio of sapwood to leaf area; Cruiziat et al., 2002), and  $H_f$  is the mean foliage  
 198 height  $H(1 - f_c/2)$ . The constraint that the initial sapwood area must be equal to the stem cross-sectional area  
 199 leads to the following identity:  $L c v_H = 1$ .

200 NPP is allocated to stem increment ( $dW_s/dt$ ), foliage increment ( $dW_f/dt$ ), fine-root increment ( $\zeta \sigma dW_f/dt$ ),  
 201 foliage turnover ( $W_f/\tau_f$ , where  $\tau_f$  is the turnover time of foliage), and fine-root turnover ( $\zeta \sigma W_f/\tau_r$ , where  $\tau_r$  is  
 202 the turnover time of fine roots). For simplicity, in common with many models, we do not consider allocation  
 203 to branches and coarse roots separately from allocation to stem:

$$204 \quad P_{net} = dW_s/dt + (1 + \zeta \sigma) dW_f/dt + (1/\tau_f + \zeta \sigma/\tau_r) W_f \quad (15)$$

205 From Eq. (13) and Eq. (15), the stem increment ( $dW_s/dt$ ) can now be expressed as:

$$206 \quad dW_s/dt = y A_c [P_0(1 - \exp(-kL)) - \rho_s(1 - f_c/2) H r_s/c - L \zeta r_r] \\
 207 \quad - L (\pi c/4a) [aD(1 - H/H_m) + H] (1/\sigma + \zeta) dD/dt - L A_c(1/\sigma\tau_f + \zeta/\tau_r) \quad (16)$$

208 The annual increment in ( $dD/dt$ ) and all the other diameter-related indices are simulated by combining Eq. (7)  
 209 and Eq. (16).

### 210 2.1.3 Definition of the growing season

211 The season over which GPP is accumulated (i.e. the effective growing season) is defined as running from July  
 212 in the previous year through to the end of June in the current year. This definition is consistent with the fact  
 213 that photosynthesis peaks around the time of the summer solstice (Bauerle et al., 2012) and that maximum leaf  
 214 area occurs shortly after this (Rautiainen et al., 2012). Carbon fixed during the later half of the year (July to  
 215 December) is therefore either stored or allocated to purposes other than foliage expansion. Observations of  
 216 tree radial growth show that it can occur before leaf-out (in broadleaved trees) or leaf expansion (in  
 217 needleleaved trees), thus confirming that some part of this growth is based on starch reserves from the

218 previous year (Michelot et al., 2012). This definition of the effective growing season is also supported by  
219 analyses of our data which showed that correlations between simulated and observed tree ring-widths are  
220 poorer when the model is driven by GPP during the current calendar year rather than an effective growing  
221 season from July through June.

## 222 2.2 Model application

### 223 2.2.1 Observations

224 We use site-specific information on climate and tree growth from a relatively low-elevation site (ca 128°02'E,  
225 42°20'E, 800 m a.s.l.) in mixed conifer and broadleaf virgin forest in the Changbai Mountains, northeastern  
226 China (Bai et al., 2008). This region was chosen because there is no evidence of human influence on the  
227 vegetation, and the forests are maintained by natural regeneration. Data on tree height, diameter and crown  
228 area were collected for 400 individual *Pinus koraiensis* trees from thirty-five 20m by 20m sample plots. The  
229 400 trees included all individual of this species in the 35 plots, i.e. represent a complete sampling of the  
230 variability in growth. Tree height and diameter were measured directly, and crown area measured as the area  
231 of projected ground coverage. Tree-ring cores were obtained from 46 of these individuals in 2007. The  
232 selected trees were either from the canopy layer or from natural gaps in the forest, and in both cases not  
233 overshadowed by nearby individuals in order to minimize the possible effects of competition. An attempt was  
234 made to select individuals of different diameters (diameter at breast height from 10 to 70 cm at time of  
235 sampling), broadly corresponding to the range of diameters recorded in the original sampling. The 46 trees  
236 were of different ages (ranging from < 50 to ca. 200 years at the time of sampling, 2006); subsequent analyses  
237 show there is little relationship between age and diameter at breast height. Environmental conditions (e.g. soil  
238 depth, light availability) were relatively uniform across the sampling plot. Monthly temperature, precipitation  
239 and fractional cloud cover data, from 1958 onwards, were obtained from Songjiang meteorological station  
240 (128°15'E, 42°32'E, 591.4 m a.s.l.), which is representative of the regional climate at low elevations in the  
241 Changbai Mountains.

### 242 2.2.2 Derivation of T model parameter values

243 T model parameter values were derived from measurements made at the sampling site or from the literature  
244 (Table 1). We estimated the initial slope of the height-diameter relationship ( $a$ : 116), the initial ratio of crown  
245 area to stem cross-sectional area ( $c$ : 390.43), and maximum tree height ( $H_m$ : 25.33 m) using nonlinear  
246 regression applied to diameter at breast height ( $D$ ), tree height ( $H$ ), and crown area ( $A_c$ ) measurement on all  
247 the 400 trees from the sample plots (Fig. 2). We used a value of sapwood density derived from three  
248 measurements at the sampling site (Table 1). We used values of leaf area index within the crown ( $L$ ), specific  
249 leaf area ( $\sigma$ ), foliage turnover time ( $\tau_f$ ), and fine-root turnover time ( $\tau_r$ ) for *Pinus koraiensis* from field studies  
250 conducted in northeastern China (Table 1). No species-specific information was available for the PAR  
251 extinction coefficient ( $k$ ), yield factor ( $y$ ), ratio of fine-root mass to foliage area ( $\zeta$ ), fine-root specific



