



Modelling forest management within a global vegetation model-Part 1: Model structure and general behaviour

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Title: Modelling forest management within a global vegetation model – Part 1: model structure and general behaviour

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12 Abstract

13 This article describes a new Forest Management Module (FMM) that explicitly simulates forest stand growth and management within a process-based global vegetation model (GVM) called ORCHIDEE. The net primary productivity simulated by ORCHIDEE is used as an input to the FMM module. The FMM then calculates stand and management characteristics such as stand density, tree size distribution, tree growth, the timing and intensity of thinnings, wood extraction and litter generated after thinning. Some of these variables are then fed back to ORCHIDEE. These computations are made possible with a distribution-based modelling of individual tree size. The model derives natural mortality from the relative density index (*rdi*), a competition index based on tree size and stand density. Based on the common forestry management principle of avoiding natural mortality, a set of rules is defined to calculate the recurrent intensity and frequency of thinning and forestry operations during the stand lifetime. The new coupled model is called ORCHIDEE-FM (Forest Management).

14 The general behaviour of ORCHIDEE-FM is analysed for a broadleaf forest in north-eastern France. Flux simulation throughout a forest rotation compare well with literature values, both in absolute values and dynamics.

15 Results from ORCHIDEE-FM highlight the impact of forest management on ecosystem C-cycling, both in terms of carbon fluxes and stocks. In particular, the average Net Ecosystem Productivity (NEP) of $225 \text{ gC.m}^{-2}.\text{yr}^{-1}$ is close to the biome average of $311 \text{ gC.m}^{-2}.\text{yr}^{-1}$. The

NEP of the “unmanaged” case is 40% lower, leading us to conclude that management explains 40% of the cumulated carbon sink over 150 years. A sensitivity analysis reveals 4 major avenues for improvement: a better determination of initial conditions, an improved allocation scheme to explain age-related decline in productivity, and an increased specificity of the self-thinning curve and the biomass-diameter allometry.

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Keywords: forest management; global vegetation model; ORCHIDEE; carbon cycle

18

19 Introduction

Global Vegetation Models (GVMs) simulate the carbon, energy and water budgets of ecosystems on a grid. In their representation of forests, individual tree characteristics, and the processes which control them, are generally ignored. To some very rare exceptions (eg. Sato et al., 2007), most GVMs simulate the functioning of an “average tree” for forest ecosystems in each grid point and discard the effects of forest management.

In their global applications (eg. Sitch et al., 2008), GVMs usually calculate biomass to be in steady state equilibrium with climate. Discarding forest management has hitherto precluded a realistic estimation of biomass stocks in GVMs: the steady state assumption leads to overestimated biomass (Ciais et al., 2008), and to underestimated carbon sink due to forest re-growth (Desai et al., 2007; Schaefer et al., 2008; Carvalhais et al., 2010). A GVM intercomparison (Viovy et al., 2010) further indicates huge between-model differences for

aboveground biomass (ranging $0.5 - 10 \text{ kg C m}^{-2}$) simulations, illustrating the fact that GVM results are seldom evaluated against fine scale biomass data.

Replacing a forest by an average tree in a GVM raises two spatial scaling issues. The first issue is that stands of different ages coexist within the same grid point. This sub-grid heterogeneity problem can be tackled by modelling explicitly different age classes existing within each point (Zaehle et al., 2006; Shevliakova et al., 2009). The second scaling issue is that trees of different sizes coexist within the same forest stand. Forest management, which reacts in practice to the size and density of trees, is delicate to simulate in this context. For instance, Zaehle *et al.* (2006) decided in the LPJ model to remove a percentage of wood biomass in each grid point based upon a simple age criteria, based on 'top-down' timber harvest statistics only available at country scale. As a result, the carbon budget of regions where forests are intensively managed, such as Europe, cannot be confidently reproduced (Lindner et al., 2004; Zaehle et al., 2006). Another drawback of ignoring within-stand heterogeneity in GVM is that estimates by these models are difficult to relate with the most abundant source of validation and parameterization data: plot measurements from forest inventories (Valentine and Mäkelä, 2005). Indeed, forest inventories measure variables such as tree density, basal area, or standing volume, which depend on processes that call for an explicit description of within-stand heterogeneity (Dhôte, 1999).

Intensive efforts were made to simulate vegetation dynamics and individual tree characteristics through gap models (Pacala et al., 1996; Pretzsch et al., 2002; Lischke et al., 2006) and growth and yield models (Hoffmann, 1995; Dhôte and Hervé, 2000; Masera et al., 2003). Gap models

were originally developed by ecologists to simulate species succession in a newly opened gap. They represent mortality processes at tree level (Bugmann, 2001). By contrast, growth and yield models were originally developed by foresters to predict the number and size of the stems that a stand will yield. Their representation of mortality processes focuses on emergent properties at the scale of the stand (Saint-Andre et al., 2008). Both types of models are often spatially limited by the need for a local calibration of productivity which, together with rotation length, has been shown to contribute most to simulation uncertainty at regional scale (Bottcher et al., 2008). Therefore, they both need specific adaptations to be included in GVM.

This paper describes a new forest growth and management module (FMM) that is inspired from the forest growth and yield model FAGACEES (Dhôte and Hervé, 2000). It sets focus on the characteristics of individual trees within a forest stand, and can incorporate management rules based on actual forestry operations. The FMM is designed to be portable into a GVM, but it can also be applied to yield tables data, e.g. for cross validation. The FMM can simulate clear cuts, intermediate thinnings and natural mortality due to competition (self-thinning). We have coupled the FMM to a GVM called ORCHIDEE (Krinner et al., 2005).

In the following, the structure and functioning of the FMM model and its coupling to ORCHIDEE are described. Test simulations are performed for a virtual broadleaf forest in North-eastern France in order to illustrate the general response of ORCHIDEE-FM, and to assess the differences between ORCHIDEE-FM and the standard version of the ORCHIDEE GVM which simulates equilibrium biomass levels in unmanaged forests. For the ORCHIDEE-FM simulations, we consider an “unmanaged” scenario and a “managed” scenario. The expected improvements

are benchmarked using carbon stocks, carbon fluxes, and stand characteristics. The sensitivity of the ORCHIDEE-FM model to varying parameters values is evaluated. Obviously, a single example site offers an illustration of the behaviour of the FMM, but does not constitute a rigorous assessment of model performance. A follow-up paper (Bellassen et al., Part 2, this issue) presents the validation of ORCHIDEE-FM against a variety of stand-scale and continental-scale datasets provided by forest inventories, yield tables and permanent monitoring plots.

20 Model structure

20.1 Modelling strategy

Management processes can be modelled at different levels of complexity. Most often in GVMs, a constant proportion of standing biomass is simply removed from the system (Zaehle et al., 2006). Franklin et al. (2009) establish a synthetic set of equations averaging management processes, with the explicit objective of being easily added to GVMs. At a higher level of complexity, Moorcroft et al. (2001) and Sato et al. (2007) represent the evolution of each tree crown on a daily time-step, which enables them to compute photosynthesis and mortality at the tree scale. In this continuum, we opted for an intermediate level of complexity. As Sato et al. (2007), we compute the distribution of individual tree characteristics such as circumference and height, and use this information to simulate stand-scale mortality and the repartition of stand-scale growth among individual trees. We therefore move from the “average tree” modelling strategy of Zaehle et al. (2006) to an “average stand” modelling strategy similar to Desai et al. (2007). Trees of different sizes are simulated within each grid cell, and their

evolution from an initial size distribution represents the average stand composition in the cell for a series of stand ages. This fine-scale representation allows an easy comparison to real tree stands, as well as useful information for upcoming developments on wood products and physical interactions of forests with the atmosphere. Though desirable, a process-based tree-scale computation of photosynthesis and mortality is currently incompatible with the computing constraints of half-hourly flux simulation in a fully coupled Earth System Model such as IPSL-CM4 (Marti et al., 2010).

The management module (FMM) provides an explicit description of the characteristics (basal area and height) of each tree in an “average hectare”, representative of a given age-class at the resolution at which ORCHIDEE operates (typically 10-50 km² to allow for regional to global simulations). In Europe, even-aged high forests are the most common forest type (Vetter et al., 2005; Vallet et al., 2006), and their management is generally aimed at avoiding natural mortality from competition by selectively felling trees (Nabuurs et al., 2002). This is the default type of forest management simulated by the FMM, although its simulation of self-thinning and clear cutting makes it applicable to other regions by disengaging the “intermediate thinning” option.

The FMM is inspired from an existing forestry model, FAGACEES (Dhôte and Hervé, 2000). All the equations of the FMM, be they adapted from FAGACEES or not, are fully described below.

20.2 Structure of the ORCHIDEE Global Vegetation Model

20.2.1 Standard structure of ORCHIDEE

The ORCHIDEE global vegetation model (“Organizing Carbon and Hydrology In Dynamic Ecosystems”) is designed to operate from regional to global scales (Krinner et al., 2005). It is a process-driven model, composed of two main components

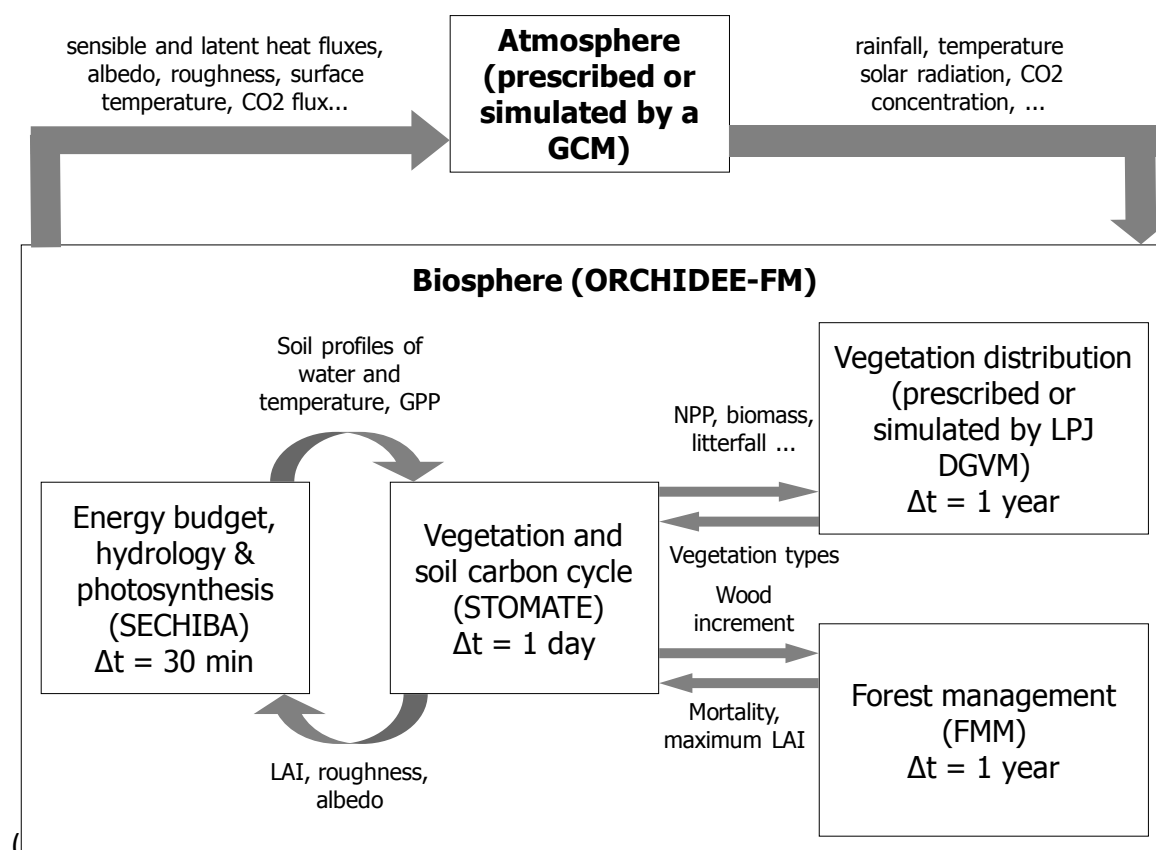


Figure 1). SECHIBA computes the energy and hydrology budget on a half-hourly basis, together with the gross primary production (GPP). These results are fed to STOMATE, the carbon cycle component. STOMATE simulates the carbon cycle on a daily basis: GPP is allocated to the different organs, and then respired by the plant or by soil micro-organisms when parts of the

plant die. These processes determine several stand-scale characteristics such as leaf area index (LAI) and canopy roughness, which are fed back to SECHIBA as they impact the energy and water budget. The equations of ORCHIDEE are given by Ducoudre et al. (1993), Krinner et al. (2005) and in <http://orchidee.ipsl.jussieu.fr/>.

ORCHIDEE requires seven climatic driving variables on a half-hourly timescale: air temperature, precipitation, specific humidity, wind speed, pressure, short wave and long wave incoming radiation. Other pedo-climatic inputs such as CO₂ concentration, soil water holding capacity, and soil texture are used at lower time-resolutions. The meteorological variables can be prescribed from climate datasets in so-called “offline” simulations. But ORCHIDEE can also be dynamically linked to the atmosphere and ocean components of the IPSL-CM4 earth system model (Marti et al., 2010) in so-called “online” simulations.

As in most global biogeochemical models, the vegetation is classified into Plant Functional Types (PFT), with 13 different PFT over the globe. Distinct PFTs follow the same set of governing equations, but with different parameter values, except for the calculation of the growing season onset and termination, which involve PFT-specific processes (Botta et al., 2000). Only the European woody PFTs (temperate needleleaf evergreen, temperate broadleaf evergreen, temperate broadleaf summergreen, boreal needleleaf evergreen, boreal broadleaf summergreen and boreal needleleaf summergreen) are of interest for this study.

20.2.2 Specific add-ons to the standard version

Explicit modelling of forest stand growth and management within the ORCHIDEE framework could not be achieved without the addition of several processes to the standard version: age-

related decline in net primary production (NPP), age-related limitation of LAI in young stands, age-related allocation ratio between stem and coarse roots, branch mortality and a coarse woody debris litter compartment.

Age-related decline in NPP

NPP has long been shown to decline in older forest stands, even if the processes underlying this decline are still subject to controversy (Gower et al., 1996; Magnani et al., 2000; Murty and McMurtrie, 2000; Lefsky et al., 2005). Three main hypothesis have been laid out to explain this phenomenon: an increase of autotrophic respiration as the tree gets bigger, a decrease of nitrogen availability from the initial litter input of tree fall or harvest residues, and hydraulic constraints on photosynthesis efficiency (Gower et al., 1996; Ryan et al., 2006). The first process is already represented in ORCHIDEE but is not sufficient to simulate a decrease of NPP with age. The two other processes were empirically added to ORCHIDEE through the introduction of a new limiting factor to photosynthesis efficiency, $decl_{factor}$ Eq. (1).

$$\left(\begin{array}{l} \text{If } age_{stand} > decl_{start} : V_{max} = decl_{factor} \times V_{max}_{std} \\ \text{And : } decl_{factor} = \max \left(decl_{max}, \frac{age_{stand} - decl_{start}}{decl_{end} - decl_{start}} \right) \end{array} \right) \quad (1)$$

where V_{max} is the photosynthesis efficiency, $decl_{factor}$ is the age-related decline factor, V_{max}_{std} is the standard value of V_{max} in ORCHIDEE, age_{stand} is the age of the stand, $decl_{max}$ is the maximum age-related decline factor, $decl_{start}$ is the age at which age-related decline starts and $decl_{end}$ is the age at which age-related decline ends.

The age-dependency of $decl_{factor}$ was calibrated on the age-related decline of aboveground wood increment from a database of European yield tables (JRC, 2009, see appendix 7.1 for details).

Age-related limitation of LAI in young stands

ORCHIDEE is highly dependent on a PFT-specific parameter setting the maximal LAI value (lai_{max}) that a PFT can reach (Jung et al., 2007). As the creation of new leaves is time and energy consuming, and because structural constraints do not always allow young trees to close the canopy, stand LAI in forests does not reach its maximum value before 10-15 years (Ovington and Madgwick, 1957; Vieira et al., 2003; Hurtt et al., 2004). This process is negligible for the standard version of ORCHIDEE which represents a steady-state equilibrium, but gets important in ORCHIDEE-FM where early stand development stages are also simulated. Therefore, lai_{max} is made dependant on age during the first years Eq. (2):

$$lai_{max} = lai_{max_std} \times \min \left(\sqrt{\frac{age_{stand}}{15}}, 1 \right) \quad (2)$$

where lai_{max} and lai_{max_std} are the maximal LAI value in $m^2 m^{-2}$ in respectively ORCHIDEE-FM and the standard version of ORCHIDEE and age_{stand} is the age of the stand in years.

Age-related allocation ratio between stem and coarse roots

The root/shoot ratio of trees has been shown to decrease with age (Mokany et al., 2006). The introduction of age in ORCHIDEE-FM allows simulating this pattern by decreasing the belowground-to-aboveground-wood allocation ratio with age Eq. (3):

$$\frac{alloc_{ab}}{alloc_{be}} = alloc_{min} + (alloc_{max} - alloc_{min}) \times \left(1 - e^{-\frac{age_{stand}}{demi_{alloc}}} \right)$$

(3)

where $alloc_{ab}$ and $alloc_{be}$ are respectively the allocation to aboveground and belowground sapwood in $gC\ m^{-2}$, $alloc_{min}$, $alloc_{max}$, and $demi_{alloc}$, are the minimum, maximum, and half-life of the aboveground/belowground sapwood allocation ratio and age_{stand} is the age of the stand in years.

Moreover, the allocation to fruits, set at 10% of NPP by Krinner et al. (2005) was reverted to 0.5%, a value more consistent with field estimates (Granier et al., 2008).

Branch mortality

Branches are usually not harvested (IFN, 2006), although the rising demand for biomass may change this in the future (European Commission, 2005). In the perspective of coupling ORCHIDEE with a forest management module, it is thus necessary to differentiate stem and branches within the aboveground biomass compartment. This is done by setting a constant PFT-specific branch/stem ratio ($branch_{ratio}$) and a constant sapwood/heartwood ratio in branches ($branch_{sap/heart}$). Two processes can lead to branch mortality: branch turnover as the tree grows, and tree mortality due to thinning (natural or anthropogenic). Since branch turnover is only one of the two processes driving branch mortality in our model, we set the branch turnover rate ($branch_{turn}$) toward the lower end of the 0.02-0.05 $year^{-1}$ range of literature values for other models (Lloyd and Farquhar, 1996; Masera et al., 2003).

