

Article

The Role of Respiration in Estimation of Net Carbon Cycle: Coupling Soil Carbon Dynamics and Canopy Turnover in a Novel Version of 3D-CMCC Forest Ecosystem Model

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Abstract: Understanding the dynamics of organic carbon mineralization is fundamental in forecasting biosphere to atmosphere net carbon ecosystem exchange (NEE). With this perspective, we developed 3D-CMCC-PSM, a new version of the hybrid process based model 3D-CMCC FEM where also heterotrophic respiration (R_h) is explicitly simulated. The aim was to quantify NEE as a forward problem, by subtracting ecosystem respiration (R_{eco}) to gross primary productivity (GPP). To do so, we developed a simplification of the soil carbon dynamics routine proposed in the DNDC (DeNitrification-DeComposition) computer simulation model. The method calculates decomposition as a function of soil moisture, temperature, state of the organic compartments, and relative abundance of microbial pools. Given the pulse dynamics of soil respiration, we introduced modifications in some of the principal constitutive relations involved in phenology and littering sub-routines. We quantified the model structure-related uncertainty in NEE, by running our training simulations over 1000 random parameter-sets extracted from parameter distributions expected from literature. 3D-CMCC-PSM predictability was tested on independent time series for 6 Fluxnet sites. The model resulted in daily and monthly estimations highly consistent with the observed time series. It showed lower predictability in Mediterranean ecosystems, suggesting that it may need further improvements in addressing evapotranspiration and water dynamics.

Keywords: forest ecosystem; Fluxnet; soil respiration; net ecosystem exchange; phenology

1. Introduction

Global concerns over increasing levels of greenhouse gas concentrations, particularly carbon dioxide (CO_2), have pushed research efforts to better investigate biogeochemical carbon (C) flux dynamics and patterns between atmosphere and biosphere, and to upscale C flux estimates from

site-specific to regional, continental and global scales. Increased atmospheric concentration of CO₂, combined with increasing temperatures and size variations of ecosystem C pools, are responsible for year-to-year terrestrial ecosystem carbon flux perturbations, through the variation of both photosynthetic and respiration rates [1].

In the last decades, the eddy covariance (EC) technique has provided long-term continuous measurements of net carbon ecosystem exchange (NEE), water vapor and energy, within the global network of EC flux towers (FLUXNET) distributed over major terrestrial ecosystems. The availability of EC measure of NEE contributed to quantify and to determine seasonal and inter-annual variability of ecosystem C budgets at EC tower site-specific scale [2–4]. Observed NEE does not directly quantify the two major components of ecosystem C flux balance represented by ecosystem respiration (R_{eco}) and gross primary productivity (GPP). Thus, flux partitioning algorithms have been developed to partition eddy covariance NEE into photosynthetic uptake and respiratory release [5,6]. At the same time, EC flux measurements provide key information for the parameterization, calibration and validation of process-based forest ecosystem models (FEMs) contributing to large-scale estimates of main ecosystem C pools.

The implementation of both forest process-based models (PBMs) [7–12] and functional–structural tree models (FSTMs) [13–16], based on the widely used light use efficiency (LUE) approach [17], has contributed to understanding and upscaling the main physiological processes supporting ecosystem C uptake. Although most of forest ecosystem models provide reliable estimates of forest growth, limitations for NEE estimates are related to the uncertainty in R_{eco} estimation [18,19]. Hence, the implementation of biogeochemical models integrating soil respiration models and FSTMs is a great opportunity to reliably estimate NEE [20–22].

Although soil respiration and soil organic matter (SOM) decomposition depend mainly on abiotic factors, such as temperature and soil moisture [23–25], a key role is played by soil organic carbon (SOC) stock size, microbial pools [26,27], and dynamics of SOC supply to soil with littering [28]. Leaf and fine root senescence is of primary importance in determining dynamics in heterotrophic respiration [29,30], and exhibits seasonal patterns and intra-seasonal pulses [31,32]. These pulses are mostly driven by phenological transitions through stages of dormancy, active growth, and senescence. For example, supply of dead leaves in soil is strongly dependent on a tree's leaf-fall strategy. This is also true when predicting tree respiration, which is directly related to their growth and nitrogen content, which depends on spring phenology. Unfortunately, processes involving budburst and senescence are still partly obscure. PBMs usually represent these processes simplistically; these simplifications may lead some terrestrial ecosystem models to result in biased predictions [33–35].