252 respiration rate ( $r_r$ ), or sapwood specific respiration rate ( $r_s$ ). We therefore used published values for other  
253 species of evergreen needleleaf trees, taken from papers that summarise results from a range of field  
254 measurements. Most of the published values for these parameters fall in a relatively narrow range (Table 1).  
255 The uncertainty in fine-root specific respiration rate is not given in the original source paper (Yan and Zhao,  
256 2007) but the average value is consistent with other studies (e.g. Zogg et al., 1996). The published values for  
257 sapwood-specific respiration rate in pines show considerable variability, ranging from 0.5~10, or even 20  
258 nmol/mol/s (Landsberg and Sands, 2010). Analyses (see Sect. 3.1) show that the model is sensitive to the  
259 specification of sapwood respiration. We therefore selected the final value for this parameter based on  
260 calibration of the simulated mean ring width against observations, constrained by the published range of  
261 values for sapwood respiration rate.

### 262 2.3 Model application

263 We applied the model to simulate the growth of 46 individual *Pinus koraiensis* trees from the study site  
264 between 1958-2006. The 46 trees were of different ages (ranging from <50 to ca 200 years at the time of  
265 sampling, 2006) and different diameters (diameter at breast height from 10 to 70 cm at time of sampling).  
266 Environmental conditions (e.g. soil depth, light availability) were relatively uniform across the sampling plot.  
267 The start date for the simulations was determined by the availability of local climate data. Site latitude,  
268 elevation, and observed monthly temperature, precipitation, fraction of cloud cover were used as input for the  
269 P model. Each tree was initialised at its actual diameter at 1958, calculated from the measured diameter in  
270 2007 and measured radial growth between 1958 and 2007. The model was initially run with a fixed CO<sub>2</sub>  
271 concentration of 360ppm. To examine the impact of changing atmospheric CO<sub>2</sub> levels on tree growth, we  
272 made a second simulation using the observed monthly CO<sub>2</sub> concentration between 1958 and 2006 (310 - 390  
273 ppm: NOAA ESRL).

### 274 2.4 Statistical methods

275 For statistical analyses and comparison with observations, the individual trees were grouped into three  
276 cohorts, based on their age in 1958: young (0-49 years), mature (50-99 years), and old (>100 years).  
277 Individual trees within each cohort exhibit a range of diameters: young ca 20-37 cm, mature: 9-59 cm, and  
278 old: 25-40 cm. These differences in size will affect the expression of ontogeny within each cohort. The mean  
279 and standard deviation (SD) of year-by-year diameter growth was calculated for each age cohort from the  
280 observations and the simulations. The Pearson correlation coefficient and root mean squared error (RMSE)  
281 were used to evaluate the degree of agreement between the observations and simulations. We used generalised  
282 linear modelling (GLM: McCullagh, 1984) to analyse the response of tree growth to the major climate factors  
283 and age. The GLM approach is helpful for separating the independent influence of individual factors on tree  
284 growth, given the inevitable existence of correlations between these factors.

285

## 287 3.1 Simulated ring width versus observation

288 There are only small differences between different age cohorts in the mean simulated ring width, with a mean  
289 value of 1.43 mm for young trees, 1.31 mm for mature trees, and 1.37 for older trees. These values are  
290 comparable to the mean value obtained from the observations (1.48mm, 1.29mm, 1.34mm respectively).  
291 However, the general impact of ageing is evident in the decreasing trend in ring widths between 1958 and  
292 2007 within any one cohort (Fig. 3). The slope is stronger in the observations than in the simulations,  
293 indicating that the model somewhat underestimates the effects of ontogeny.

294 There is considerable year-to-year variability in tree growth. The simulated interannual variability (standard  
295 deviation) in simulated ring width is similar in all the age cohorts (0.265 mm in the young, 0.265 mm in the  
296 mature, and 0.264 mm in the old trees). This variability is somewhat less than shown by the observations,  
297 where interannual variability is 0.274mm, 0.367mm, 0.245mm respectively in the young, mature and old  
298 cohorts. The RMSE is 0.263mm, 0.332mm, and 0.284mm respectively for young, mature and old age cohorts.  
299 The correlation between the observed and simulated sequence in each cohort is statistically significant ( $P =$   
300 0.000, 0.001, 0.009 respectively for young, mature and old age cohorts).

301 Despite the fact that the model reproduces both the mean ring width and the interannual variability in tree  
302 growth reasonably well, the range of ring widths simulated for individual trees within any one cohort is much  
303 less than the range seen in the observations. This is to be expected, given that individual tree growth is  
304 affected by local factors (e.g. spatial variability in soil moisture) and may also be influenced by ecosystem  
305 dynamics (e.g. opening up of the canopy through the death of adjacent trees) – effects that are not taken into  
306 account in the model.

## 307 3.2 Parameter sensitivity analysis

308 To evaluate the sensitivity of the model to specification of individual parameters, we ran a series of  
309 simulations in which individual parameter values were increased or decreased by 50% of their reference  
310 value. For each of these simulations, the T model was run for 500 years using constant potential GPP (the  
311 mean GPP during the period 1958-2006).