224 *Coarse woody litter compartment*

225 Litter and soil carbon dynamics in the standard version of ORCHIDEE are derived from an older
226 version of the CENTURY model (Parton et al., 1988). As it was designed for grasslands, this
227 version of CENTURY only has two litter compartments: structural and metabolic. The structural
228 compartment represents the stalk of herbaceous vegetation that decomposes fairly rapidly
229 compared to woody debris. In the standard version of ORCHIDEE at steady state, this leads to
230 an underestimation of the litter pool but has little impact on fluxes as the woody litter input is
231 almost constant over time. This impact is much stronger when the forest management module
232 (FMM) is activated, as woody litter inputs are irregular and potentially large: if only a few stems
233 die after a self-thinning event, all branches and coarse roots are laid off to the decomposing
234 woody litter pool. It was thus necessary to add a coarse woody litter compartment which
235 decomposes more slowly (Lloyd and Farquhar, 1996), with a maximum turnover rate (τ_{cwd}) set
236 lower than the 4.08 year^{-1} of structural litter. Due to moisture, temperature and lignin content
237 limitations however, the actual turnover rate is much lower than its theoretical maximum of
238 0.75 year^{-1} (Table 1), averaging 0.03 year^{-1} for coarse woody debris. This value is consistent with
239 observed and simulated residence time of around 30 years (Olsson et al., 1996; Schelhaas et al.,
240 2004; Nagy et al., 2006).

20.3 *Structure of the forest management module (FMM)*

20.3.1 General structure

The general structure of the FMM, represented in Figure 2, is inspired from the FAGACEES stand-level model (Dhôte and Hervé, 2000). The FMM runs on an annual time-step, can be coupled to ORCHIDEE, and simulates three main processes: the annual distribution of cumulated stand wood increment to individual trees, the natural mortality due to self-thinning, and the timing and intensity of intermediate thinnings or clear cuts.

20.3.2 Individual growth of trees

The first step of the FMM is to allocate the yearly wood increment calculated by ORCHIDEE to a population of individual trees, here described by the distribution of their circumferences.

Initial distribution of tree circumferences

After a clear cut, the initial circumference distribution has to be prescribed. The initial number of trees is set to a default $n_{maxtrees}$ and the initial distribution of circumferences follows a truncated exponential law of parameter λ (Lanier, 1994; Dhôte and Le Moguédec, 2003):

$$\lambda = \frac{\sqrt{2}}{\pi \times Dg_{init}}$$

(4)

where the parameter Dg_{init} is the initial quadratic mean diameter.

Details on the algorithm producing the exponential distribution are given in appendix 25.2.

258 *Allocation of stand-level wood increment to individual trees*

259 To simulate competition for resources – such as light, water and nutrients – between trees, and
 260 the resulting heterogeneity in tree diameters, larger trees are assumed to grow faster in basal
 261 area (Ryan et al., 2006). The individual growth function (Eq. 5) is taken from Deleuze (2004):

$$\delta ba_i = \frac{\gamma}{2} \times \left(circ_i - m\sigma + \sqrt{(m\sigma + circ_i)^2 - 4\sigma \times circ_i} \right) \quad (5)$$

263 where δba_i is the annual increase in basal area of tree i in square meters, $circ_i$ is the
 264 circumference of tree i in meters. γ , σ and m are respectively the slope, threshold and
 265 smoothing parameters (see Figure 3): trees whose circumference is lower than σ barely grow, γ
 266 is the slope of the δba_i vs $circ_i$ relationship above σ .
 267 σ is a function of tree density within the stand, calibrated with data from permanent
 268 monitoring plots (Dhôte and Hervé, 2000).

$$\ln(\sigma) = a_\sigma \times \ln(circ_{med}) + b_\sigma \quad (6)$$

270 where $circ_{med}$ is the median circumference of trees in meters, and a_σ and b_σ are parameters.
 271 The main conceptual difference between the FMM and FAGACEES comes from how γ and σ are
 272 computed. In FAGACEES, γ represents intersite variability in stand-level growth increment and
 273 is therefore calibrated on a site-per-site basis. σ is then adjusted so that the individual
 274 circumference growths computed by Eq. (5) are consistent with total stand growth. In
 275 ORCHIDEE-FM however, the inter-site variability in stand-level growth increment is computed
 276 by ORCHIDEE. The FMM estimates σ from the median circumference from Eq. (6), and then

computes γ so that the individual circumference increments computed by Eq. (5) are consistent with the ORCHIDEE-prescribed stand woody growth. The site-by-site adjustment of γ is therefore done by iteratively computing a value of γ that yields exactly the aboveground wood increment given by ORCHIDEE (σ and m being fixed). Solving for γ requires a tree level biomass-circumference allometry relationship, given by Eq. (7) (Zianis and Mencuccini, 2004):

$$\text{biomass}_i = a_{bc} \times \text{circ}_i^{b_{bc}} \quad (7)$$

where biomass_i is the dry aboveground biomass of tree i in kg and circ_i is the circumference of tree i in centimetres.

20.3.3 Self-thinning curves

Self-thinning curves and natural mortality

Natural mortality processes in forest stands have been studied for a long time. The FMM uses the well-established Reineke rule to test whether self-thinning occurs (Eq. 8). It happens when stand density exceeds the maximum density corresponding to its quadratic mean diameter (Reineke, 1933).

$$\text{dens}_{\max} = \frac{\alpha_{st}}{Dg^{\beta_{st}}} \quad (8)$$

where dens_{\max} is stand maximum density in ind ha^{-1} (individuals per hectare), α_{st} and β_{st} are parameters, and Dg is the quadratic mean diameter (Eq. 9) in m.

$$Dg = \sqrt{\frac{\sum_i \text{diam}_i^2}{\text{dens}}}$$

(9)

where diam_i is the diameter of tree i in m and dens is the actual density of the stand.

Yang et al. (2002) showed that these relationships were not dependent on site quality, but could vary between species. This argues for a PFT-specific parameterization of Eq. (8).

Relative density index (rdi) and anthropogenic thinning

When the stand has reached a high enough dominant height – defined as the average height of the 100 tallest trees per hectare – h_{start} , human intervention through commercial thinning becomes feasible (Lanier, 1994; Grote and Erhard, 1999). In order to test this condition, the height of each tree is estimated from an allometric relationship. From the five allometric relations analysed by Newton and Amponsah (2007), a model of intermediate complexity was chosen and calibrated on data from the French national forest inventory (IFN, 2008):

$$\text{height}_i = 1.3 \times \alpha \times \text{ba}_j^\chi \times \left(1 - \exp \left(-\delta \times \text{dens}_j^\phi \times \frac{100}{2\pi} \text{circ}_i \right)^\varphi \right)$$

(10)

where height_i and circ_i are respectively the height and circumference of tree i in meters, and ba_j and dens_j are the basal area and tree density of the stand, respectively in $\text{m}^2 \text{ha}^{-1}$ and ind ha^{-1} .

α , χ , δ , Φ , and φ are parameters. Details on calibration are given in appendix 25.3.

Then, in order to avoid natural mortality and maximize wood exploitations, foresters are

assumed to remove trees from the stand by thinning, in order to maintain a lower density than

311 $dens_{max}$. To simulate this behaviour, we define the relative density index (rdi) as the ratio of
 312 actual to maximal density Eq. (11).

$$rdi = \frac{dens}{dens_{max}} \quad (11)$$

314 where rdi is the relative density index, and $dens$ and $dens_{max}$ are respectively the actual and
 315 maximal tree density of the stand in $ind\ ha^{-1}$.

316 Throughout the rotation, rdi is kept close to a targeted value rdi_{target} that depends on
 317 management practices: the lower the rdi_{target} , the more intensive the management and the
 318 lower the stand density. δrdi determines the leeway allowed around rdi_{target} : when rdi reaches
 319 $rdi_{target} + \delta rdi$, the stand is thinned until it is scaled back to $rdi_{target} - \delta rdi$ (see Figure 3). The final
 320 harvest occurs when stand density falls below $dens_{target}$ or when stand age reaches age_{target}
 321 (Lanier, 1994).

322 **20.3.4 Harvest**

323 *Tree removal*

324 In order to determine which trees are felled during a thinning event, a probability of death τ_i is
 325 attributed to each tree Eq. (12). A felling algorithm is then applied so that the rdi gets back to 1
 326 (self-thinning) or $rdi_{target} - \delta rdi$ (anthropogenic thinning) while respecting the tree-level
 327 probability of death τ_i .

$$\left(\begin{array}{l} \text{If } th_{strat} \geq 0: \tau_i = \tau_{min} + (\tau_{max} - \tau_{min}) \times \left(\frac{circ_{max} - circ_i}{circ_{max} - circ_{min}} \right)^{th_{strat}} \\ \text{If } th_{strat} < 0: \tau_i = \tau_{min} + (\tau_{max} - \tau_{min}) \times \left(\frac{circ_i - circ_{min}}{circ_{max} - circ_{min}} \right)^{|th_{strat}|} \end{array} \right) \quad (12)$$

where τ_i and $circ_i$ are probability of death and the circumference of tree i in meters, $circ_{min}$ and $circ_{max}$ are the minimum and maximum circumference in the stand in meters, and τ_{min} and τ_{max} are respectively the minimum and maximum probabilities of death.

The value of the parameter th_{strat} sets the thinning strategy: if $th_{strat} > 0$, a “thinning from below” strategy is simulated, with smaller trees preferentially thinned to obtain larger logs in the future. If $th_{strat} < 0$, a “thinning from above” strategy is simulated, with larger trees preferentially thinned thus freeing resources for smaller trees (for an illustration of this range of possible thinning strategies, see appendix 25.4).

Final harvest

Final harvest occurs at age_{target} or if a thinning event is predicted when stand density is below $dens_{target}$. All trees are cut and a new stand begins to grow in order to simulate multiple rotations over long time periods. Stems are exported while branches and coarse roots move to the litter pool as coarse woody debris. All leaves and fine roots go to the structural and metabolic litter pools, following the standard proportions set by ORCHIDEE.

20.4 Coupling: interaction between wood increment and forest management

The only input from ORCHIDEE to the FMM is the mean annual stand-level wood increment, allocated in the different biomass compartments (aboveground vs belowground, sapwood vs heartwood vs carbohydrate reserves).

The FMM feeds back three variables to ORCHIDEE: LAI, biomass and litter (see

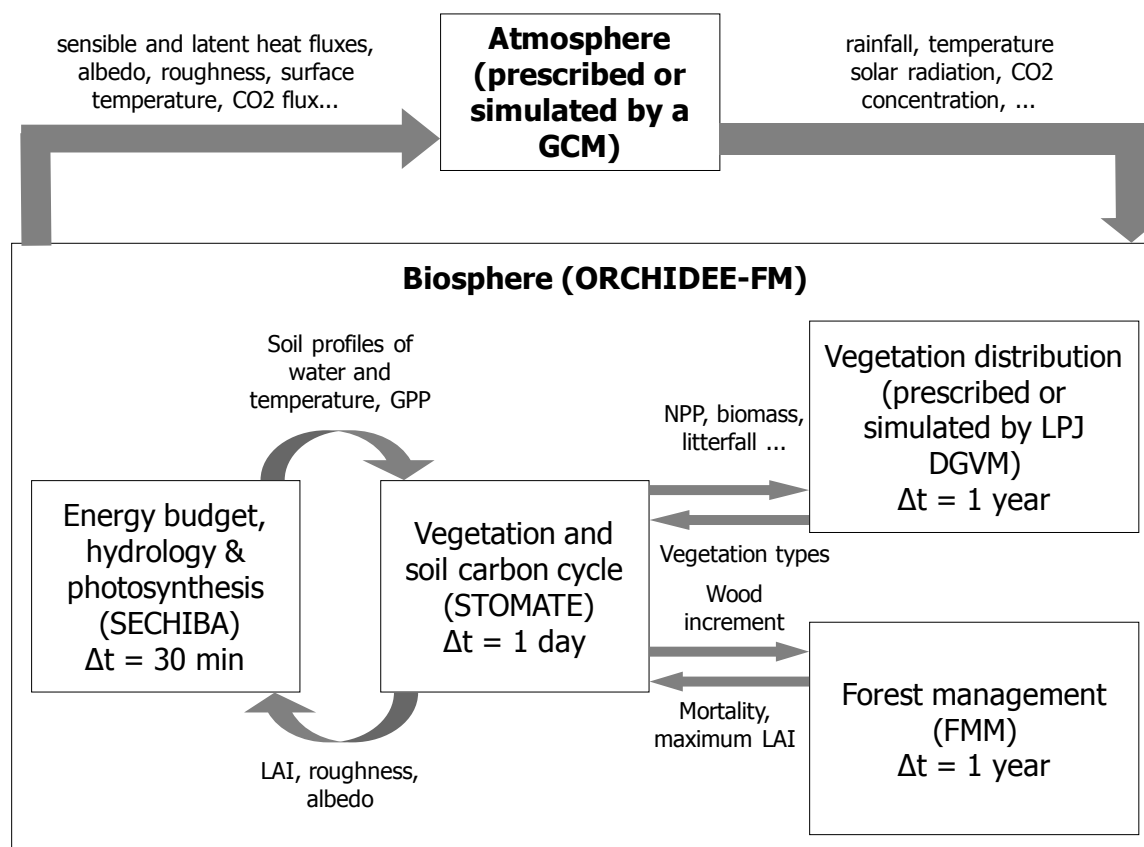
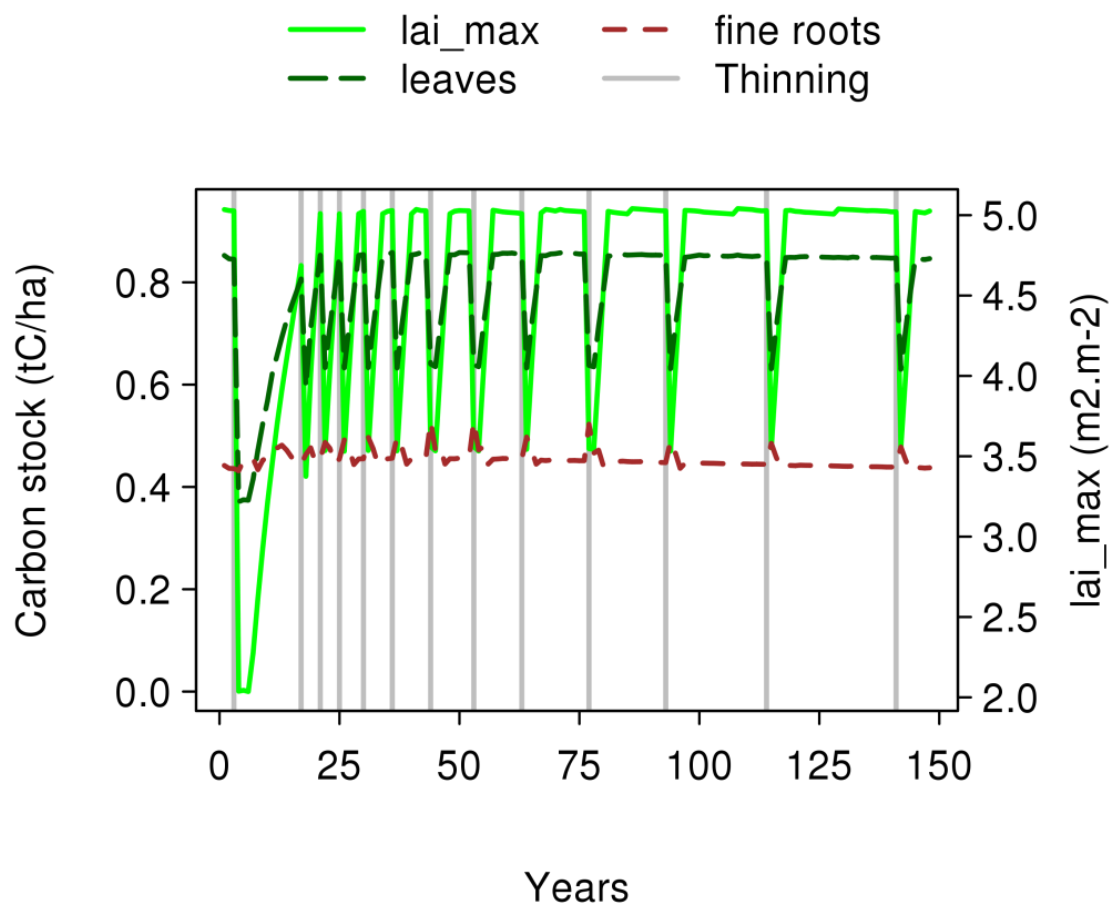


Figure 1). The soil carbon budget is indirectly impacted by the explicit simulation of branch and tree mortality in the FMM as living biomass turns – abruptly in the case of clearcuts and anthropogenic thinnings – into litter.

353 Feedback of the FMM on the leaf area index (LAI)

354 The stand-level LAI is modified by the FMM in two cases: when trees are too young to close the
 355 canopy (see part 20.2.2), and after a thinning event. Thinnings have been shown to temporarily
 356 decrease the maximum value that LAI can reach (lai_{max}), until growing branches fill the gaps.
 357 When the FMM predicts a thinning event, lai_{max} is decreased by a fixed proportion δlai_{max} , and
 358 recovers gradually within 3 years (Le Dantec et al., 2000; Bouriaud, 2003; Vesala et al., 2005).
 359 This evolution of LAI is displayed in



360

361 Figure 4.

Feedback of the FMM on biomass and litter

As detailed in part 20.3.4, the thinnings and final harvests simulated by the FMM have three types of impacts on the biomass which is fed back to ORCHIDEE:

- When self-thinning occurs, all biomass corresponding to the thinned individuals goes to the litter compartments.
- When anthropogenic thinning occurs, the stems of the thinned individuals are extracted from the stand, while the rest of their biomass (branches, roots, foliage) goes to litter compartments.
- During final harvest, all stems are exported out of the stand whereas branches, roots and foliage go to the litter pool. To close the carbon budget in simulations, the biomass corresponding to the initial circumference distribution is deducted from the old stand before harvest and allocated to the new one.

These feedbacks on biomass impact NPP as autotrophic respiration decreases. The resulting effect of the simultaneous decreases in GPP and autotrophic respiration after thinning will be discussed in the results part.

20.5 *Parameterization*

Most parameters are derived from literature, and empirical studies are preferred to modelling studies where available. Parameters for which values are available and different for broadleaves and coniferous are attributed PFT-specific values (Table 1). When the literature does not provide precise values, the French National Forest Inventory dataset (IFN, 2008) and a compilation of European yield tables (JRC, 2009) are used for calibration. The values of all

parameters specific to this version of ORCHIDEE and its associated FMM, together with their source, are summarized in Table 1.

20.6 Simulations

Three simulation set-ups are used to illustrate the impact of the FMM on the long term dynamics of carbon stocks and fluxes within the ORCHIDEE-FM framework. The first one is a control simulation using ORCHIDEE without the FMM (ORCH-STD). For the two others, the FMM is activated. In the “unmanaged case” (ORCH-FM_u), anthropogenic thinning is disabled and only self-thinning occurs. In the “managed case” (ORCH-FM_m), the full version of ORCHIDEE-FM is used.

This last set-up is also used for a sensitivity analysis of 16 key parameters. One after another (OAT approach), parameters are increased by 50% and decreased by 50%. These variations are not intended to represent a realistic range of variation or error in the parameters, but to test the response of the model to a strong variation in individual parameters.