For these reasons, we developed the 3D-CMCC-PSM (3D-CMCC-Phenology and soil Model), a new version of the hybrid process-based model 3D-CMCC FEM proposed by Collalti et al. [36,37], where (1) spring phenology was directly taking into account tradeoffs between growth and Non Structural carbon (NSC) demand; (2) phenological transitions and supply of Fresh organic matter (FOM) to soil were more explicitly represented; and (3) heterotrophic respiration (R_h) was explicitly simulated to dynamically quantify stock changes of 7 different SOC pools mediated by the amount of active microbial C pools, following the rationale proposed in the DNDC [38]. The aim of this study was to: (1) test the performance of the modified model version, comparing model NEE estimates against independent time series for 6 Fluxnet sites, representing different forests in different climatic areas, distributed over a wide latitudinal gradient amongst European EC sites; and (2) quantify uncertainty associated to 3D-CMCC-PSM constitutive relations structure and parameterization.

2. Materials and Methods

2.1. Study Area and Data

Eddy covariance data were collected from FLUXNET [39]. We chose the 6 sites to represent a climatic and longitudinal transect through Europe (Figure 1), so that the model could be tested on

different critical boundary conditions (Table 1). We used EC data from different time series from daily to annual, processed using the method described in [40]. Gross primary production (GPP) and ecosystem respiration (R_{eco}) were partitioned using [6]. Information about forest structure and total SOC at the beginning of the simulation was collected from literature (e.g., [41,42]) and PIs information. Sites were chosen to represent 3 diverse forest ecosystems, dominated by different species composition from deciduous broadleaved, DBF (i.e., *Fagus sylvatica* L.), evergreen broadleaved, EBF (i.e., *Quercus ilex* L.), and evergreen needle leaved, ENF (i.e., *Pinus sylvestris* L. and *Picea abies* L.), representing the most common European forest species from boreal to mediterranean ecoregions across Europe.

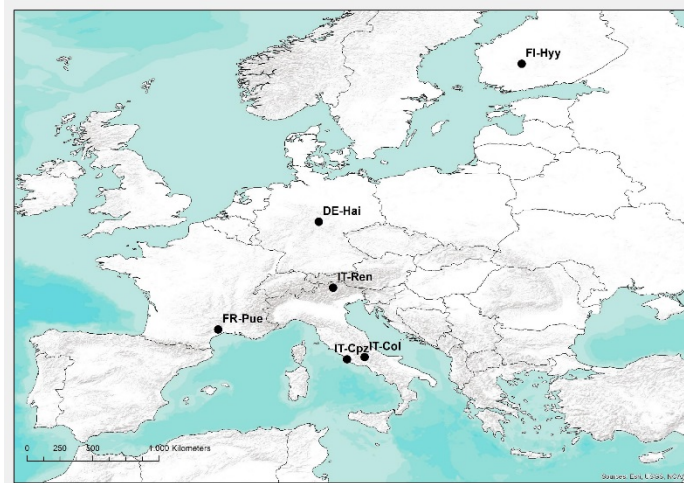


Figure 1. Location of the 6 Fluxnet sites used to evaluate 3D-CMCC-PSM.

2.2. Model Description

The 3D-CMCC FEM is a stand-scale process-based model (PBM) designed to simulate C and water cycle in natural and managed forest ecosystems (for a full description see [36,37]). Several eco-physiological processes were modeled at species-specific level, and at a variable temporal scale (from daily to annual) depending on the process to simulate (Figure 2). Model outputs were generally represented at hectare scale, while processes were simulated at different spatial scales from cellular (e.g., stomatal conductance), to canopy (e.g., transpiration), to individual tree, up to stand level.

Carbon assimilation was modeled for sun and shaded leaves using the LUE approach [43]: potential C assimilation was constrained by environmental and stand structural (e.g., tree age) scalars [44]. Autotrophic respiration (R_A) was explicitly modeled as the sum of growth and maintenance respiration (R_G and R_M , respectively). The first was computed as a fixed ratio of new growth tissues (30%) and the latter was based on nitrogen content in stems, branches, leaves, fine and coarse roots, non-structural carbon (NSC), and fruit tree pools. Carbon allocation among these pools was controlled by species-specific parameters, phenology, light and water availability. Water cycle was modeled calculating the daily balance between precipitation, canopy transpiration, evaporation, soil evaporation, and runoff. Meteorological variables used to force the model were: global solar radiation ($\text{MJ m}^{-2} \text{day}^{-1}$), maximum and minimum air temperature ($^{\circ}\text{C}$), relative humidity (%), and daily cumulated precipitation (mm day^{-1}). To be initialized, the model required knowing stand structural characteristics such as: species composition, stand density, diameter at breast height (DBH), tree height and age. Soil initialization required the estimation of total organic carbon (TOC) in the different SOC pools, as described in the following section.