312 The model simulates a rapid initial increase in ring width, with peak ring widths occurring after ca 10 years,  
313 followed by a gradual and continuous decrease with age (Fig. 4). The model is comparatively insensitive to  
314 uncertainties in the specification of fine-root specific respiration rate ( $r_r$ ), fine-root turnover time ( $\tau_f$ ), and  
315 specific leaf area ( $\sigma$ ), while leaf area index within the crown ( $L$ ), ratio of fine-root mass to foliage area ( $\zeta$ ) and  
316 fine-root turnover time ( $\tau_r$ ) have only a moderate effect on the simulated amplitude of ring width. The largest  
317 impacts on the amplitude of the simulated ring width are from initial slope of height-diameter relationship ( $a$ ),  
318 initial ratio of crown area to stem cross-sectional area ( $c$ ), and sapwood density ( $\rho_s$ ). Maximum tree height

319 ( $H_m$ ) and sapwood-specific respiration rate ( $r_s$ ) have the greatest influence on the shape of the simulated  
320 ageing curve. These two parameters also have a large impact on the amplitude of the growth of old trees. The  
321 parameters values for  $a$ ,  $c$ ,  $H_m$  and  $\rho_s$  are derived from observations, with uncertainties much less than 50%  
322 (Fig. 2). Thus, the sensitivity of the model to these parameters is not important. However, model sensitivity to  
323 sapwood respiration ( $r_s$ ) both in terms of the shape of the ageing curve and the amplitude is of greater  
324 concern, given the large range of values in the literature. Although some part of the uncertainty in the  
325 specification of sapwood respiration may be due to differences between species, the difficulty of measuring  
326 this trait accurately also contributes to the problem. For the final model, we tuned  $r_s$  against the ring-width  
327 observations. The best match with the observations was obtained with a value 1.4 nmol/mol/s, which is within  
328 the range of published values for pines (see summary in Landsberg and Sands, 2010 ).  $r_s$  is the only parameter  
329 that was tuned.

### 330 3.3 Controls on tree growth

331 The GLM analysis revealed a strong positive relationship between  $PAR_0$  and tree growth, while moisture  
332 stress (as measured by  $\alpha$ , an estimate of the ratio of actual to potential evapotranspiration) was shown to have  
333 a less steep but still positive effect (Fig. 5 and Table 2). The observed partial relationship between mean  
334 annual temperature and tree growth is negative. The Changbai Mountains are at the southern end of the  
335 distribution of *Pinus koraiensis* in China, which makes it plausible that tree growth would be inhibited during  
336 warmer years.

337 The model reproduces these observed relationships between climate factors and tree growth. The slope of the  
338 observed positive relationship with  $\alpha$  is statistically indistinguishable from the modelled slope, but the  
339 observed positive relationship with  $PAR_0$  is weaker, and the negative correlation with mean annual  
340 temperature is stronger, in the observations than in the simulations. These differences between observations  
341 and simulations could reflect the influence of an additional climate control, related to both  $PAR_0$  and  
342 temperature (e.g. cloud cover). The difference between observed and simulated effects of temperature may  
343 also be because although simulated growth is inhibited by low temperatures (through the computation of  
344  $PAR_0$ ), the current model does not include any mechanism for inhibition due to heat stress at high air and leaf  
345 temperatures.

346 The GLM analysis also showed that age, as represented by the three age cohorts, has an impact on ring width:  
347 young trees have greater ring widths than mature trees, while old trees have somewhat greater ring widths  
348 than mature trees. This pattern is seen in both the observations and simulations, although the differences  
349 between the young and mature cohorts are slightly greater in the observations.

350 The overall similarity in the observed and simulated relationships between growth rates and environmental  
351 factors confirms that the T model performs realistically. The observed relationships are considerably noisier

352 than the simulated relationships (Fig. 5, Table 2), reflecting the fact that growth rates are affected by small-  
353 scale variability in environmental conditions as well as time-varying competition for light.

#### 354 3.4 Simulated CO<sub>2</sub> effect on tree growth

355 Elevated levels of CO<sub>2</sub> are expected to have a positive impact on tree growth (Hyvönen et al., 2007; Donohue  
356 et al., 2013; Hickler et al., 2008; Boucher et al., 2014). This positive response to [CO<sub>2</sub>] is seen in the  
357 comparison of the fixed [CO<sub>2</sub>] and real [CO<sub>2</sub>] simulations (Fig. 5). In the first part of the simulation, prior to  
358 1980, the actual [CO<sub>2</sub>] is lower than the level of 360 ppm used in the fixed [CO<sub>2</sub>] experiment: this results in  
359 lower growth rates. The 50 ppm difference between the two experiments at the beginning of the simulation  
360 results in a difference in ring width of 0.242 mm. After 1980, when the actual [CO<sub>2</sub>] was higher than 360  
361 ppm, the tree growth in the simulation with realistic [CO<sub>2</sub>] is enhanced. The 30 ppm difference at the end of  
362 the simulation results in a difference in ring width of 0.101 mm. Overall, the change in [CO<sub>2</sub>] between 1958  
363 and 2006 results in a positive enhancement of tree growth of ca 0.343 mm/yr. However, this difference is very  
364 small compared to the impact of ageing (> 1 mm from observations) or to the differences resulting from the  
365 interannual variability of climate (1.212 mm) on tree growth.

366

#### 367 4 Discussion

368 We have shown that radial growth (ring width) can be realistically simulated by coupling a simple generic  
369 model of GPP with a model of carbon allocation and functional geometric tree growth with species-specific  
370 values. The model is responsive to changes in climate variables, and can account for the impact of changing  
371 CO<sub>2</sub> and ontogeny on tree growth. Although several models draw on basic physiological and/or geometric  
372 constraints in order to simulate tree-ring indices (Fritts, 2012; Vaganov et al., 2006; Rathgeber et al., 2005;  
373 Misson, 2004), and indeed the two approaches have been combined to simulate between-site differences in  
374 ecosystem productivity and tree growth (Härkönen et al. 2010; Härkönen et al. 2013), this is the first time to  
375 our knowledge that the two approaches have been combined to yield an explicit treatment of individual tree-  
376 growth processes, tested against an extensive ring-width data set.