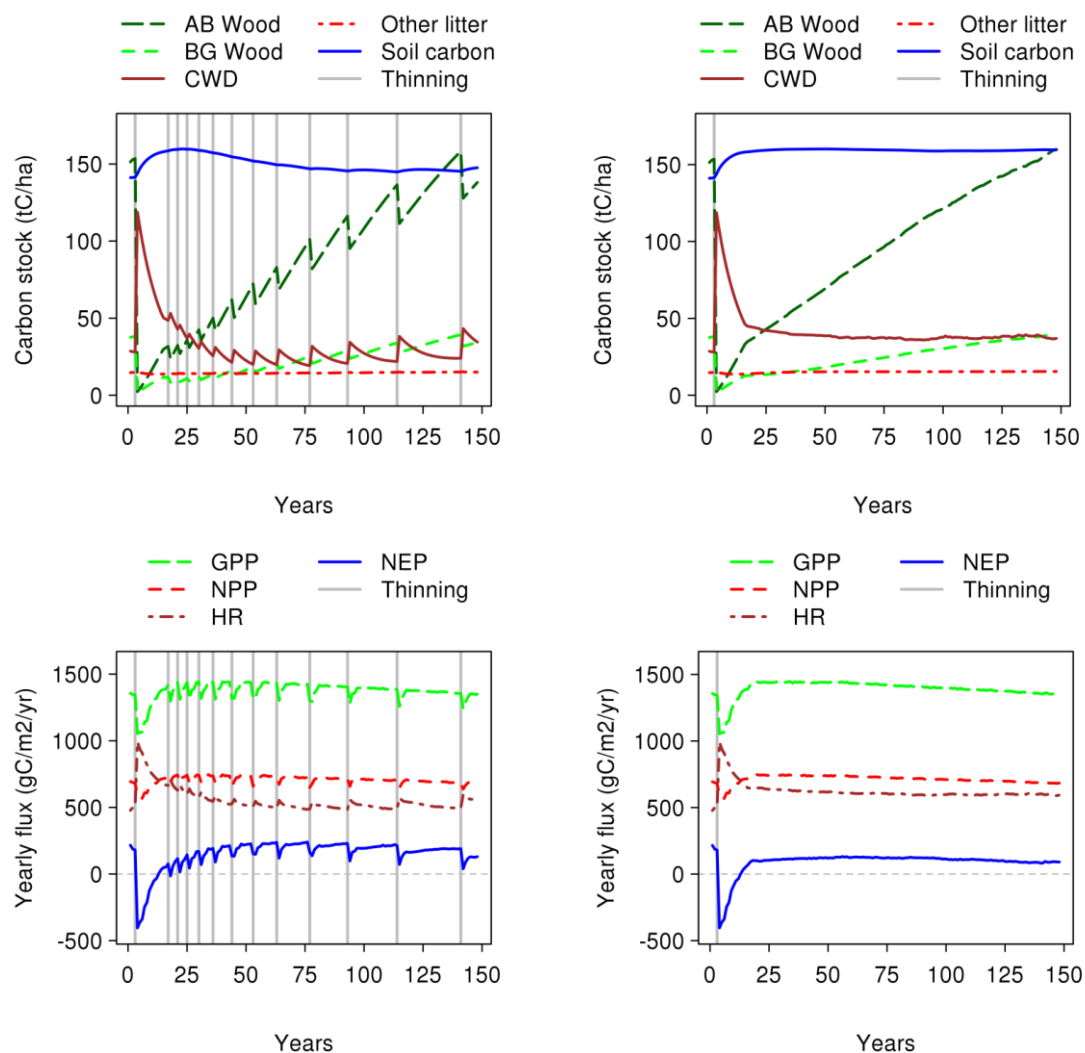
We selected a grid cell near Nancy (48.125°N7.125°E) and a plant functional type (temperate broadleaf) for which the standard version of ORCHIDEE has already been validated (Loustau, 2004). To facilitate the interpretation of carbon dynamics, we use a single year of climate that is repeated over one rotation (approx. 150 years). The selected year was 1997, close to average climate of the grid cell in terms of temperature and precipitation. Climate data comes from the 0.25° resolution REMO reanalysis (Kalnay et al., 1996; Vetter et al., 2008). CO₂ concentration is set at 380 ppm.

403 A model “spinup” is performed before any simulation to define the initial state of carbon and
404 water pools. For both the “managed” and “unmanaged” case, this “spinup” consists of
405 repeatedly running ORCHIDEE-FM for the climate of year 1997 and a CO₂ concentration of 380
406 ppm until all ecosystem carbon and water pools reach cyclical steady state equilibrium (see
407 appendix 25.5 for details). The conditions of the stand before the last clearcut are used as initial
408 conditions, and the simulation begins with a clearcut. For the control case, the “spinup” is a
409 repeated run of the standard version of ORCHIDEE instead of ORCHIDEE-FM.

21 Results

21.1 *Stand-scale results*

21.1.1 Carbon stocks

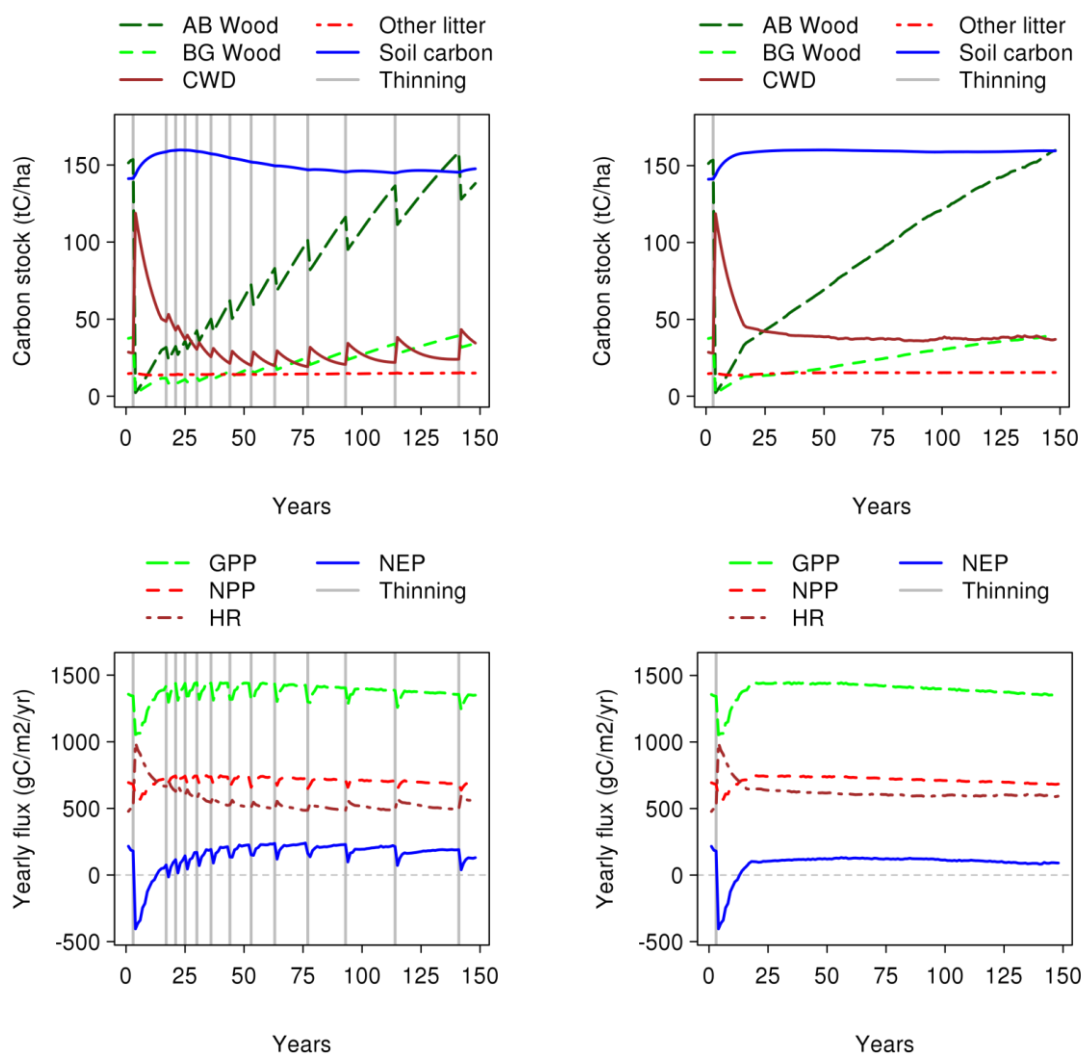


a. "Managed" case (ORCH-FM_m)

b. "Unmanaged" case (ORCH-FM_u)

Figure 5 shows the evolution of the different carbon pools during a rotation. The first year clear cut of the preceding rotation puts almost all the 30 tC ha^{-1} of belowground wood and roughly a

416 third of the 150 tC ha^{-1} of aboveground wood into litter as coarse woody debris (CWD). The
 417 decomposition of this CWD litter drives the slow initial increase in soil carbon towards 160 tC
 418 ha^{-1} . Then, as trees grow, woody biomass follows a steadily increasing trend punctuated by
 419 temporary drops after each thinning. As the initial source of litter inputs diminishes, soil carbon
 420 peaks around year 20, and then decreases. In the “unmanaged case”, where only self-thinning
 421 is allowed



a. “Managed” case (ORCH-FM_m)

b. “Unmanaged” case (ORCH-FM_u)

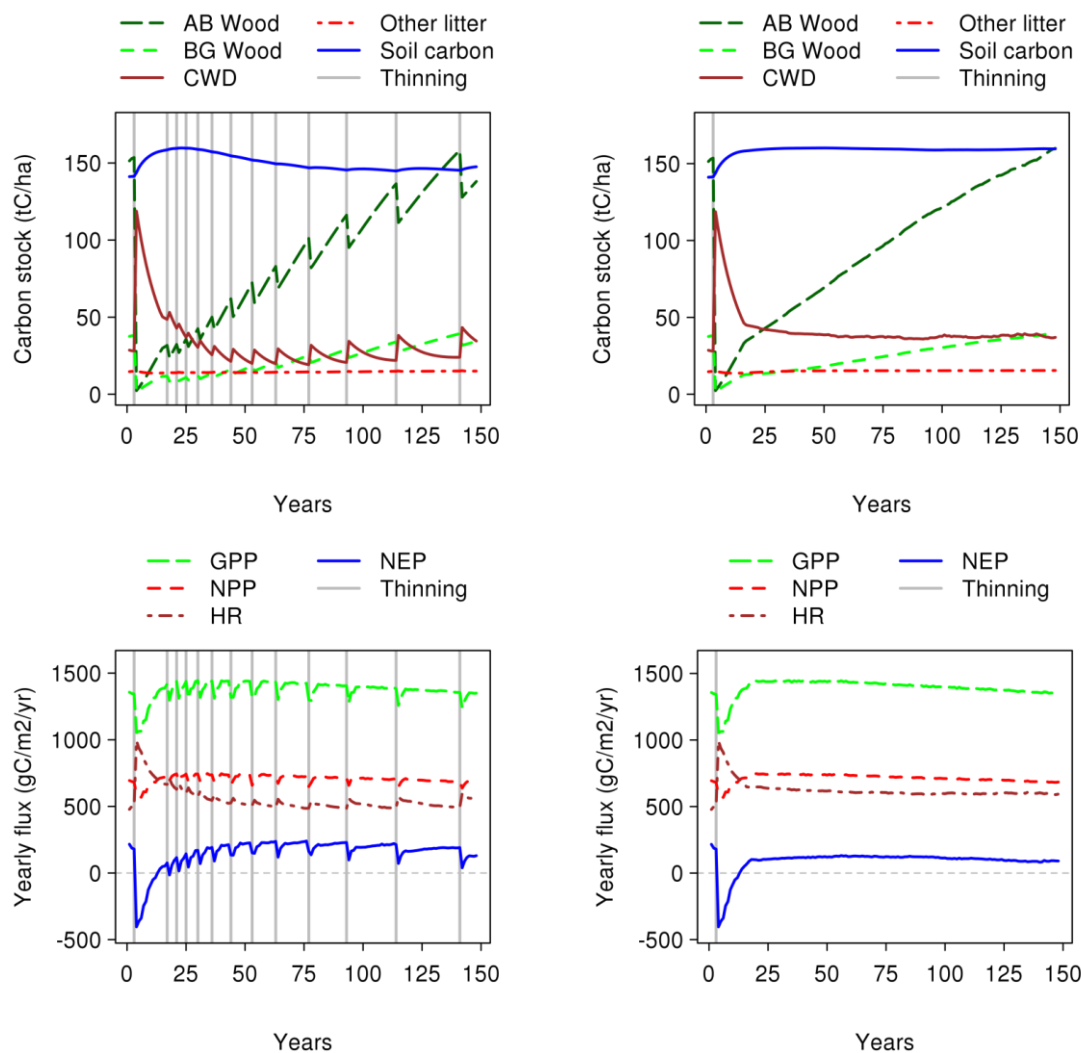
422 (

423 Figure 5), the evolution of most stocks is similar, though smoothed as they do not undergo the
424 periodic disturbance of anthropogenic thinning. Two exceptions are CWD and soil carbon which
425 keep being fed by non-exported dead stems, and reach a different equilibrium.

426 Figure 6b shows that above-ground biomass is only slightly lower (-10 tC ha^{-1} on average) when
427 the forest is regularly thinned. The main difference between the “managed” (ORCH-FM_m) and
428 “unmanaged” (ORCH-FM_u) cases in terms of biomass is seen in the coarse woody debris
429 compartment which, continuously fed by dying trees in the unmanaged case, is $10\text{-}20 \text{ tC ha}^{-1}$
430 higher. The comparison with the control (Figure 6a) highlights the 30% lower value of soil
431 carbon under management. The aboveground biomass catches up with the control value
432 towards the end of the rotation, after around 130 years.

433

21.1.2 Carbon fluxes

a. "Managed" case (ORCH-FM_m)b. "Unmanaged" case (ORCH-FM_u)

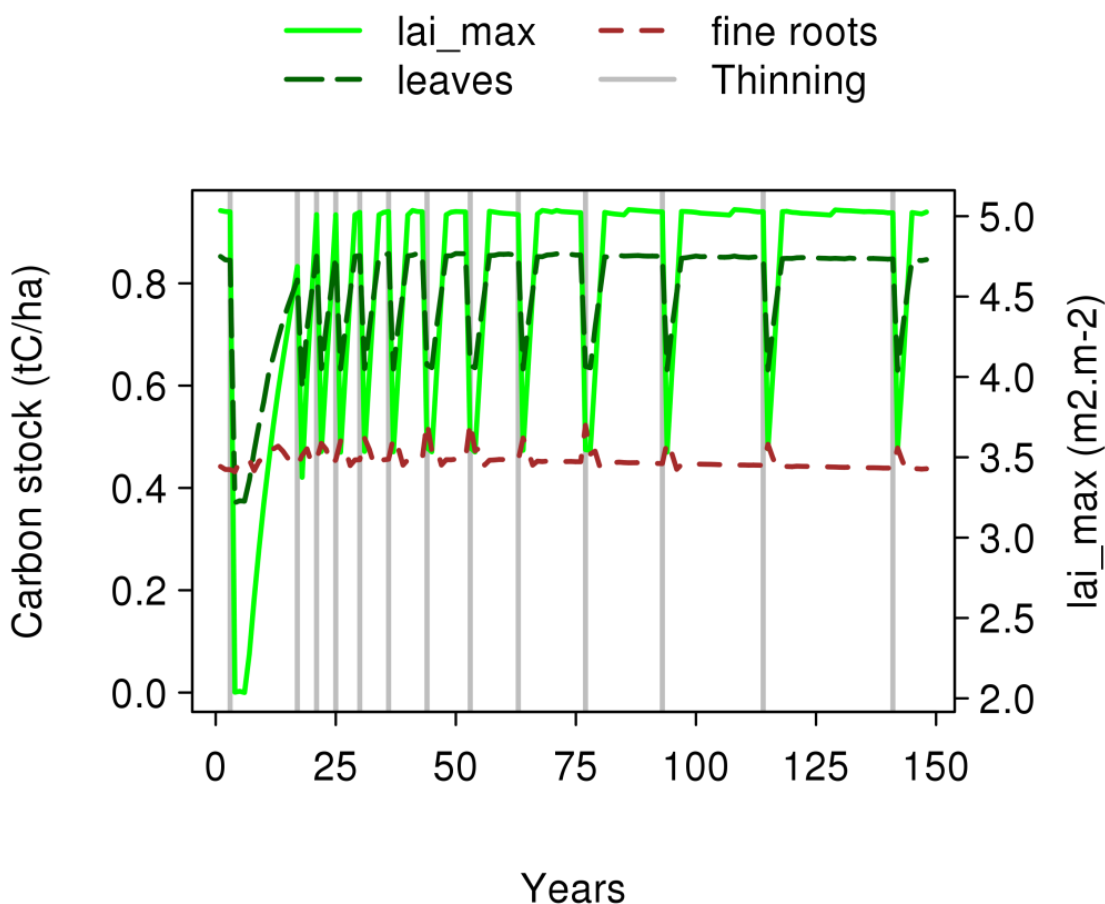
434

435 Figure 5 illustrates how the different carbon fluxes are affected by forest management. In the

436 "anthropogenic thinning" case, gross primary production (GPP) and net primary production

437 (NPP) are increasing progressively to reach 1 450 gC m⁻² yr⁻¹ and 700 gC m⁻² yr⁻¹ respectively

438 during the first 15 years, together with a gradual increase in LAI



439 (

440 Figure 4). After 50 years, the age-related decline of photosynthesis efficiency weighs on GPP
 441 and NPP, both of which slowly decline by 7-9%. Heterotrophic respiration (HR) displays a strong
 442 peak close to $1\,000\text{ gC m}^{-2}\text{ yr}^{-1}$, almost twice the level of its long-term average of $500\text{ gC m}^{-2}\text{ yr}^{-1}$.
 443 This peak is due to the slash inputs from the clear cut which ends the previous rotation, as all
 444 branch and belowground biomass is turned into litter. For the rest of the rotation,
 445 heterotrophic respiration follows a slightly decreasing trend as the coarse woody debris
 446 compartment is fed by anthropogenic thinnings. Net ecosystem productivity sums up these

evolutions: starting with an all-time low (source of CO₂ to the atmosphere) at -400 gC m⁻² yr⁻¹, it becomes positive (sink) after 10-15 years, peaks at 225 gC m⁻² yr⁻¹ and starts to decrease after 50 years as NPP decreases while HR increases.

Compared to an “unmanaged” scenario (Figure 6b), the main differences lie in GPP and HR. In the self-thinning scenario, GPP is smoother, without the small post-thinning decreases, and HR is kept higher as no exported stem is removed from litter inputs. This explains the consistently higher NEP in the managed case. The comparison between the managed case and the control (Figure 6a) highlights the influence of age-related decline of GPP and NPP in the FMM compared to the standard ageless version of ORCHIDEE.

The age-related decline in NPP leads to a parallel decline in wood increment. A similar pattern is observed on neighbouring national inventory plots, although the age-related decline in wood increment starts earlier in the model (Figure 7).