Table 1. Site descriptions and stand initialization data. Mean temperature (T) and Precipitation (prec.) are annual averages collected at site.

| Site Name | Coords (Lat°/Lon°) | Climate | Species Composition | Mean Annual T (°C) | Mean Annual Prec. (mm year ⁻¹) | Mean DBH (cm) | Tree Height (m) | Stand Age (years) | Stand Density (trees ha ⁻¹) |
|------------------------|-----------------------|---------------|----------------------------------|-----------------------|---|------------------|--------------------|----------------------|--|
| Collelongo (ITCol) | 41.8/13.5 | Temperate | <i>Fagus sylvatica</i> L. (DBF) | 6.3 | 1180 | 20.27 | 19.84 | 100 | 900 |
| Hainich (DEHai) | 51.0/10.4 | Temperate | <i>Fagus sylvatica</i> L. (DBF) | 8.3 | 720 | 30.8 | 23.1 | 120 | 334 |
| Hyytiälä (FIHyy) | 61.8/24.2 | Boreal | <i>Pinus sylvestris</i> L. (ENF) | 3.8 | 709 | 10.3 | 6.5 | 39 | 1796 |
| Renon (ITRen) | 46.5/11.4 | Temperate | <i>Picea abies</i> L. (ENF) | 4.7 | 809 | 16.98 | 11.32 | 50 | 767 |
| Castelporziano (ITCpz) | 41.7/12.3 | Mediterranean | <i>Quercus ilex</i> L. (EBF) | 15.6 | 780 | 16 | 12.5 | 45 | 458 |
| Puechabon (FRPue) | 43.7/3.6 | Mediterranean | <i>Quercus ilex</i> L. (EBF) | 13.5 | 883 | 7 | 6 | 59 | 6149 |

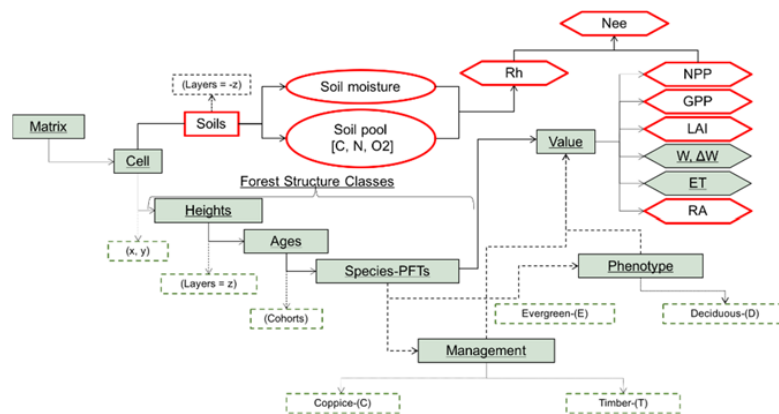


Figure 2. 3D-CMCC version of PSM main flowchart modified from [36]. Red-circled boxes represent the pools and variables introduced or modified by 3D-CMCC-PSM.

2.3. Model Improvements

2.3.1. Soil Carbon Dynamics

The most recent 3D-CMC FEM model version (v.5.1, [37]) lacks in representing SOC dynamics, preventing any estimation of NEE. With that perspective, we developed a simplified version of the method described in [38] to quantify, dynamically, changes of 7 different SOC pools mediated by the amount of active microbial C pool (Figure 3).