377 Our simulations suggest that after a brief but rapid increase for young plants, there is a general and continuous  
378 decrease in radial growth with age (Fig. 4). This pattern is apparent in individual tree-ring series, and is  
379 evident in the decreasing trend in ring widths shown when the series are grouped into age cohorts (Fig. 3). It is  
380 a necessary consequence of the geometric relationship between the stem diameter increment and cross-  
381 sectional area: more biomass is required to produce the same increase in diameter in thicker, taller trees than  
382 thinner, shorter ones. However, we find that ring widths in old trees in our study region are consistently wider  
383 than in mature trees, and this property is reproduced in the simulations (Fig. 5). This situation arises because  
384 the old trees are on average smaller than the mature trees at the start of the simulation (in 1958). Thus, while  
385 the difference between average ring-widths in the mature and old cohorts conforms to the geometric

386 relationship between stem diameter increment and cross-sectional area, it is a response that also reflects  
387 differences in the history of tree growth at this site which determined the initial size of the trees in 1958. Lack  
388 of climate data prior to 1958 or detailed information about stand dynamics precludes diagnosis of the cause of  
389 the growth history differences between mature and old trees.

390 Studies attempting to isolate the impact of climate variability on tree growth, including attempts to reconstruct  
391 historical climate changes using tree-ring series, often describe the impact of ageing as a negative exponential  
392 curve (Fritts, 2012). However, our analyses suggest that this is not a good representation of the actual effect of  
393 ageing on tree growth, and would result in masking of the impact of climate-induced variability in mature and  
394 old trees. The simulated NPP of individual trees always increases with size (or age). This is consistent with  
395 the observation that carbon sequestration increases continuously with individual tree size (Stephenson et al.,  
396 2014).

397 We have shown that total PAR during the growing season is positively correlated with tree growth at this site.  
398 This is not surprising given that PAR is the primary driver of photosynthetic carbon fixation. However, none  
399 of the empirical or semi-empirical models of tree growth uses PAR directly as a predictor variable; most use  
400 some measure of seasonal or annual temperature as a surrogate. PAR is determined by latitude and cloudiness.  
401 Although temperature varies with latitude and cloudiness, it is also influenced by other factors including heat  
402 advection. Temperature changes can impact the length of the growing season, and hence have an impact on  
403 total growing-season PAR, but this is a trivial effect over recent decades. In fact, we show that mean annual  
404 temperature *per se* is negatively correlated with tree growth at this site. Given this decoupling, and the  
405 potential that longer-term changes in cloudiness will not necessarily be correlated with changes in temperature  
406 (Charman et al., 2013), we strongly advocate the use of growing-season PAR for empirical modelling as well  
407 as in process-based modelling.

408 We found no age-related sensitivity to inter-annual variability in climate: the interannual variability in ring  
409 width is virtually identical between age cohorts. The strength of the relationship with individual climate  
410 variables is also similar between the three age cohorts. It is generally assumed that juvenile and old trees are at  
411 greater risk of mortality from environmental stress than mature trees (e.g. Lines et al., 2010; McDowell et al.,  
412 2008). This may be true in the case of extreme events, such as wildfires or windthrow, or pest attack. Our  
413 results suggest that although climate variability has an important effect on tree growth it is not an important  
414 influence on mortality.

415 We have assumed that the period contributing to growth (i.e. the effective growing season) in any year  
416 includes carbon stores generated during the second half of the previous year. The total foliage area determines  
417 the radial area of the stem, and once this is achieved NPP is allocated either to fine root production or stored  
418 as carbohydrate for use in stem growth in the early part of the subsequent year. This is consistent with  
419 observations that radial growth begins before leaf-out (Michelot et al., 2012) and that maximum leaf area is  
420 generally achieved by mid-summer (Rautiainen et al., 2012), and the MAIDEN model also allows tree growth

421 to be influenced by a fixed contribution from the previous year's growth (Misson, 2004). Defining the  
422 effective growing season as being only the current growth year had no impact on the influence of climate on  
423 ring widths, or the shape of the aging curve. It did, however, produce a considerably lower correlation  
424 between simulated and observed inter-annual variability in growth. Since tree ring-width reflects the  
425 integrated climate over the "effective growing season", reconstructions of climate variables reflect conditions  
426 during that season, not simply during the current calendar year.

427 The high degree of autocorrelation present in tree-ring series is often seen as a problem requiring pre-  
428 treatment of the series in order to derive realistic reconstructions of climate variables (e.g. Cook et al., 2012;  
429 Anchukaitis et al., 2013; Wiles et al., 2014). However, spatial or temporal autocorrelation is a reflection of the  
430 causal mechanism underpinning the observed patterning. Here we postulate that the mechanism that gives rise  
431 to the temporal autocorrelation in tree-ring series is the existence of carbon reserves that are created in one  
432 year and fuel early growth in the next. If a large reserve of carbon is created in the second half of the growing  
433 season, because of favourable conditions, this will offset poor conditions in the following year. However,  
434 large reserves may not be necessary if conditions during the subsequent growing year are very favourable.  
435 The fact that the relative influence of one year on the next can vary explains why the measured autocorrelation  
436 strength in a given tree-ring series varies through time.