21.1.3 Stand characteristics

Table 2 gives the stand characteristics simulated by the model in the “managed scenario” (ORCH-FM_m). The prescribed initial density of 10 000 trees.ha⁻¹ is already reduced to 4 100 trees.ha⁻¹ after 20 years, and continues to decrease towards 150 trees.ha⁻¹ after 140 years. Basal area, standing volume, and average height all keep increasing as the stand ages, though the increase is faster during younger ages. The exported volume to total volume produced ratio increases rapidly with the first thinnings to reach 0.55 in the long term. The time interval between two thinnings also increases over time from 4 years after 20 years, up to 27 years around the end of the rotation. Finally, as the stand ages, the average circumference gets closer

to the circumference of the largest tree in the stand, reflecting the progressive change in circumference distribution (see 21.2 and appendix 25.6). All these values are within the ranges given by yield tables (JRC, 2009).

21.2 *Tree-scale results: distribution in circumference classes*

Figure 8 shows the evolution of tree circumference. The difference between before and after thinning distributions illustrates the thinning processes whereby smaller trees are preferentially thinned. As the stand ages, the circumference distribution shifts from a decreasing exponential with a majority of smaller trees towards a majority of larger trees. This is consistent with the evolution described by the local forestry guide for this type of management (Asael, 1999).

21.3 *Sensitivity analysis*

The sensitivity of stand variables to a selected set of parameters is illustrated in Figure 9. The parameters listed on the left are increased by 50% (upper part of the figure) or decreased by 50% (lower part of the figure). The model is shown to be little sensitive to the initial distribution ($dens_{init}$, $circ_{init_{min}}$, p_{max}). The most sensitive parameters are the ones dealing with the relative density index (σ , rdi_{lim} , rdi_{target} , and $selfth_{curve}$). Most variables are also very sensitive to allometric equations, and in particular the allometry between circumference and biomass Eq. (7).

22 Discussion

22.1 *Carbon stocks and fluxes*

Carbon stocks and fluxes are all within the range of reported values for temperate broadleaves (Pregitzer and Euskirchen, 2004; Luyssaert et al., 2007). The 7% difference in standing aboveground wood between “unmanaged” and “managed” cases (Figure 6b) is smaller than existing estimates of 25%-50% for moderate to high thinning regimes (Lanier, 1994; Vetter et al., 2005). The simulated thinning regime is indeed quite extensive, with a target *rdi* of 0.75. Two other explanatory factors, the uncertainty of the self-thinning parameters and the absence of thinning-related mortality, are further discussed in the context of the sensitivity analysis below.

The flux dynamics throughout the rotation also compares well with previous studies. As in Thornton, et al. (2002), CWD decomposition drives HR, and therefore determines how quickly the stand turns into a carbon sink after a clear cut. Both the amplitude of the source and the time of recovery are within the ranges of existing modelling studies, respectively of -500 to -1000 gC m⁻² yr⁻¹ and 10-20 years (van Oene et al., 2000; Thornton et al., 2002; Turner et al., 2005). This is also consistent with the empirical range of 700 to 1300 gC m⁻² yr⁻¹ for the HR of temperate forests aged between 0 and 10 (Pregitzer and Euskirchen, 2004).

As found by Lloyd and Farquhar (1996), an important part of the vegetation sink is due to the lag between NPP and litterfall. The role of management however is not negligible. In the “unmanaged scenario” (ORCH-FM_u), the cumulated NEP over a rotation of 150 years – 13500

505 gC m^{-2} – makes up only 60% of the cumulated NEP – $22\,500 \text{ gC m}^{-2}$ – in the “managed scenario”
 506 (ORCH-FM_m). As much as 40% of the sink of the “managed” scenario can therefore be
 507 attributed to management. This ability of ORCHIDEE-FM to simulate a positive NEP – i.e. a net
 508 sink – through forest growth is an important improvement for the null average of the standard
 509 steady-state simulation. While ORCHIDEE has long been able to simulate climate-related inter-
 510 annual variability and long-term trends in NEP, the absence of a management-driven sink has
 511 been singled out as a capital weakness of the model (Ciais et al., 2008; Luyssaert et al., 2010).
 512 While the results from ORCHIDEE-FM thus confirm the recent empirical findings of positive NEP
 513 in old forests (Field and Kaduk, 2004; Ciais et al., 2008; Luyssaert et al., 2008) with about 150 gC
 514 $\text{m}^{-2} \text{ yr}^{-1}$ at 150 years in the ORCH-FM_m simulation, this result has to be interpreted with caution.
 515 The narrowing of the gap between NPP and HR is mainly due to the parameterized age-related
 516 decline in NPP. This age-related decline of 9% at 150 years is found to be on the lower end of
 517 the empirical range of 0-76% (Gower et al., 1996), and much lower than the modelled value of
 518 72% (Magnani et al., 2000). Yet, the decline of 50% in aboveground wood increment that
 519 follows from it is consistent with yield tables (JRC, 2009), IFN data (IFN, 2008), and other
 520 modelling studies (Zaehle et al., 2006). This suggests that our empirical approach to age-related
 521 decline of stand NPP leads to a higher than observed wood increment decline to NPP decline
 522 ratio. A higher age-related value of $decl_{factor}$ – more consistent with estimates of age-decline in
 523 NPP but less consistent with estimates of age-related decline in wood increment – would give a
 524 lower, if not negative, value of NEP for old forests. This contradiction calls for an improvement
 525 in the allocation framework of ORCHIDEE. Attempting such an improvement will be most

meaningful when a future inclusion of the nitrogen cycle allows for more variation in the allocation to leaves.

Note that the simulated effect of a thinning is a decrease in NPP, which means that the effect of GPP decrease overcomes the effect of harvest on autotrophic respiration. Finally, the increase in HR and decrease in NPP creates a temporary but strong $150 \text{ gC m}^{-2} \text{ yr}^{-1}$ decrease of NEP following thinnings. Empirical evidence regarding the effect of a partial and temporary defoliation – such as a defoliation due to thinning – on NEP is mixed: Vesala et al. (2005) and Granier et al. (2008) find no significant effect while Allard et al. (2008) attributes a 25% decrease in NEP to an insect-induced defoliation. In particular, a compensating increase in understory GPP, which has been shown to occur in a Finnish forest (Vesala et al., 2005), would be missed by ORCHIDEE-FM which does not represent the understory. For these reasons, the simulated effect of thinnings on NEP has to be interpreted with caution.

22.2 *Parameterisation and sensitivity analysis*

22.2.1 Initial distribution of trees

As these parameters ($dens_{init}$, $circ_{init_{min}}$, p_{max}) are probably the least well known, the small sensitivity of model results to them is an important result. Nevertheless, the high uncertainty associated with these parameters means they could vary by more than 50%. A narrower initial distribution for example – with a p_{max} increased five-fold, leading to an initial maximum circumference decreased by 27% – leads to a narrower distribution throughout the whole rotation (see appendix 25.2.2). Unfortunately, measurements in densely stocked young stands

are challenging and the literature on stand characteristics during the very first years after harvest is scarce. The only reference we have is for initial biomass. At 2.5 tC ha^{-1} – or about 1.5% of before-cut biomass – the value simulated by ORCHIDEE-FM is close to the 1% of before-cut biomass used by Vetter et al. (2005).

22.2.2 Accuracy of the thinning parameters

Parameters dealing with the relative density index (σ , rdi_{lim} , rdi_{target} , and $selfth_{curve}$) are shown to be among the most sensitive in the FMM. These parameters, though better known than those governing initial distribution, still carry a relatively high uncertainty: rdi_{lim} and rdi_{target} are quite specific to the modelling strategy of the FMM, and therefore not often reported in the literature. σ and $selfth_{curve}$ have a wider theoretical interest (Jack and Long, 1996; Dhôte, 1999), but reviews of estimates for a wide range of species and climate conditions are still lacking. Such studies could greatly improve the accuracy of the FMM.

In the meantime, in order to ensure that our default values are not erroneous, we analysed the thinning pattern that follows from these parameters. The cumulated thinned volume to total volume produced before clear cut ratio, for example, is close to 0.55. This thinning pattern is on the higher end of the 0.3-0.5 range of previous European-scale modelling studies (Nabuurs et al., 2000; Nabuurs et al., 2002), but in the middle of the 0.5-0.6 range of relevant French yield tables at the end of the rotation (Vannière, 1984). This comparison shows that the thinning pattern simulated by the FMM is realistic, though the average European practice may yield lower thinned volumes. Taken together with the small difference in standing biomass between the “managed” and “unmanaged” simulations (see 22.1), this observation calls for a re-

evaluation of the self-thinning curve towards denser stands if the self-thinning scenario is to be used at European scale.

22.2.3 Allometries

The literature is abundant on the topic (Zianis and Mencuccini, 2004), but also points to species-specific variations (Vallet et al., 2006). Adding height as an explanatory variable for biomass has also been shown to improve the fit significantly (Joosten et al., 2004; Vallet et al., 2006). Refining this allometry, for example by the assimilation of remotely sensed height and/or biomass would therefore be a promising avenue of improvement for ORCHIDEE-FM.

22.2.4 Correlated effects and threshold effects

More than sensitivity alone, Figure 9 points to couples of parameters that have similar effects on model results, and to parameters exhibiting a non-linear effect:

- Branch ratio and branch turnover have the same impact on most variables through branch mortality. They only differ by their impact on the exported volume to total volume produced ratio which is only affected by branch ratio.
- Similarly, modulating the self-thinning equation ($selfth_{curve}$) or the rdi_{target} have the same qualitative impact on most variables as they both determine the acceptable tree density to quadratic mean diameter ratio.
- When the circumference threshold σ above which basal area increase takes off (see Figure 3 and equation 2) is increased by 50%, it becomes higher than most tree circumferences. As most trees are below the threshold, they all receive a more or less

equal share of the wood increment, which results in a narrow circumference distribution. This explains the higher minimum circumference and lower maximum circumference observed on Figure 9.

22.3 *Modelling strategy*

22.3.1 **Model coupling: averaged runs vs full-coupling**

In the Ecosystem Demography model, Moorcroft *et al.* (2001) do not opt of a full coupling between the GVM and a small-scale gap model. They derive the predictions of their gap model along the two most important variables, namely tree size and age since last disturbance, and apply the simplified derived function to their GVM. This approach makes sense when the small-scale model is stochastic in order to obtain the deterministic solutions expected from large-scale GVMs while keeping computing time manageable. In this study however, we adopted a full-coupling strategy between ORCHIDEE and the FMM yet on annual time scale, more akin to Friend *et al.* (1997). This strategy makes it easier to analyse the effect of management at large scales: it is possible to cut off some processes and/or amplify others, and directly keep track of the result at large scales. As the FMM is strictly deterministic, a single run per location and per age class is sufficient, helping to minimize additional computing time (8 seconds – 0.5% – more than the standard version of ORCHIDEE per rotation and per site).

22.3.2 **Model limitations and non-simulated processes**

The FMM ignores several minor stand-scale processes involved in stand dynamics over the long term and after anthropogenic thinning. While being negligible over a standard rotation, the

absence of natural regeneration in the FMM would lead to unrealistic results over the long term if no clear cut is prescribed: left to itself, the FMM would end up with a single enormous tree after a millennium. This problem also precludes the FMM from simulating uneven-aged types of management such as the selective logging widely practiced in primary tropical forests. In temperate regions however, this management type remains uncommon (Jaccaud, 2007). Regarding anthropogenic thinning, only two processes are simulated by the FMM: the biomass transfers linked to the felling of trees and the recovery of the maximum leaf area index as the branches of surviving trees fill the gaps. Other processes have been shown to occur after an anthropogenic thinning: some mortality in damaged but unharvested trees, a possible boost in productivity, a possible change in assimilate allocation and some adjustment in biomass-circumference allometries (Mitchell, 2000; Petritsch et al., 2007; Nabuurs et al., 2008). As the quantification of these processes is still very uncertain, they are ignored in the FMM.

23 Conclusion

This study describes the structure and typical results of the new ORCHIDEE-FM model. This model calculates stand and management characteristics such as stand density, tree size distribution, tree growth, the timing and intensity of thinnings, wood removals from the stand and litter generated after thinning. The general pattern simulated for a grid cell in north-eastern France was shown to be consistent with existing studies on carbon fluxes and stocks, both in absolute values and dynamics. In particular, they confirm the possibility that forests could still act as carbon sinks after a hundred years. Anthropogenic thinning leads to biomass export from the stand and decreases the litter substrate for respiration, thus explaining 40% of

the sink throughout the rotation. A thorough model-data comparison is the object of a follow-up article, at three different scales: tree, stand and regional (Bellassen et al., Part 2, this issue). The sensitivity analysis reveals 4 major leads for improvement. Two lie in the model structure itself: an in-depth study of the impact of the initial tree circumference distribution and a review of the allocation framework of ORCHIDEE to strike a better balance between age-related decline in NPP and age-related decline in wood increment. The other two require the assimilation of local information: both the self-thinning curve and the circumference-biomass allometry have been shown to be very sensitive parameters in the FMM. The most promising way of increasing their accuracy would be to fit them locally based on the dominant species, tree height and/or soil fertility. We suspect that the use of remote sensing data could bridge the gap between the large scale of GVMs and the smaller scale at which this type of information is usually collected.

Overall, our investigation supports the notion that including forest management in DGVMs will reveal a more realistic picture of biosphere-atmosphere interactions, future carbon sequestration and vulnerability of land carbon pools to climate change than focusing solely on natural forests at equilibrium.

24 Acknowledgements

We want to acknowledge the contribution of Antoine Colin (IFN), Daniel Rittié (INRA-LERFoB), and Maurizio Teobaldelli (JRC) without whom the work on the datasets they manage would have been both impossible and meaningless. We also want to thank Eric Dufrêne (CNRS-ESE) and Soenke Zaehle (MPI) for their useful suggestions on the model structure, and Sebastiaan

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25 Appendixes

25.1 Age-related decline in NPP

The age-related decline factor of photosynthesis efficiency decreases linearly with age, down to a maximum age-related decline factor of $decl_{max}$ (cf. Eq. 1, main text).

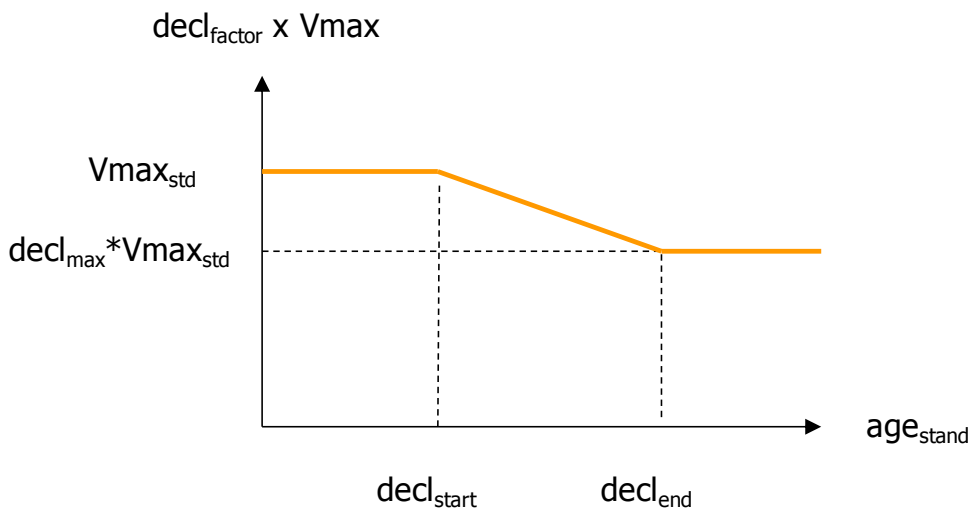


Figure A 1 illustrates this process.

25.2 *Initial distribution*

25.2.1 Algorithm

The initial tree circumference distribution follows a truncated exponential law of parameter λ resulting from the following algorithm:

- A minimum initial circumference, circ_init_{\min} , is selected.
 - The exponential distribution is truncated so that unlikely values do not appear.
- circ_init_{\max} , the maximum initial circumference, is selected so that:

$$P(X < \text{circ_init}_{\max}) = 1 - p_{\max} \text{ with } p_{\max} \text{ set at } \frac{100}{n_{\max\text{trees}}} \quad (\text{A1})$$

- The $[\text{circ_init}_{\min}, \text{circ_init}_{\max}]$ interval is divided into 20 intervals.
- The number of trees n_i in each interval $[a, b]$ is proportional to $P(a \leq X < b)$:

$$n_i = (\exp(-\lambda b) - \exp(-\lambda a)) \times n_{\max\text{trees}} \quad (\text{A2})$$

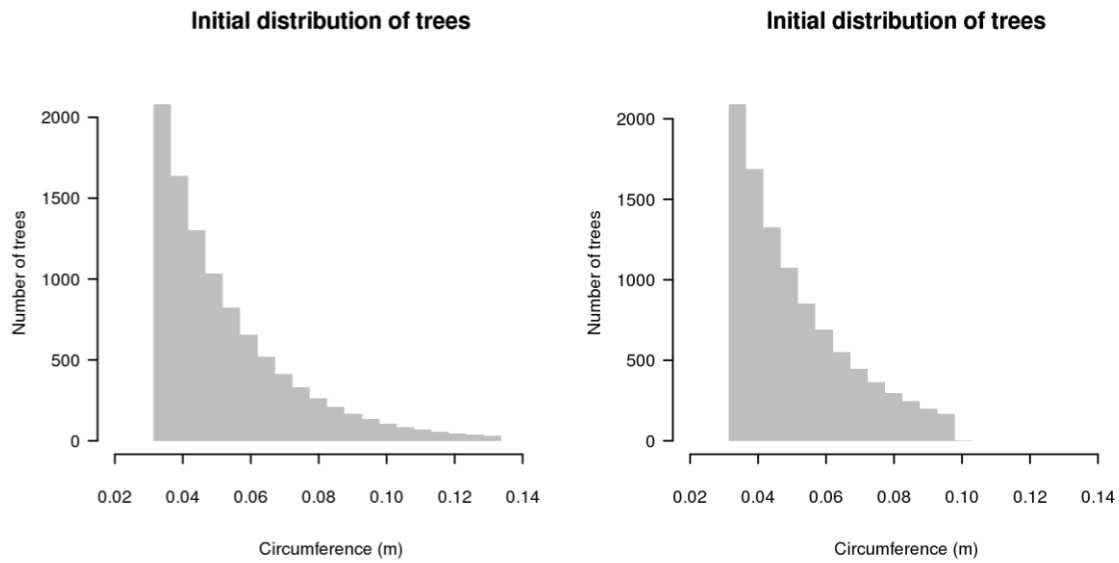
- These numbers are rounded to the closest integer, and the number of trees of each intervals are adjusted so that:

$$\sum_i n_i = n_{\max\text{trees}} \quad (\text{A3})$$

- Tree circumferences are then equally distributed in each interval:

$$\forall j \in n_i \text{ circ}_j = a + b \times \frac{j}{n_i} \quad (\text{A4})$$

The resulting distribution is illustrated in



a. $p_{\max} = 100/n_{\text{maxtrees}}$

b. $p_{\max} = 500/n_{\text{maxtrees}}$

Figure A 2.