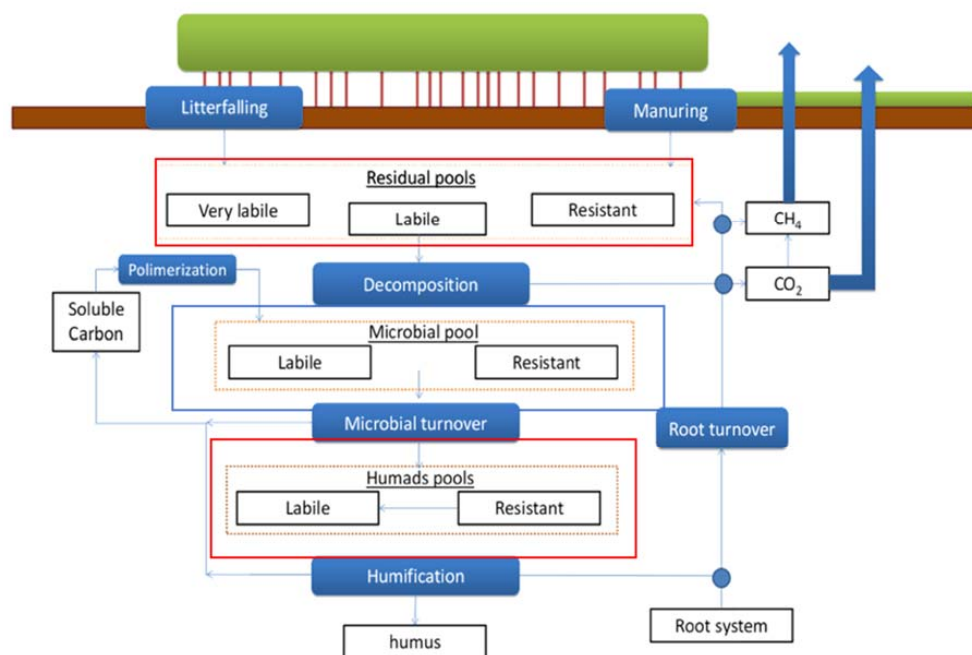


Figure 3. Soil Carbon dynamics in 3D-CMCC-PSM. The three macro pools are highlighted by red boxes (dead C pool) and blue box (live C pool, i.e., microbial). Blue-filled boxes represent the processes simulated by the soil model.

The litter C decomposed by microbial activity is partly mineralized as CO₂, partly stored into microbial metabolic biomass (labile), partly in structural microbial biomass (resistant), and partly transformed in other organic compounds [38]. SOC stability is also related to its chemical recalcitrance, its accessibility, and interaction with clays [45]. We divided SOC in 7 pools to take these differences

into account. Humic pool (we use the term Humads, to be consistent with [38]) was divided into a more labile (labile Humads, which stands for Humic acid), an intermediate (resistant Humads, which stands for Fulvic acid) and a more resistant sub-pool (Humus, which stands for Humine).

We assumed that microbial C use efficiency was different for different pools, but constant within each pool [44]. Microbial activity is strictly related to micro-environmental conditions too. Given these premises we modeled C dynamics among soil pools using the following two partial differential equations:

$$\begin{cases} \frac{\partial C_p}{\partial t} = C_L(t) - \beta C_p(t)B(t) \\ \frac{\partial B}{\partial t} = \beta C_p(t)B(t) - \alpha B(t) \end{cases} \quad (1)$$

with $\frac{\partial C_p}{\partial t}$ being the CO₂ efflux produced from a specific C pool decomposition, $C_L(t)$ is the amount of new carbon entering the specific soil pool, $C_p(t)$ the amount of carbon in that C pool, $B(t)$ the microbial biomass competing for $C(t)$. α and β respectively represented the microbial turnover and SOM consumption factors. α was treated as a constant value, as in [38].

The consumption factor β was estimated as a function of both SOM stability and micro-environmental conditions:

$$\begin{cases} \beta(C_{pool}) = \mu_T \mu_M \mu_A k_{pool} \\ \mu_T = -0.014 T^2 + 0.099 T + 0.02 \\ \mu_M = -2.85 \theta^3 + 1.49 \theta^2 + 1.77 \theta - 0.03 \\ \mu_D = 0.6 z^{-0.136} (-0.02 \text{ clay}\% + 0.03) \\ \mu_A = \mu_D \frac{\log(0.14 \text{ clay}\%^{-1})}{2.3026} + 1 \end{cases} \quad (2)$$

where μ_T represented the temperature factor, μ_M the moisture factor, μ_A the accessibility, k_{pool} the recalcitrance, and μ_D a clay dependent depth factor [46]. The k_{pool} factor was treated as a specific parameter depending on each compartment's biomass. T represented temperature in Celsius degrees, θ stands for water moisture, z the mean depth of the soil, $\text{clay}\%$ is the percentage of clays in soil. Humads were decomposed using the same rationale, but had slower rates. Inert organic matter (IOM) was calculated following [47].