437 The T model is sensitive to the values adopted for some parameters, specifically the initial slope of height-  
438 diameter relationship ( $a$ ), the initial ratio of crown area to stem cross-sectional area ( $c$ ), maximum tree height  
439 ( $H_m$ ), sapwood density ( $\rho_s$ ), sapwood specific respiration rate ( $r_s$ ), leaf area index within the crown ( $L$ ), ratio of  
440 fine-root mass to foliage area ( $\zeta$ ) and fine-root turnover time ( $\tau_r$ ). Several of these parameters are easily  
441 derived from observations (e.g.  $a$ ,  $c$ ,  $H_m$ ,  $\rho_s$ ,  $L$ ) and provided that sufficient site-based observations are  
442 available should not pose a problem for applications of the model. However, the model is also sensitive to less  
443 easily measured parameters, including sapwood respiration, root respiration and the ratio of fine roots to  
444 leaves. Estimates of values for root respiration and root mass to foliage area in the literature do not show  
445 substantial differences, and we therefore used an average value to parameterize our model. This approach  
446 could be used for other applications. We parameterized fine-root turnover rates based on observations on  
447 *Pinus koriensis* from Changbai. While this obviated the need for tuning in the current application, lack of data  
448 on fine-root turnover rates in other regions (or for other species) could pose problems for future applications  
449 of the model. The model is also highly sensitive to the parameter value used for sapwood respiration and the  
450 range of reported values is large (Table 1). Because of this, we derived a value for sapwood respiration by  
451 tuning the model to obtain a good representation of average ring width. This is the only parameter that  
452 requires tuning in the current version of the T model. Although sapwood respiration is difficult to measure, it  
453 would certainly be better if more measurements of sapwood respiration were available, as this would remove  
454 the need for model tuning.

455 Our modelling approach integrates the influence of climate, [CO<sub>2</sub>] and ontogeny on individual tree growth.  
456 Such a model is useful to explore the response of tree growth to potential future changes in climate, and the  
457 impact of changes in tree growth on carbon sequestration. We also envisage that it could also be used to  
458 investigate the impact of past climate changes on tree growth. Reconstructions of temperature changes beyond  
459 the recent observational period, used as a baseline for the detection of anthropogenic influences on the climate  
460 system, are largely derived from statistical reconstructions based on tree-ring series (Jones et al., 1998; Esper  
461 et al., 2002; Hegerl et al., 2006; Mann et al., 2008; Ahmed et al., 2013). However, as we show here,  
462 temperature is neither the only, nor the most important, influence on tree growth. This may help to explain  
463 why correlations between ring-widths and climate at individual sites appear to have broken down in recent  
464 decades (the so-called divergence problem: D'Arrigo et al., 2008). The availability of a robust model to  
465 investigate tree growth could help to provide better reconstructions of past climate changes (see e.g. Boucher  
466 et al., 2014) as well as more plausible projections of the response of tree growth to continuing climate change  
467 in the future.

468

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666

667 Table and Figure Captions

668 Table 1. Parameter description and the derivation of parameter values.

669 Table 2. GLM analysis of tree growth response to the climatic factors and age, based on simulations and  
670 observations. The dependent variable is mean ring width series (1958 to 2006) for each age cohort (young,  
671 mature, old). The independent variables are the growing-season total annual photosynthetically active  
672 radiation (PAR), mean annual temperature (MAT), the ratio of actual to potential evapotranspiration ( $\alpha$ ), with  
673 age-cohort treated as a factor.

674 Figure 1. Model application flow. We combined the simple light-use efficiency and photosynthesis model (P  
675 model) with a carbon allocation and functional geometric tree growth model to simulate tree growth (e.g. ring  
676 width). The inputs to the P model are latitude, elevation, [CO<sub>2</sub>], and monthly temperature, precipitation, and  
677 fractional cloud cover. Potential gross primary productivity (GPP), output by the P model, drives the T model,  
678 together species-specific parameter values.

679 Figure 2. Estimation of parameter values for the application of the T model. Diameter at breast height ( $D$ ),  
680 tree height ( $H$ ), and crown area ( $A_c$ ) of the 400 trees from the sample plots were used for the estimation of the  
681 initial slope of height-diameter relationship ( $a$ ), and (asymptotic) maximum tree height ( $H_m$ ). Relationships  
682 among crown area ( $A_c$ ) diameter at breast height ( $D$ ) and height ( $H$ ) (Eq. 7) are used to estimate the initial  
683 ratio of crown area to stem cross-sectional area ( $c$ ).

684 Figure 3. Comparison between simulations and observations for the three age-cohorts (young: 0-49 year,  
685 mature: 50-99 year, and old >100 year). Each tree was initialised at its actual diameter at 1958, calculated  
686 from the measured diameter in 2007 and measured radial growth between 1958 and 2007. The black line is  
687 the mean of observations within each age-cohort, and grey bars are the standard deviation (SD) of individuals  
688 within each age-cohort. The blue line and bars are the mean and standard deviation from the simulations.

689 Figure 4. Parameter sensitivity analyses for the T model. A constant input of gross primary productivity (GPP)  
690 (mean during 1958-2006) was used to drive the T model to simulate tree growth for 500 years following  
691 establishment. The black line was obtained with the reference value of each parameter. The effects of an  
692 increase (150% of reference value, blue line) and a decrease (50% of reference value, red line) are also shown.

693 Figure 5. Tree growth response to climate and tree age: partial residual plots based on the GLM analysis  
694 (Table 2), obtained using the visreg package in R, are shown.