25.2.2 Impact of a more condensed initial distribution

If p_{\max} is increased 5-fold, circ_init_{\max} is decreased by 27%, leading to a more condensed distribution (see

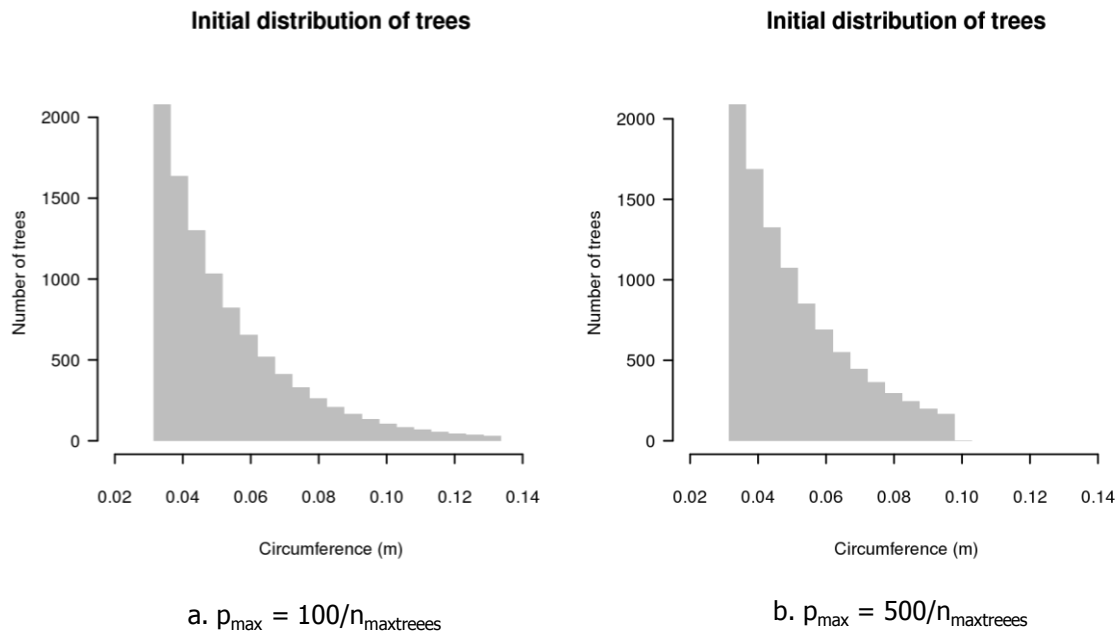
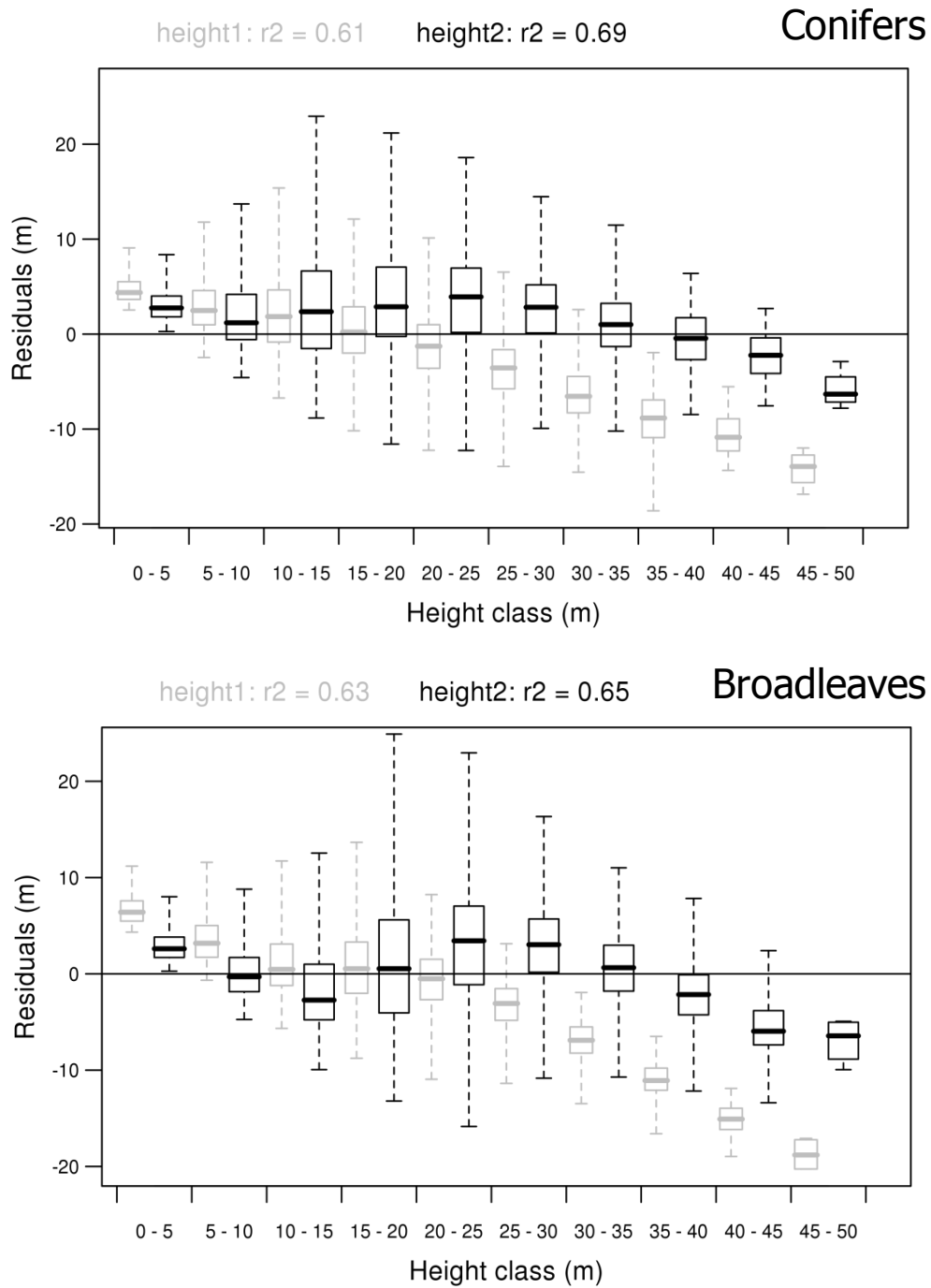


Figure A 2b).

25.3 Calibration of the height-circumference allometry

To calibrate the height-circumference allometry, we restricted the national inventory data set (IFN, 2008) along the following stand criteria: high forests, dedicated to wood production, with a known tree density, a closed canopy, and a basal area greater than $10 \text{ m}^2 \text{ ha}^{-1}$. Broadleaf and needleleaf stands were fitted separately. The Gauss-Newton non-linear algorithm was then used to fit the allometry.



690

691 Figure A 3 shows that the residuals of the allometric model used in the FMM, “height model 2”,

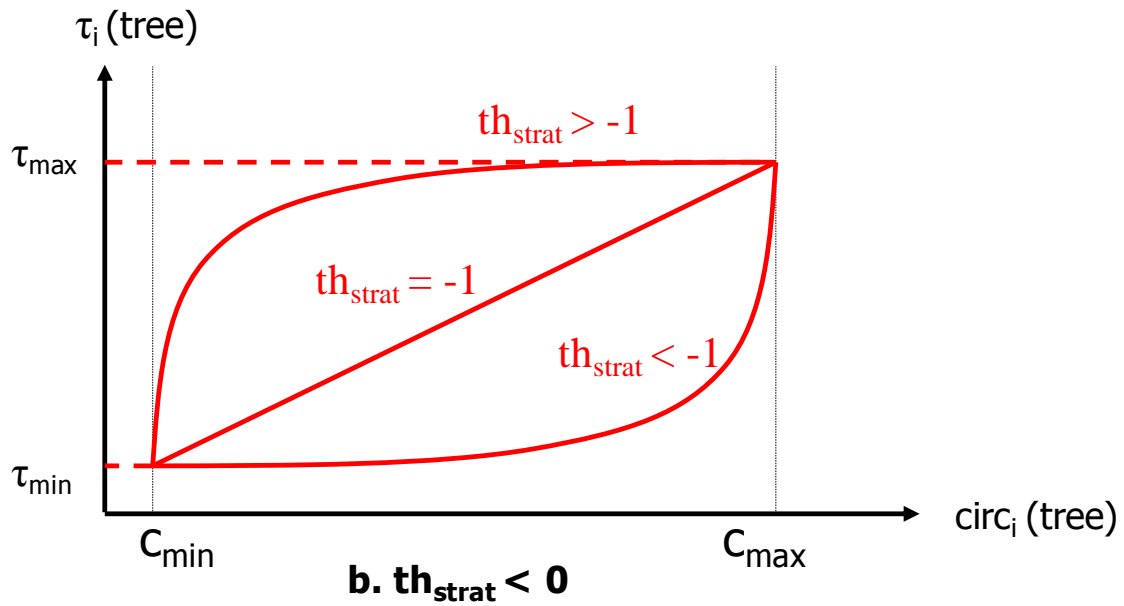
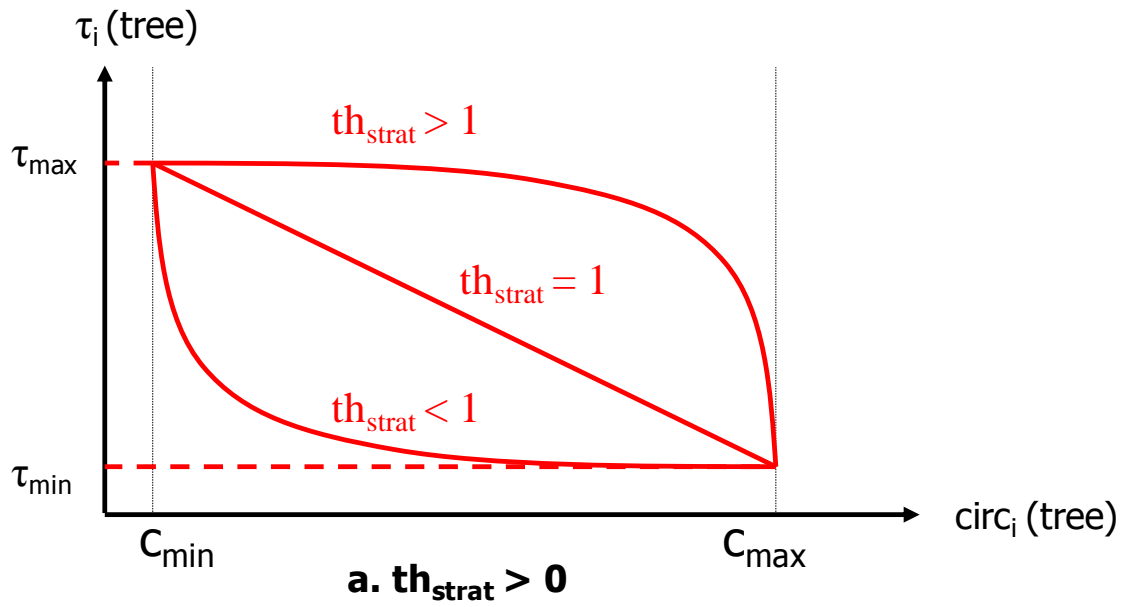
692 are less biased for large trees than those of a simpler model, “height model 1” (Eq. (A5)).

$$\text{height}_i = 1.3 + \alpha \times \left(1 - \exp(-\delta \times \text{circ}_i)^\varphi\right) \quad (\text{A5})$$

where height_i and circ_i are respectively the height and circumference of tree i in m , and α , δ , and φ are parameters.

25.4 Thinning strategy (th_{strat})

In order to determine which trees are felled during a thinning event (be it natural or anthropogenic), a probability of death τ_i is attributed to each tree Eq. (A4). The value of the parameter th_{strat} sets the thinning strategy: if $th_{strat} > 0$, a “thinning from below” strategy is simulated, with smaller trees preferentially thinned to obtain larger logs in the future. If $th_{strat} < 0$, a “thinning from above” strategy is simulated, with larger trees preferentially thinned thus giving way to smaller trees.

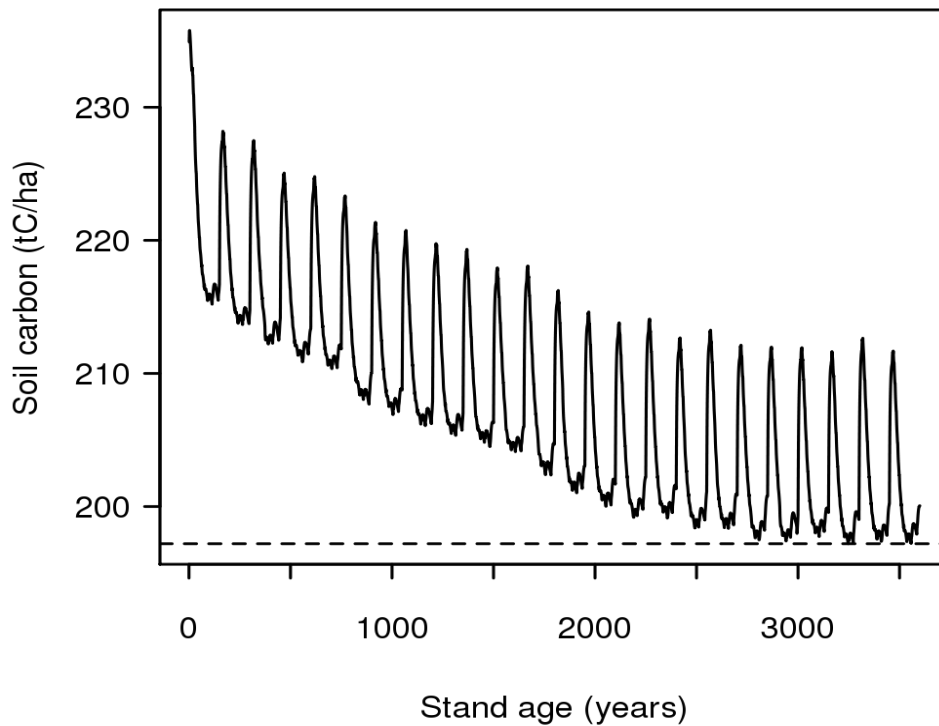


703

704 Figure A 4 illustrates this algorithm.

25.5 *Initial conditions (“spinup”)*

As computing time is increased when ORCHIDEE is coupled to the FMM, the “spinup” is performed in two steps. First, ORCHIDEE without the FMM is repeatedly run for the 1997 climate and a CO₂ concentration of 380 ppm until all ecosystem carbon and water pools reach their steady state equilibrium. Using this first steady state as initial conditions, ORCHIDEE is then run with the FMM for seventeen rotations (that is 2550 years), using the same climatic conditions. After seventeen rotations, the soil carbon pool reaches a new cyclic steady state (see



713

714 Figure A 5). The conditions of the stand before the last clearcut are used as initial conditions for
 715 all subsequent simulations.

716 **25.6** *Evolution of tree circumference distribution over a forest rotation*

717 The simulated evolution of tree circumference distribution over a forest rotation is illustrated
 718 by

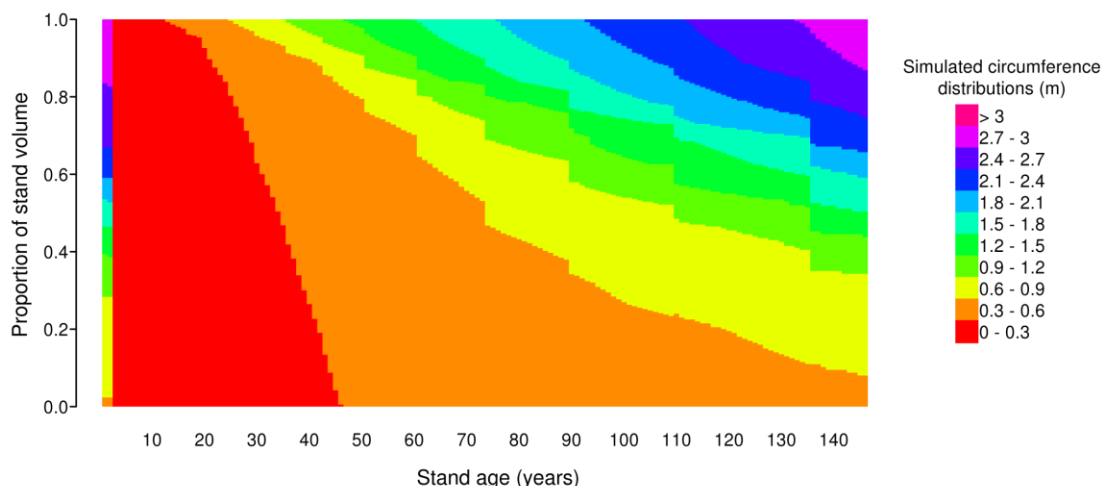


Figure A 6.