2.3.2. Deciduous Phenology

Similarly to [37] the phenology scheme was constrained in 3 and 5 sub-phases respectively for evergreen and deciduous species (Table 2). These phases were driven by photoperiod, thermal sum, and maximum leaf biomass (resulting from maximum attainable Leaf Area Index, LAI, m² m⁻²).

3D-CMCC FEM represents leaves development as a by-product of leaf biomass pool dynamics [48]. Despite being a reasonable simplification, such method leads to non-negligible excesses in growth respiration and NSC consumption during budburst. Such unrealistic demand could eventually consume all the available carbon, causing the death of the tree. For this reason, we proposed a modification of budburst phenology, to explicitly simulate the dynamic tradeoffs between demand in NSC, increase in foliar biomass, and maturation of progressively self-sufficient leaves.

Table 2. Description of the different phenological phases for deciduous and evergreen species used in 3D-CMCC PSM.

| Deciduous | | Evergreen | |
|------------------|--------------------|-----------|--------------------|
| Phase | Trigger | Phase | Trigger |
| Bud Burst | GDD threshold | Bud Burst | GDD treshold |
| Leaf Development | PeakLai/2 | PeakLai | Pipe Model |
| PeakLai | Pipe Model | Leaffall | Daylength Treshold |
| Leaffall | Daylength Treshold | | |
| Unvegetative | [49] | | |

We based the tradeoff function on the hypothesis that new leaf demand of NSC is higher during budburst, and gets progressively lower when maturing [50]. Idealistically, the total NSC mass (B_R) demanded by maturing shoots and leaves could be represented by the linear differential equation (ODE):

$$\frac{\delta B_R}{\delta t} - R_{NSC}(t) \cdot B_R(t) = 0 \tag{3}$$

where $R_{NSC}(t)$ is the instantaneous proportion of NSC demand. We assumed that C request per leaf exponentially decreases with maturity, while the total demand increases by the increasing number of leaf primordia. For simplicity, we assumed that the two components resulted in the linear reduction of $R_{NSC}(t)$. Being a fraction, the maximum value of the integral of $R_{S\&F\&R}(t)$ is equal to 1. Being the first vegetative day t_0 , and the last day of budburst (BB_T) the last possible one to reach complete leaf development, the domain of the function was $[t_0, BB_T]$:

$$1 = \int_0^{BB_T} \frac{R_{NSC_{MAX}} \cdot t}{BB_T} \delta t \rightarrow R_{NSC_{MAX}} = \frac{2}{BB_T} \rightarrow R_{NSC}(t) = \frac{2t}{BB_T^2}, \tag{4}$$

resolving the ODE, and substituting $R_{S\&F\&R}(t)$ it gives:

$$B_R(t) = \frac{B_0 \cdot e^{\frac{2t+2}{BB_T^2}} \cdot e^{-t^2/BB_T^2}}{BB_T}, \tag{5}$$

where BB_T is the parameter used in [37], $e^{\frac{2t+2}{BB_T^2}}$ is the biomass dependent and $e^{-\frac{x^2}{BB_T^2}}$ the maturity dependent factors (expressed in days). Graphically, the equation represents a skewed function of the amount of NSC allowed to be used by the trees of the specific class; the faster the leaves reach maturity, the more daily specific allocation is allowed (Figure 4).