695 Figure 6 CO<sub>2</sub> effect on tree growth. Two runs, one with a fixed 360ppm [CO<sub>2</sub>] (blue line), the other with  
696 observed monthly [CO<sub>2</sub>] (red line), are compared to show the simulated effect of [CO<sub>2</sub>] on tree growth during  
697 1958~2006.

Table 1

Parameter	Code	Value	Uncertainty or Range of value from literature	Value source: Observation or Published literature
initial slope of height-diameter relationship (-)	$a$	116	$\pm 4.35$	Observation (fig. 2)
initial ratio of crown area to stem cross-sectional area (-)	$c$	390.43	$\pm 11.84$	Observation (fig. 2)
maximum tree height (m)	$H_m$	25.33	$\pm 0.71$	Observation (fig. 2)
sapwood density ( $\text{kgC m}^{-3}$ )	$\rho_s$	200	$\pm 25$	Observation
leaf area index within the crown (-)	$L$	1.8	1.5~1.96	Chen et al., 2004
specific leaf area ( $\text{m}^2 \text{kg}^{-1} \text{C}$ )	$\sigma$	14	13.22~16.82	Huo and Wang, 2007
foliage turnover time (year)	$\tau_f$	4	-	Luo, 1996
fine-root turnover time (year)	$\tau_r$	1.04	-	Shan et al., 1993
PAR extinction coefficient (-)	$k$	0.5	0.48-0.58	Pierce and Running, 1988
yield factor (-)	$y$	0.6	0.5~0.7	Zhang et al., 2009
ratio of fine-root mass to foliage area ( $\text{kgC m}^{-2}$ )	$\zeta$	0.17	$\pm 0.198$	White et al., 2000
fine-root specific respiration rate ( $\text{year}^{-1}$ )	$r_r$	0.913	-	Yan and Zhao, 2007
sapwood specific respiration rate ( $\text{year}^{-1}$ )	$r_s$	0.044 (1.4 nmol/mol/s)	0.5~10 (20) nmol/mol/s	Landsberg and Sands, 2010

Table 2

		Intercept (mm)	PAR (mm/kmol photon m <sup>-2</sup> )	MAT (mm/°C)	$\alpha$ (mm)
	Estimation	-3.123	0.625	-0.180	0.702
Observation	Error	±0.784	±0.093	±0.042	±0.301
	p value	0.004	0.000	0.000	0.021
	Estimation	-7.139	1.056	-0.078	1.142
Simulation	Error	±0.169	±0.020	±0.009	±0.065
	p value	0.000	0.000	0.000	0.000