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968 **Tables**969 **Table 1. Parameters names and their default values**

Name	Description	Value for broadleaves	Value for coniferous	Unit	Sources	Equation	Section
$decl_{max}$	Maximum age-related decline in photosynthesis efficiency	0.95	0.90	no unit	Teobaldelli et al. 2008, Gower et al., 1996	1	2.2.2
$decl_{start}$	Age at which age-related decline of NPP starts	50	same	year	Gower et al., 1996, Magnani et al., 2000	1	2.2.2
$branch_{ratio}$	Ratio of branches over total aboveground biomass	0.38	0.25	no unit	Loustau 2004, Le Maire 2005	na	2.2.2
$branch_{turn}$	Proportion of branches dying each day	2.5	same	%·year ⁻¹	Masera 2003, van Oene 2000	na	2.2.2
$branch_{sap/heart}$	Sapwood/Heartwood ratio in branches	0.5	same	no unit	Hoffmann 1995	na	2.2.2
τ_{wd}	Maximum turnover rate of coarse woody debris	0.75	same	year ⁻¹	Olsson 1996, Schelaas 2004, Nagy 2006	na	2.2.2
$alloc_{min}$	Minimum aboveground/belowground sapwood allocation ratio	0.60	same	no unit	Mokany 2004, Nagy 2006	3	2.2.2
$alloc_{max}$	Maximum aboveground/belowground sapwood allocation ratio	0.80	same	no unit	Mokany 2004, Nagy 2006	3	2.2.2
$demi_{alloc}$	Half-life of aboveground/belowground sapwood allocation ratio increase	5.00	same	year	Mokany 2004, Nagy 2006	3	2.2.2
$n_{maxtrees}$	Initial stand density	10 000	same	ind·ha ⁻¹	Dhôte 2003, van Oene 2000	na	2.3.2
Dg_{init}	Initial quadratic mean diameter	0.01	same	m	Dhôte 2003	4	2.3.2
m	Smoothing parameter for tree growth equation (growth=f(circumference))	1.05	same	no unit	Deleuze 2003	5	2.3.2
$\rho_{density}$	Wood density	0.3	0.2	tC·m ⁻³	Hoffmann 1995, Friend 1997, FCBA 2009	na	2.3.2
a_e	Slope of the linear regression $\ln(\sigma)=f(\ln(dens))$	-0.35	same	$\ln(m) \cdot \ln(ind \cdot ha)^{-1}$	Fitted on data from Dhôte 2000	6	2.3.2
b_e	Intercept of the linear regression $\ln(\sigma)=f(\ln(dens))$	1.88	same	$\ln(m)$	Fitted on data from Dhôte 2000	6	2.3.2
a_{bc}	Coefficient of biomass-circumference allometry	$7.03 \cdot b_{bc}^{-4.76}$	same	kgDM	Zianis 2004	7	2.3.2
b_{bc}	Coefficient of biomass-circumference allometry	2.44	2.30	$\ln(kgDM) \cdot \ln(m)^{-1}$	Fitted on data from IFN 2008	7	2.3.2
α_{st}	Coefficient of self-thinning equation	min(171 582-145 248)	198 336	ind·ha ⁻¹	Dhôte 2003, Vacchiato 2008	8	2.3.3
β_{st}	Coefficient of self-thinning equation	min(1.7-1.57)	1.60	$\ln(ind \cdot ha^{-1}) \cdot \ln(m)^{-1}$	Dhôte 2003, Vacchiato 2008	8	2.3.3
α	Coefficient of circumference-height allometry	19.42	9.30	na	Fitted on data from IFN 2008	10	2.3.3
χ	Coefficient of circumference-height allometry	0.11	0.35	na	Fitted on data from IFN 2008	10	2.3.3
δ	Coefficient of circumference-height allometry	0.13	0.13	na	Fitted on data from IFN 2008	10	2.3.3
φ	Coefficient of circumference-height allometry	0.75	0.69	na	Fitted on data from IFN 2008	10	2.3.3
Φ	Coefficient of circumference-height allometry	-0.12	-0.32	na	Fitted on data from IFN 2008	10	2.3.3
rdi_{target}	Targeted value of relative density index	0.75	same	no unit	Cazin 2003	na	2.2.3
rdi_{lim}	Width of buffer within which rdi is allowed to vary between thinnings	0.05-0.1*	same	no unit	Cazin 2003	na	2.2.3
$dens_{target}$	Target density triggering a clearcut	200	100	ind·ha ⁻¹	Lanier 1994	na	2.3.4
age_{target}	Target age triggering a clearcut	150	same	years	Lanier 1994	na	2.3.4
th_{strat}	Thinning strategy index	1	same	no unit	Dhôte 2008	12	2.3.4
τ_{min}	Minimum relative mortality rate	0.01	same	no unit	na	12	2.3.4
τ_{max}	Maximum relative mortality rate	0.05	same	no unit	na	12	2.3.4
δlai_{max}	Proportional decrease of lai_{max} after thinning	30	same	%	Le Dantec 2000, Vesala 2005	na	2.4

*0.1 when density is $n_{maxtrees}$ log-linearly decreasing to 0.05 when density is denstarget

age (years)	20	40	60	80	100	120	140
density (ind/ha)	4095	1442	684	423	279	203	153
basal area (m²/ha)	18.8	24.1	26.7	29.9	31.8	33.7	35.2
average height (m)	8.0	12.2	15.6	18.4	20.7	22.5	24.0
stand volume (m³/ha)	108	187	258	328	388	441	489
exported volume / total volume ratio	0.38	0.5	0.54	0.55	0.55	0.56	0.57
thinning frequency (years)	4	8	10	16	21	27	27
average circumference (m)	0.24	0.43	0.65	0.86	1.09	1.32	1.54
minimum circumference (m)	0.18	0.29	0.36	0.43	0.49	0.54	0.58
maximum circumference (m)	0.56	1.12	1.56	1.95	2.29	2.58	2.83

Table 2. Stand characteristics at different ages in the “managed” simulation (ORCH-FM_m)

Thinning frequency is defined as the time between the two thinnings surrounding the corresponding age. Exported volume and total volume produced both refer to total wood (including branches and stem parts with diameter lower than 7 cm).

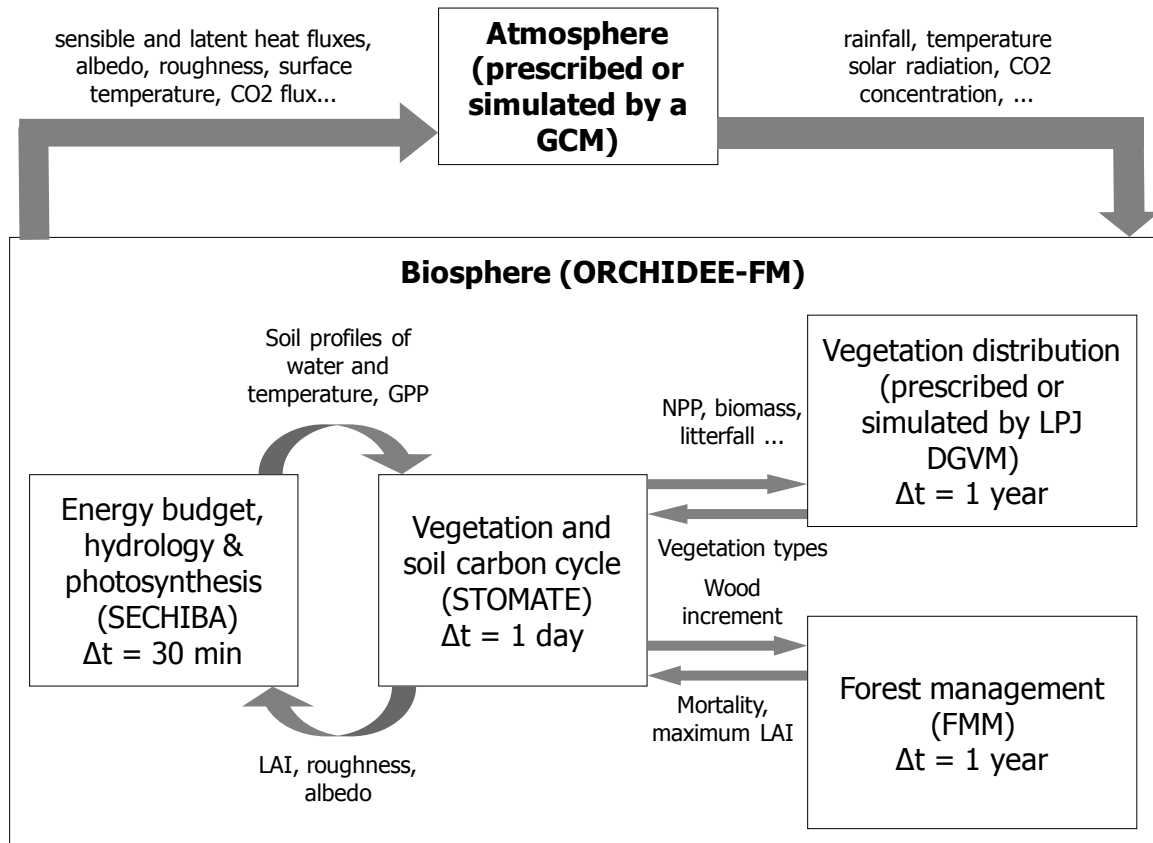
Parameters			Variables	
Symbol	Full name	Equation number	Symbol	Full name
<i>p_max</i>	Probability threshold for truncating exponential distribution	A1	<i>dens</i>	Tree density
<i>min_circ_init</i>	Minimum circumference in initial distribution	A1	<i>ba</i>	Stand basal area
<i>dens_init</i>	Initial density	na	<i>av_height</i>	Average height
<i>lambda</i>	λ parameter of initial exponential distribution	2	<i>stand_vol</i>	Standing volume
<i>height_circ</i>	Height/circumference allometry. A value greater than 1 indicated a greater height for the same circumference.	9	<i>vol_exp / vol_tot</i>	Exported volume / Total volume produce ratio
<i>circ_bm</i>	Circumference/biomass allometry. A value greater than 1 indicated a greater circumference for the same biomass	5	<i>th_int</i>	Time interval between two thinnings
<i>wood_density</i>	Wood density	na	<i>av_circ</i>	Average circumference
<i>branch_turn</i>	Branch turnover rate	na	<i>circ_min</i>	Minimum circumference
<i>branch_ratio</i>	Branch ratio	na	<i>circ_max</i>	Maximum circumference
<i>decl_max</i>	Maximum age-related decline in NPP	na		
<i>tau_spread</i>	Range between maximum and minimum relative mortality rate (τ_i)	na		
<i>th_strat</i>	Thinning strategy	10		
<i>selfth_curve</i>	Self-thinning equation. A value greater than 1 indicates that a higher density is tolerated for the same quadratic mean diameter.	6		
<i>rdi_target</i>	Targeted value of relative density index	na		
<i>delta_rdi</i>	Bandwidth around rdi_target	na		
<i>sigma</i>	Threshold of the biomass distribution equation	3		

977

978 Table 3. Full name of the parameters and variables included in the sensitivity analysis

979

980 **Figure captions (main text)**



981

982 Figure 1. Structure of ORCHIDEE

983

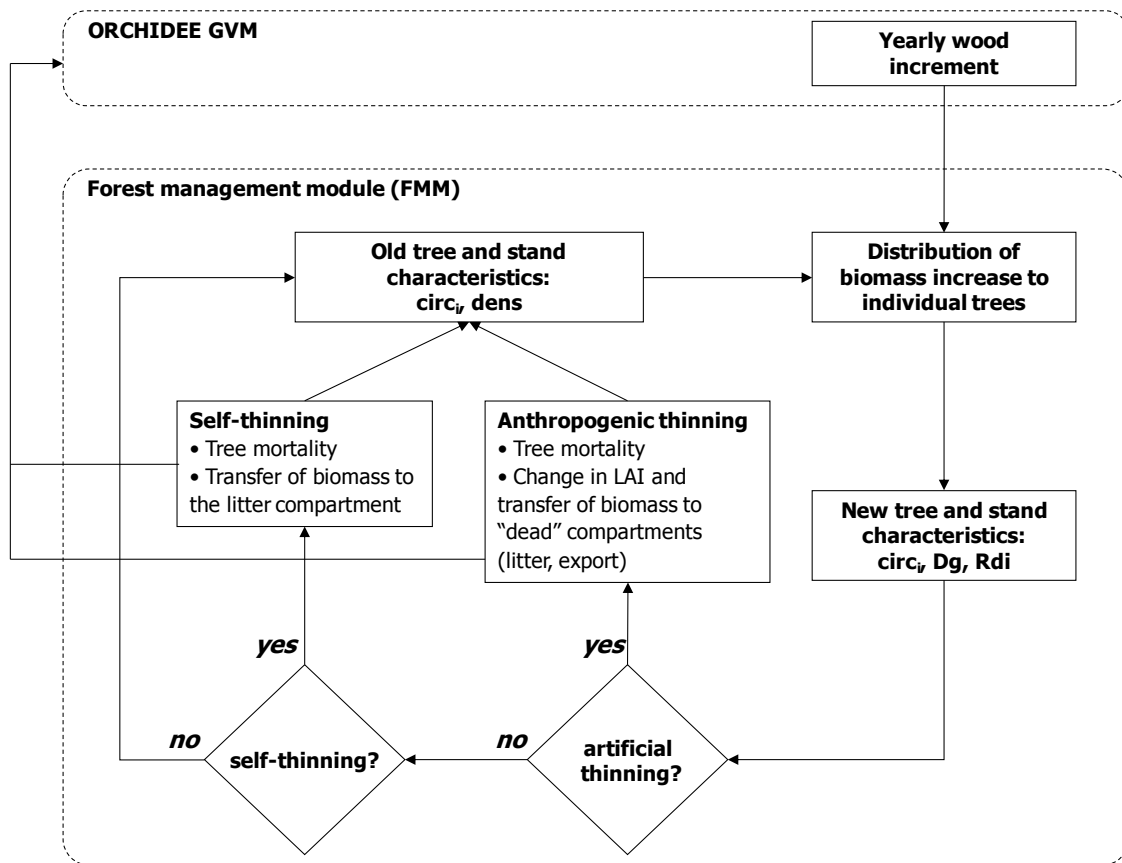


Figure 2. Conceptual diagram of the forest management module (FMM)

The FMM calculates mortality by explicitly simulating stand and tree characteristics: tree density (*dens*), the circumference of each tree (*circ_i*), quadratic mean diameter (*Dg*), relative density index (*rdi*), etc.

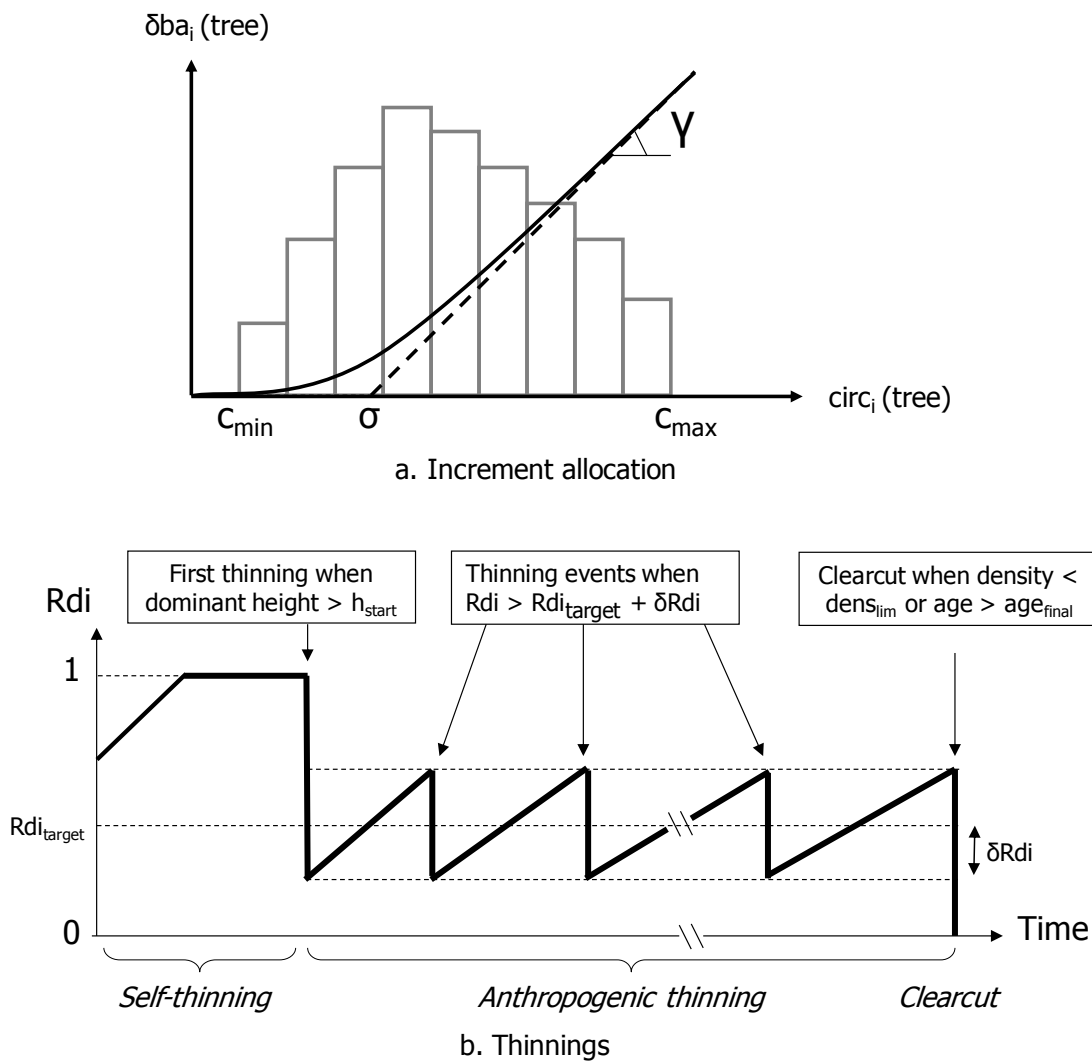


Figure 3. Increment allocation and thinnings in the FMM

- a. Increment allocation: σ is the threshold circumference for growth (the basal area increase of trees smaller than σ is close to 0) and γ is the slope of the relationship between increase in basal area and circumference. δba_i and $circ_i$ are respectively the basal area increase and circumference of tree i , and c_{min} and c_{max} are the minimum and maximum circumferences found in the plot. Larger trees get a bigger share of stand growth, and thus get a bigger increase in basal area.

b. Thinnings: the thick black line represents the evolution of rdi with time for a typical forest stand. In younger stands, self-thinning occurs to maintain the stand at its maximal carrying capacity ($rdi = 1$). Then, after a minimal height h_{start} is reached, human intervention maintains rdi around rdi_{target} . The stand is harvested when its density gets below $dens_{target}$ or its age reaches age_{final} .