Figure 1

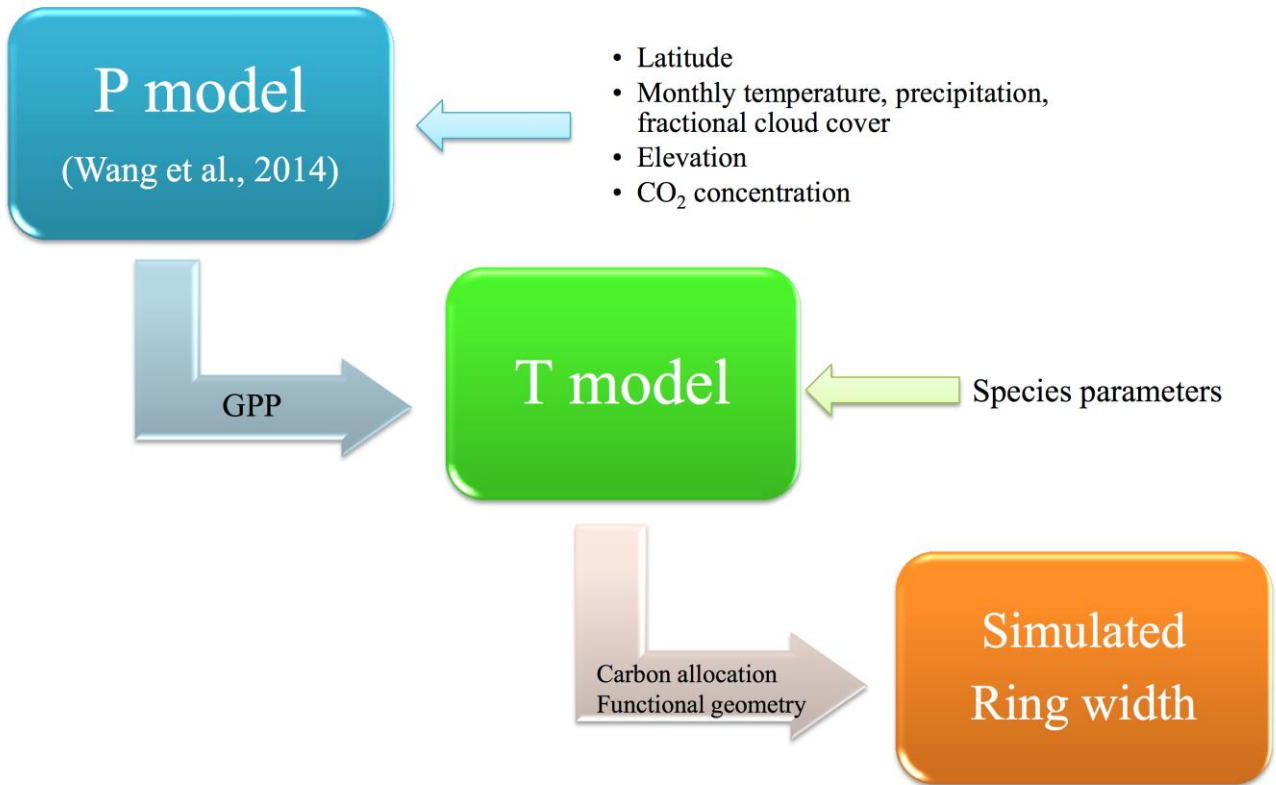




Figure 2

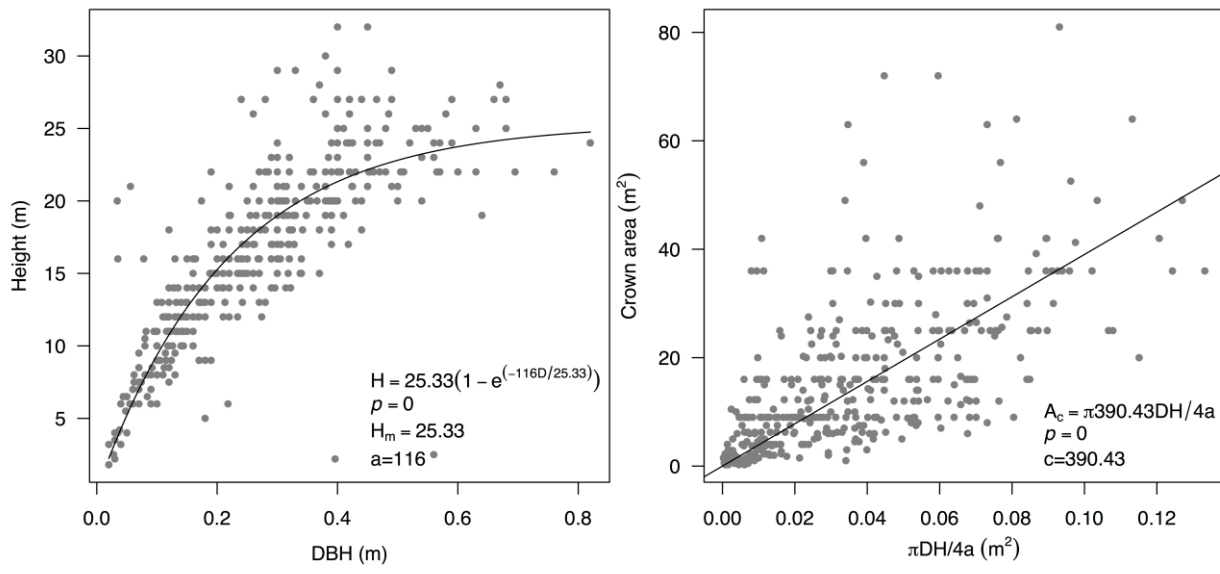


Figure 3