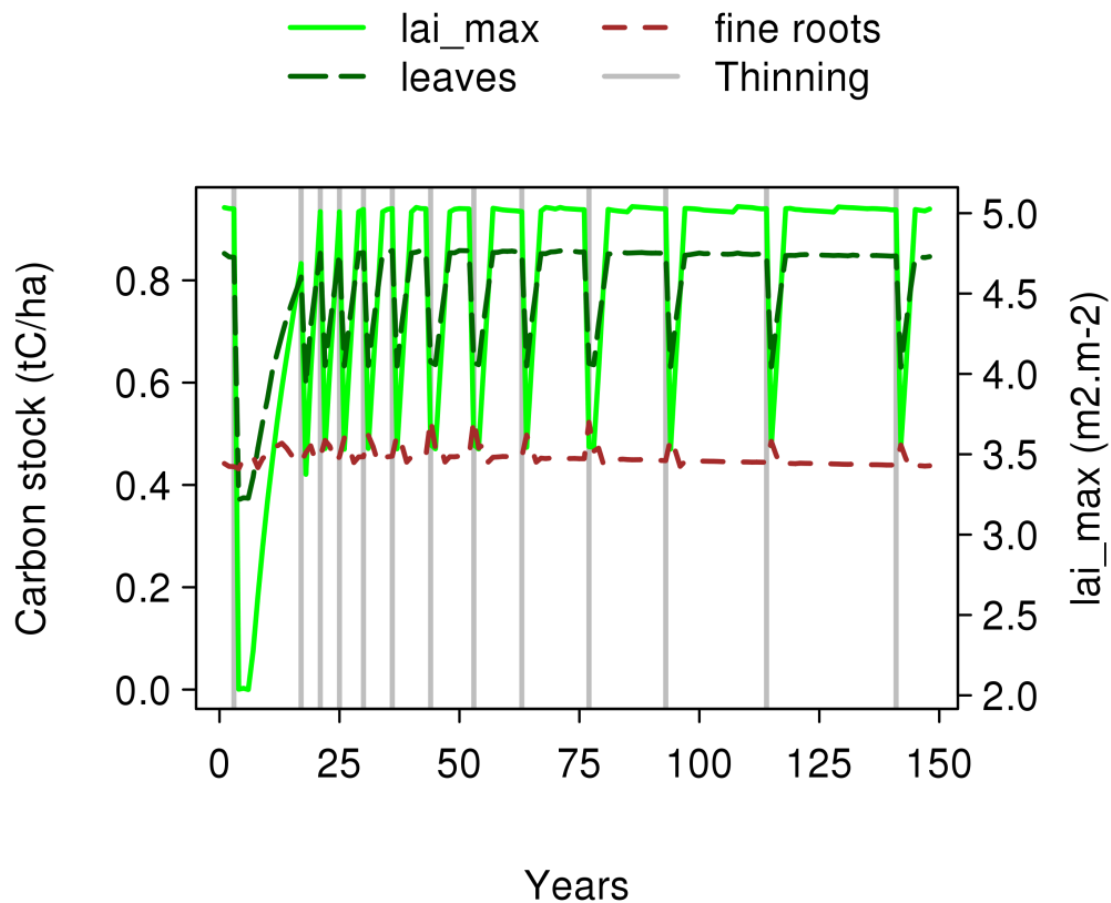
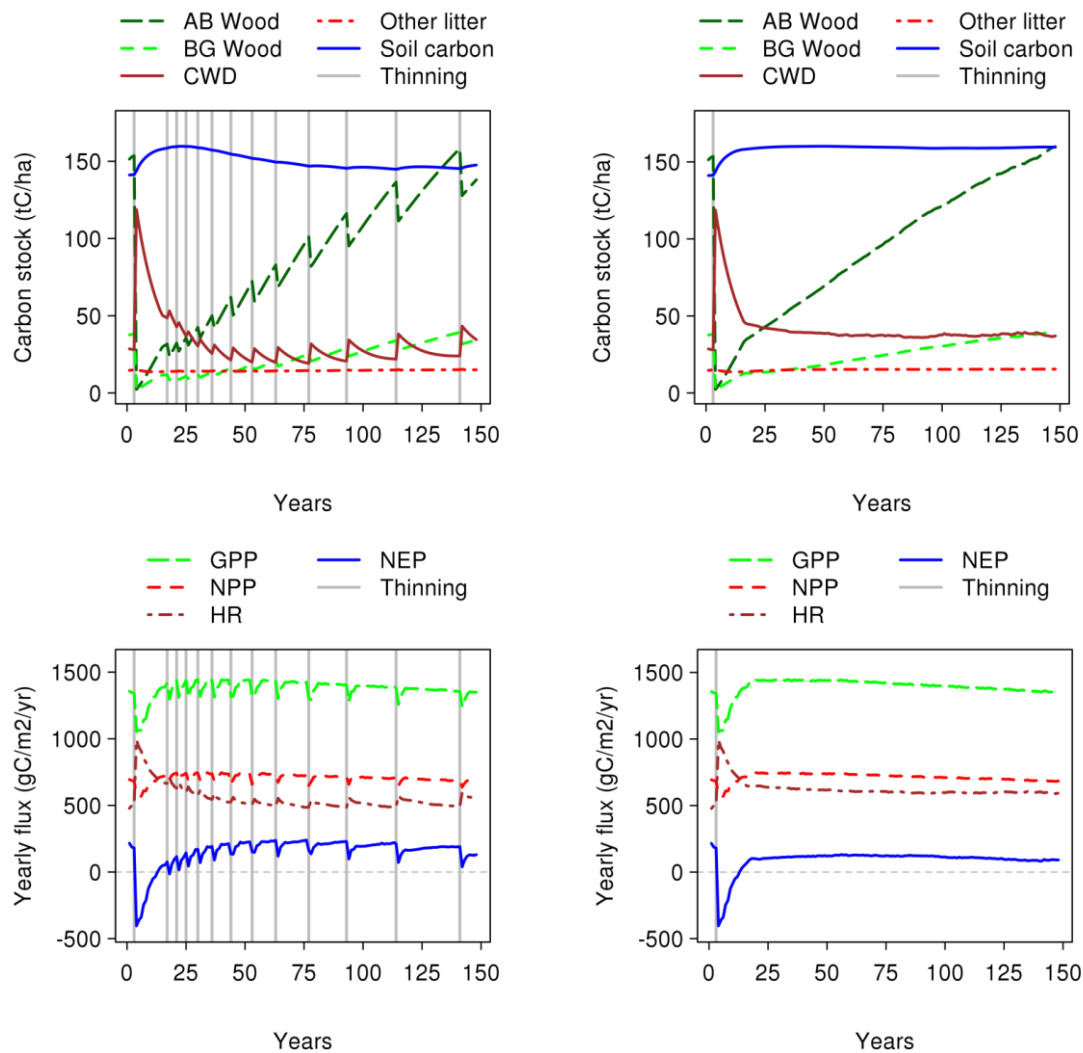


Figure 4. Lai_{max} and labile carbon stocks during a rotation period

Yearly maximum leaf area index (right axis) and yearly average carbon stocks in leaves and fine roots (left axis).

1008

a. "Managed" case (ORCH-FM_m)b. "Unmanaged" case (ORCH-FM_u)

1009

1010 Figure 5. Simulated carbon stocks and fluxes during a rotation period

1011 Yearly average in aboveground wood (AB wood), belowground wood (BG wood), coarse
 1012 woody debris (CWD), other litter (dead leaves) and soil carbon (top). Gross primary
 1013 production (GPP), net primary production (NPP), heterotrophic respiration (HR), and net
 1014 ecosystem productivity (bottom).

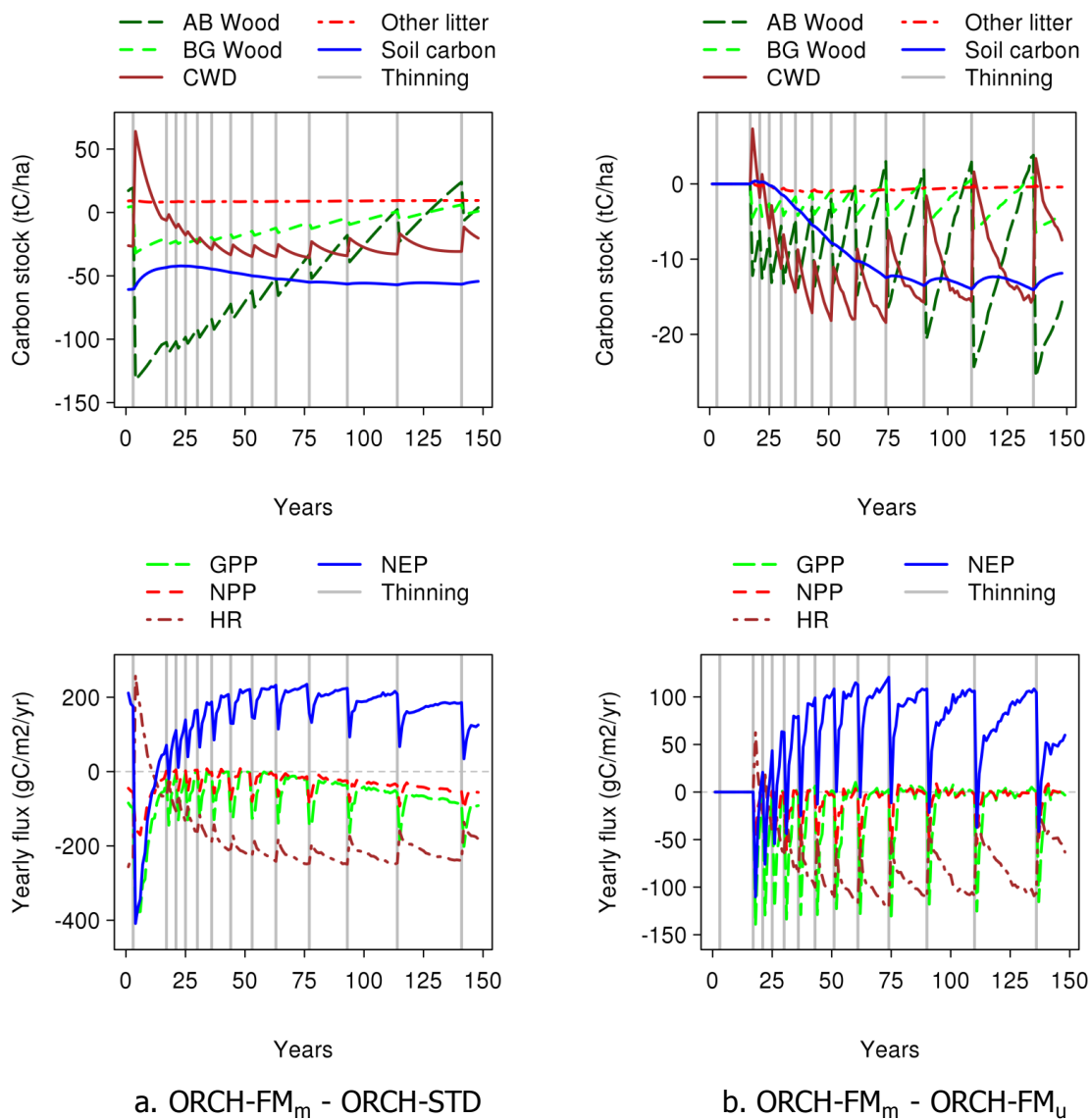
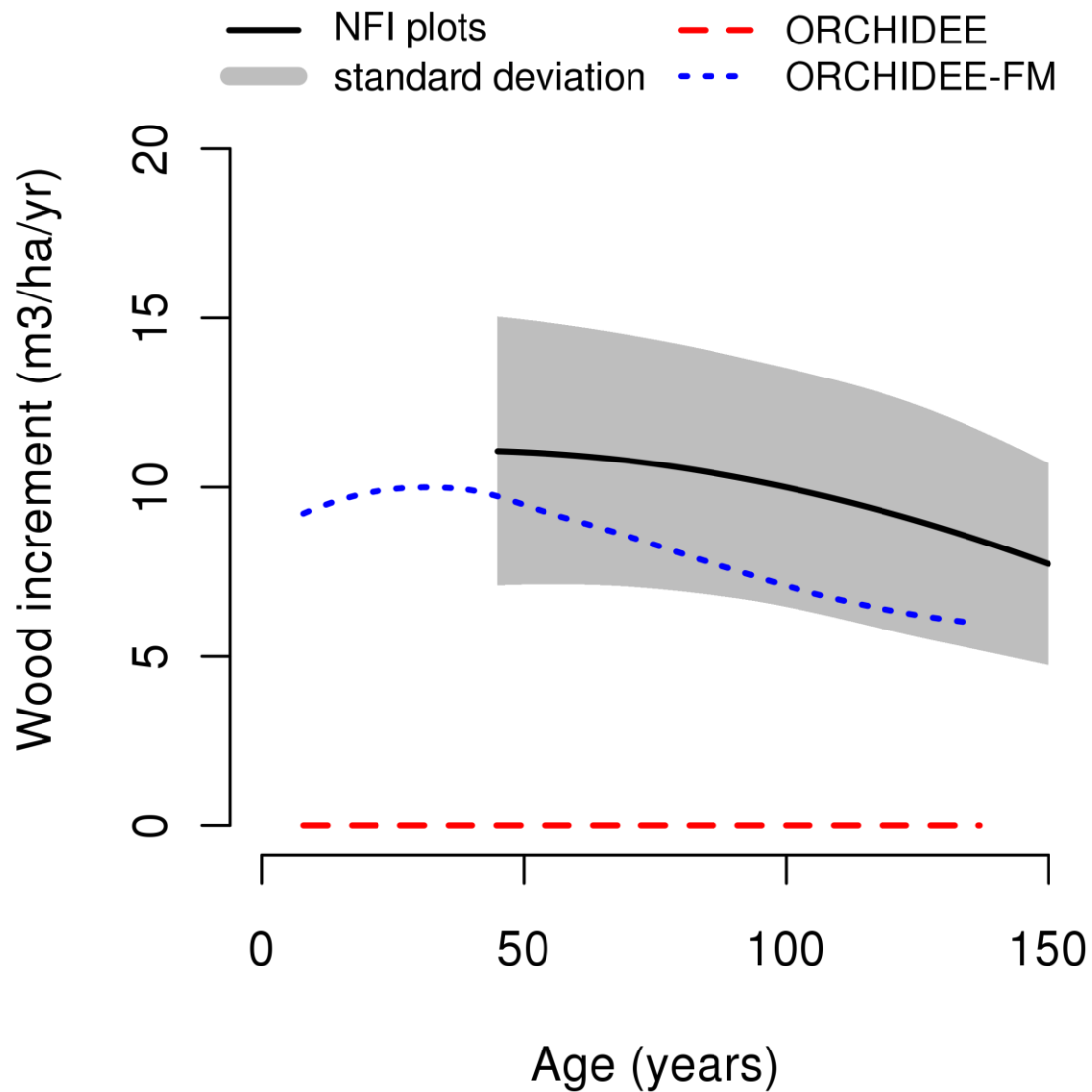


Figure 6. Comparison of ORCHIDEE-FM and ORCHIDEE.

The plotted curves represents the difference between the same variable simulated by (a) ORCHIDEE-FM “managed” (ORCH-FM_m) and ORCHIDEE (ORCH-STD) and by (b) ORCHIDEE-FM “managed” (ORCH-FM_m) and ORCHIDEE-FM “unmanaged” (ORCH-FM_u).

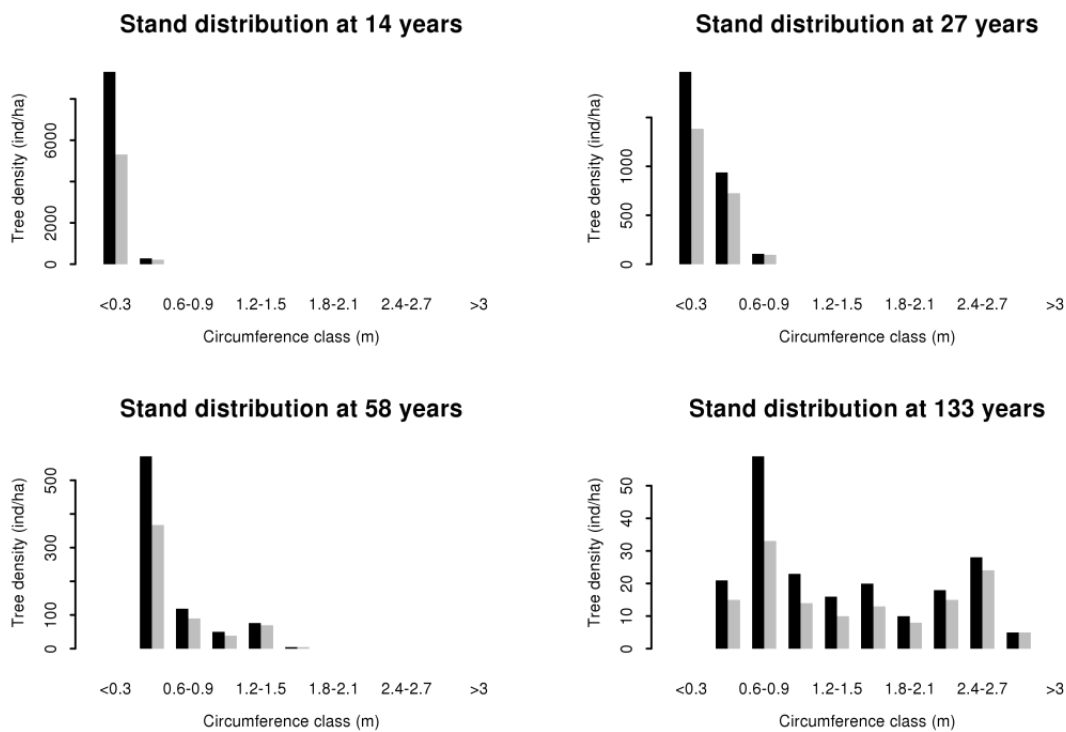
1021 Variables: yearly average in aboveground wood (AB wood), belowground wood (BG
 1022 wood), coarse woody debris (CWD), other litter (dead leaves) and soil carbon (top).
 1023 Gross primary production (GPP), net primary production (NPP), heterotrophic
 1024 respiration (HR), and net ecosystem productivity (bottom).
 1025



1026

1027 Figure 7. Simulated and observed wood increment close to Nancy

1028 The black solid line and grey area respectively give the average and standard deviation
 1029 of measured wood increment in National Forest Inventory (NFI) plots within a 50 km
 1030 radius of our selected grid cell. Measurements are pooled per age class, and the
 1031 resulting statistics per age class are smoothed using a “loess” algorithm (only age classes
 1032 with 5 or more plots are retained). The large-dashed red curve and the small-dashed
 1033 blue curve respectively give the wood increment in the ORCH-STD and ORCH-FM_m
 1034 simulations.
 1035



1036
 1037 Figure 8. Tree distribution by circumference classes
 1038 Represented before (black bars) and after (grey bars) thinning for 4 selected thinning
 1039 events.

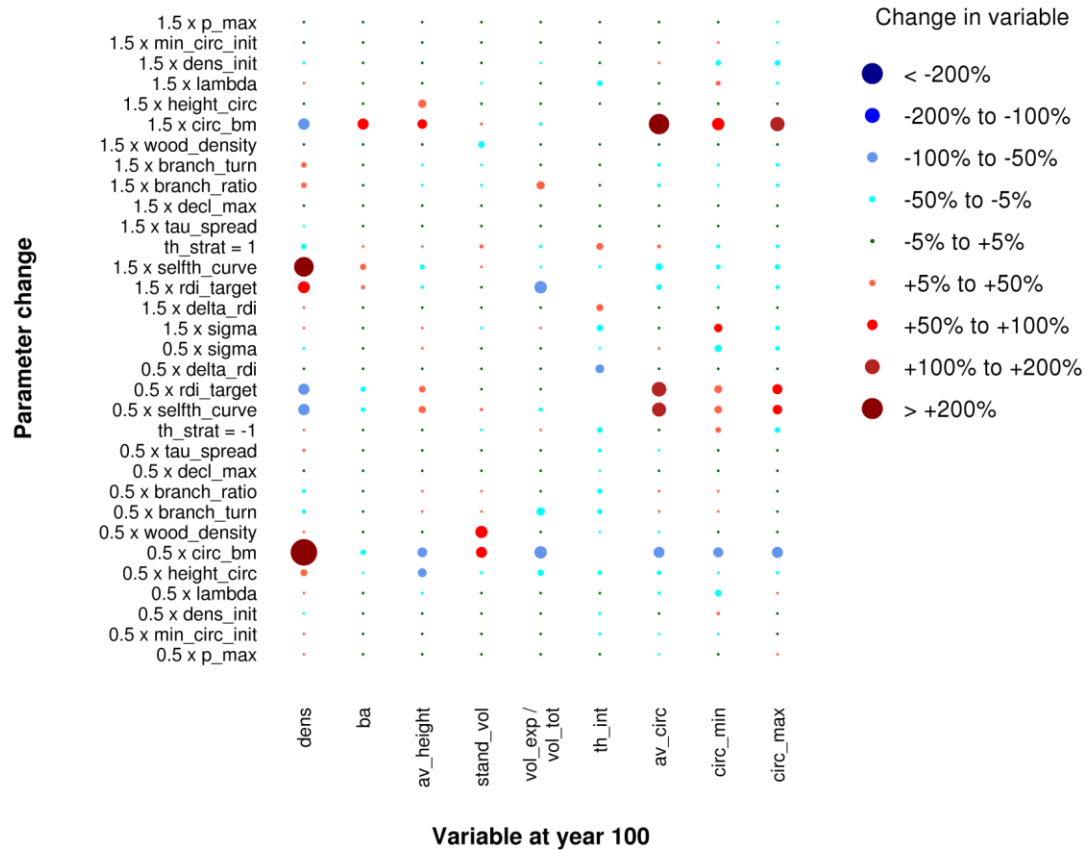
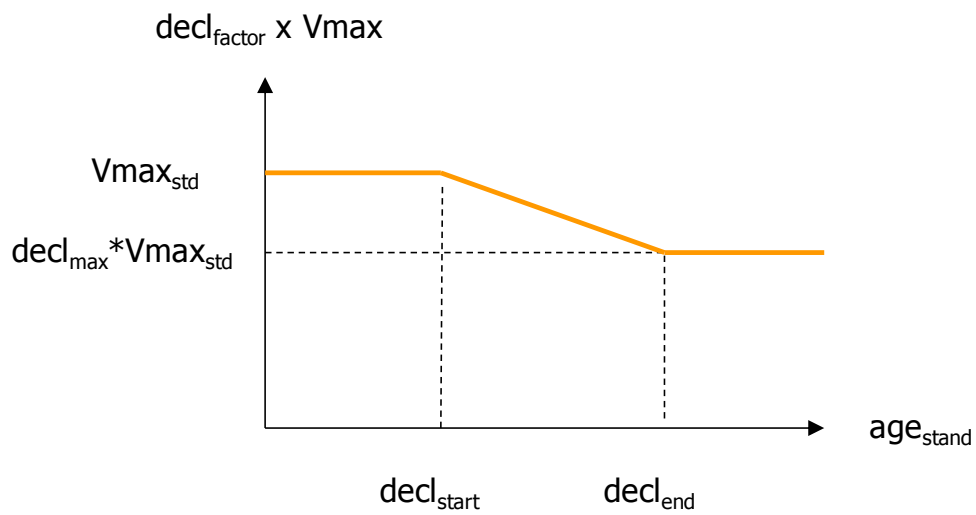


Figure 9. Sensitivity of 9 selected variables to changes in the values of 15 selected parameters

The ordinates axis indicates by how much the default parameter value is multiplied (eg. $1.5 \times \lambda$ indicates a model run with a λ increased by 50% compared to its default value given in Table 3). The impact of this parameter change on the selected variables is represented by a full circle. The area of the circle is proportional to the absolute value of the change in the selected variable. Blue circle represent decreases and red circle represent increases. Empty values correspond to infinite changes (eg.

1050 when there is no thinning between year 100 and the end of the rotation, the thinning
1051 frequency is infinite). Table 3 lists the full names of these variables and parameters.
1052 Note that parameters are classified according to their “role” in the model (grey and
1053 white highlighting).
1054

1055 **Figure captions (appendixes)**



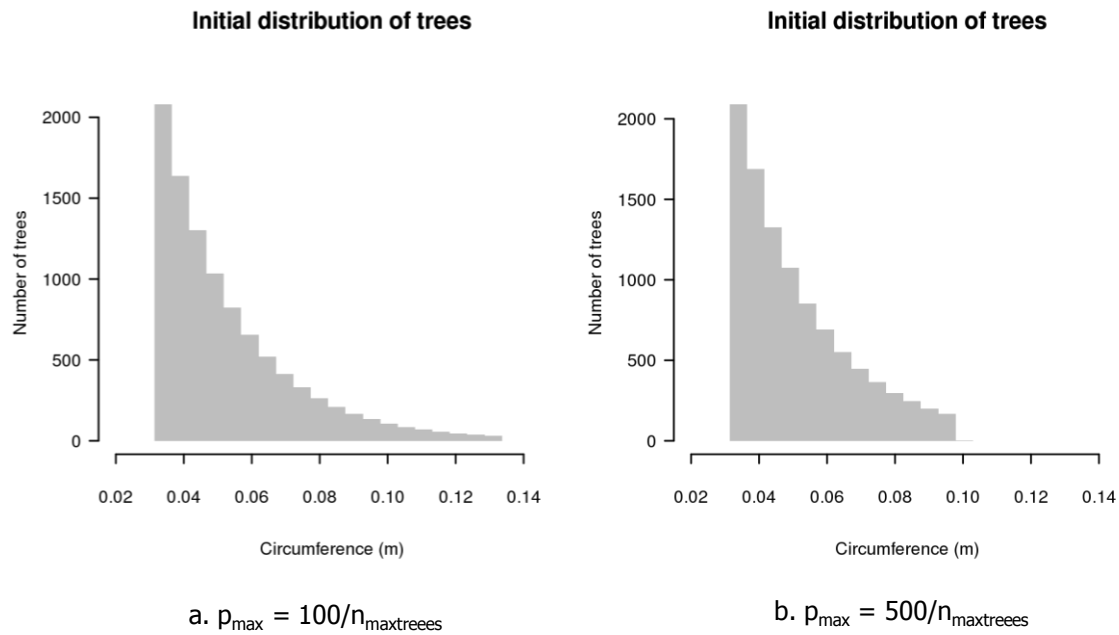
1056
1057 Figure A 1. Age-related decline of photosynthesis efficiency

1058 V_{max} is the photosynthesis efficiency, $V_{\text{max}_{\text{std}}}$ is the standard value of V_{max} in

1059 ORCHIDEE, decl is the maximum age-related decline, $\text{decl}_{\text{start}}$ and decl_{end} are respectively

1060 the ages at which age-related decline starts and saturates.

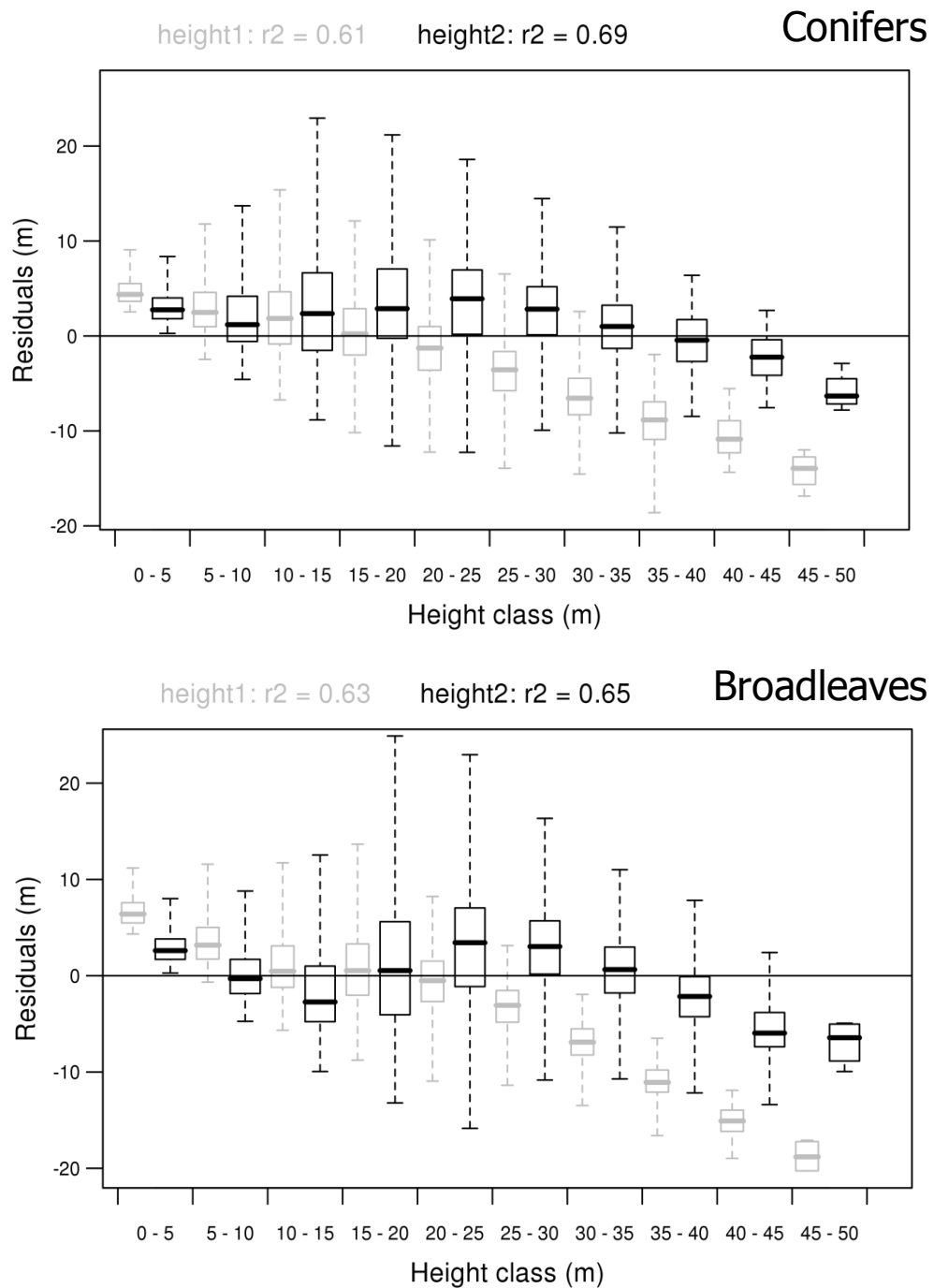
1061



1062

1063 Figure A 2. Two examples of initial distributions for the same tree density ($n_{\max\text{trees}} =$
 1064 10 000 stems per hectare): the default distribution (a) and a more condensed possibility
 1065 (b). p_{\max} is the probability threshold at which the distribution is truncated.

1066



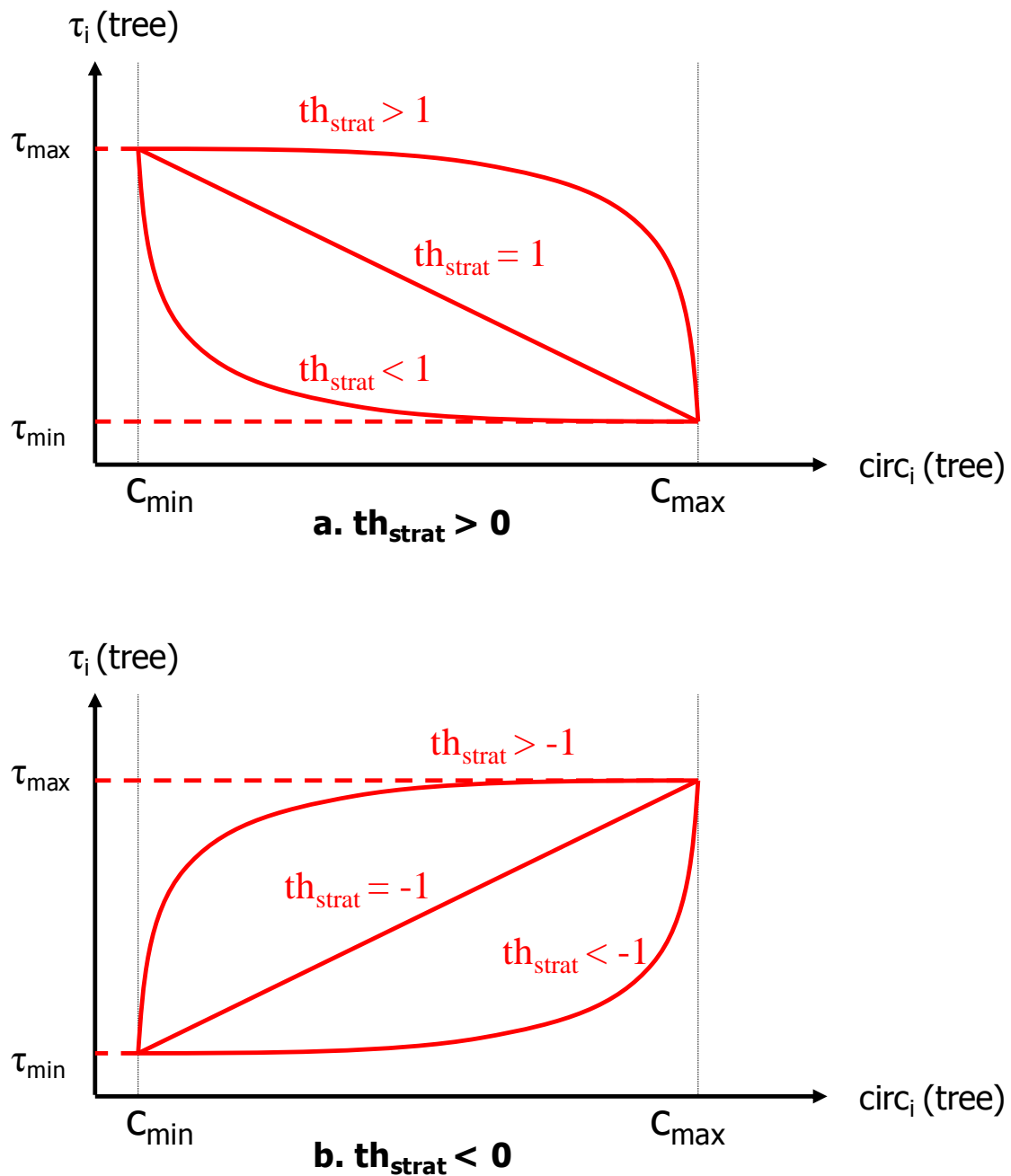
1067

1068 Figure A 3. Residuals (model – data) of the height-diameter allometry used in the FMM

1069 (height model 2) compared to a simpler allometry (height model 1) for conifers (top)

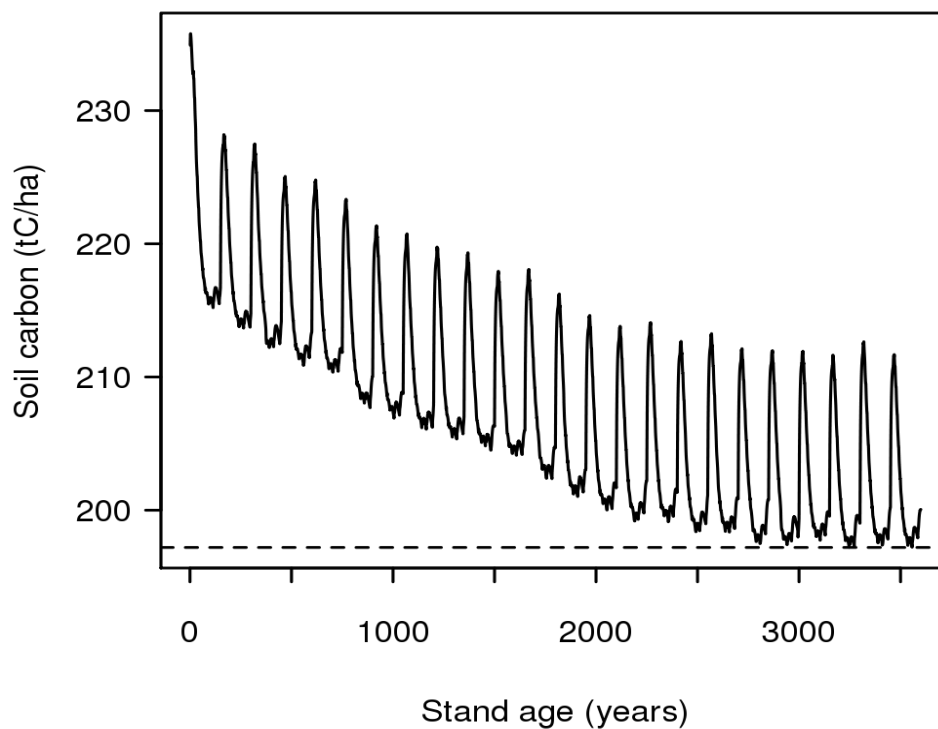
1070 and broadleaves (bottom)

1071 The whisker-plots show the median, first and third quartile, and the minimum and
 1072 maximum within a range of twice the inter-quartile value.
 1073

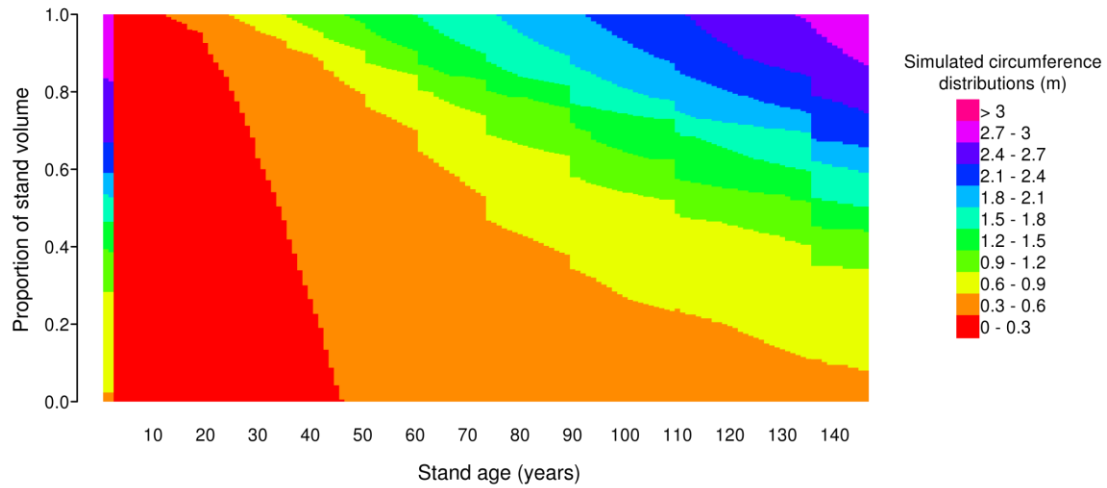


1074
 1075 Figure A 4. Thinning strategies as a function of th_{strat}

1076 $circ_i$ is the circumference of tree i , c_{min} and c_{max} are respectively the minimum and
 1077 maximum tree circumference in the stand, and τ_i is the probability of death of tree i and
 1078 τ_{min} and τ_{max} are respectively the minimum and maximum probabilities of death in the
 1079 stand. For th_{strat} , see Eq. 12.
 1080



1081
 1082 Figure A 5. Long-term soil carbon equilibrium
 1083



1084

1085 Figure A 6. Evolution of circumference distribution over one forest rotation

1086 One bar represents the simulated distribution of total stand volume between different

1087 tree circumference classes for a given stand age.