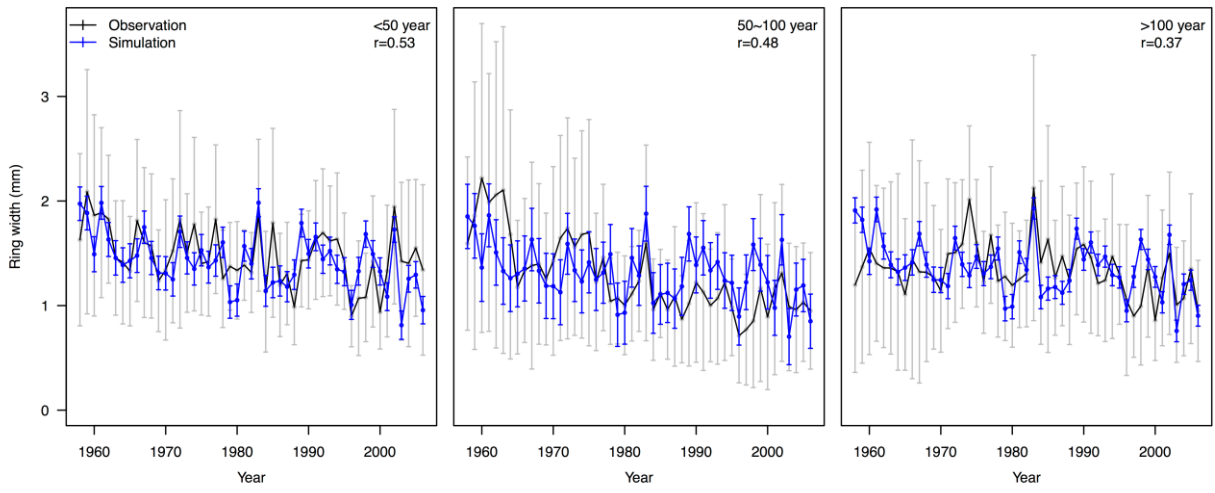


Figure 4

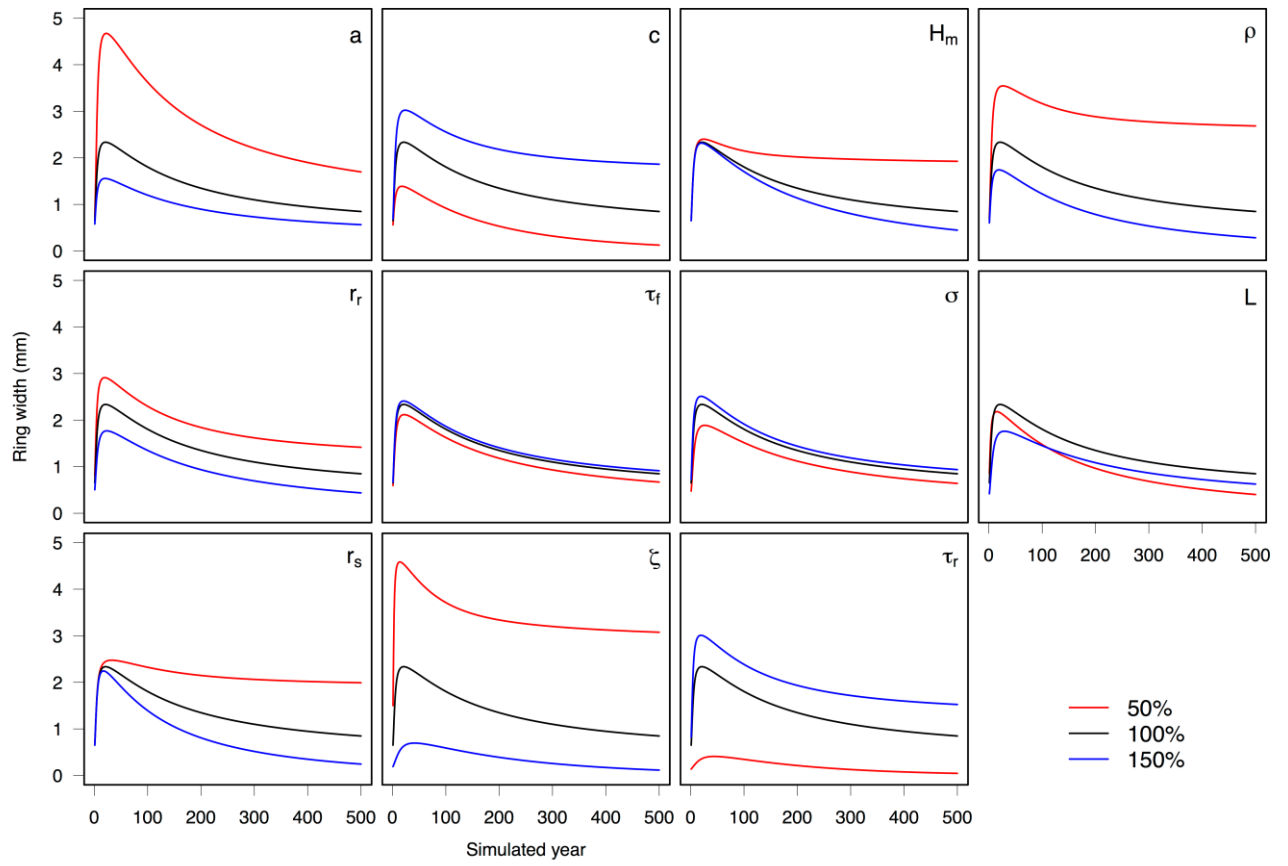


Figure 5

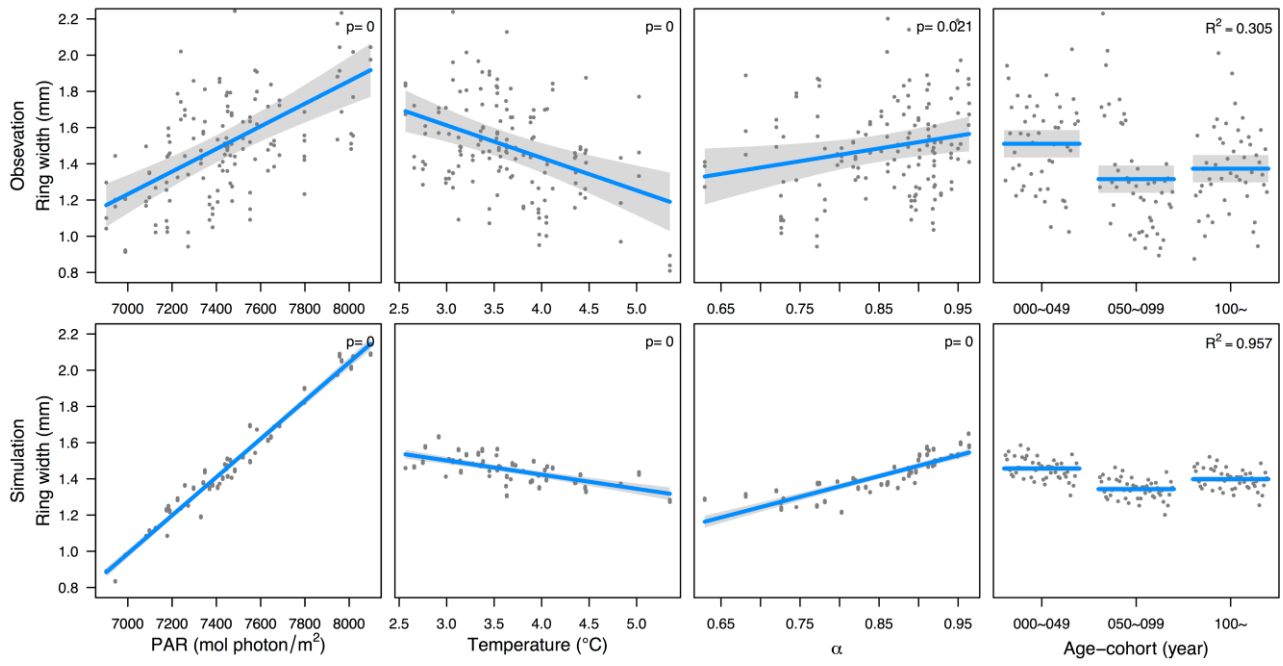


Figure 6