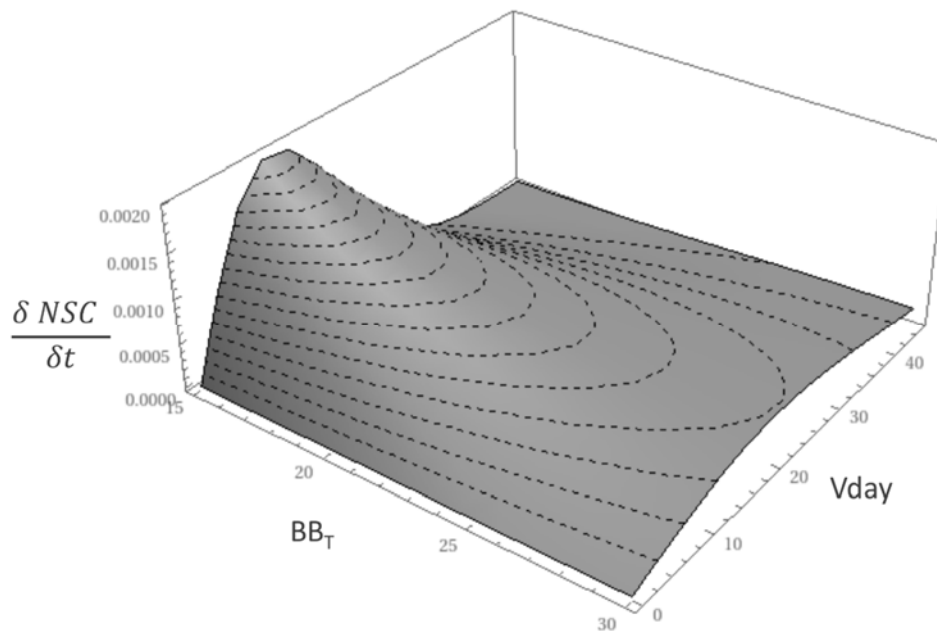


Figure 4. Graphic representation of the C tradeoff function. The axes represent respectively bud burst days (BB_T), vegetative days (Veg_day), and the fraction of total NSC invested in leaf development ($\frac{\partial NSC}{\partial t}$). The shorter BB_T , the higher the maximum NSC fraction.

Another major modification we introduced involves fall leaf yellowing and senescence. Falling leaves contribute to half of annual litter production. Correctly estimating their timing has a strong impact on both GPP and Rh dynamics [32]. The 5.1 version of 3D-CMCC-FEM represents senescence

by linearly decreasing leaves biomass of a predefined fraction until their pool is emptied. However, such method systematically overestimated leaf turnover at the beginning of the leaf fall phase, and is poorly flexible in calculating its duration. For these reasons, linear loss of leaf biomass was replaced with a sigmoid function. Assuming for hypothesis that all leaves fall by the end of senescence season, the sigmoid function was:

$$\frac{\partial L}{\partial t} = LAI(t) - \text{Max} \left(0, \frac{\alpha(h, a, sp)}{1 + e^{\frac{t-\beta(t)}{\gamma(t)}}} \right), \quad (6)$$

where $\alpha(t)$, $\beta(t)$, $\gamma(t)$ are three parameters with biological or physical meaning. In fact:

$$\begin{cases} \alpha(h, a, sp) = LAI_{0+\delta t}(h, a, sp) \\ \beta(s) = t_0(s) + \frac{\Delta t(s)}{2} \\ \gamma(s) \cong \frac{\Delta t(s)}{\ln(0.1) - \ln(10(\beta(s) - 0.1))} \end{cases}, \quad (7)$$

where $L_{0+\delta t}$ stands for LAI value at peak of green, $t_0(s)$ is the first day of senescence (triggered by a species-specific day-length parameter), $\Delta t(s)$ is the length of the senescence period (days). $\Delta t(s)$ was calculated using [49], as a function of temperature and photoperiod:

$$\begin{cases} R_{sen}(t) = (T_{Max}(s) - T(t)) \cdot \left(\frac{D_L(t)}{D_L(0)} \right)^2 \\ \Delta t(s) = \sum_{R_{Sen}(t)=0}^{R_{Sen}(t) < Y_c} R_{sen}(t) \end{cases}, \quad (8)$$

where T_{Max} is the maximum temperature at which senescence is effective, $T(t)$ is daily average Temperature, $D_L(0)$ photoperiod at the first day of senescence, $D_L(t)$ photoperiod at the i th day of the year.

2.3.3. Evergreen Phenology

Evergreen canopy turnover was modified from [37]. 3D-CMCCFEM v.5.1 assumes that evergreen leaf turnover is constant throughout the year, and that annual leaf turnover is equal to leaf biomass produced the year before [37,48]. However, leaf turnover seems to be concentrated in specific seasonal windows: (1) consistently to leaf emergence (spring); and (2) approaching of photosynthetic inefficient season (fall) [51]. For this reason, this approach may affect the ability in estimating leaf biomass and GPP intra-annual variability. To better represent leaf turnover dynamics, we developed a new framework where competition for light dynamically affected leaf turnover. We assumed the canopy to be a population of leaves optimized to intercept the highest amount of light. Since leaves cannot move from their position in the canopy, they get partially shaded by new emergent leaves when aging. Conceptually, leaves can be assumed to be in competition for one resource, light, and their turnover can be predicted by using a competition for one resource scheme as in [52]:

$$\begin{cases} \frac{\partial B_i(t)}{\partial t} B_i(t)^{-1} = f_i(R) - f_i(m_i) \\ f_i(R) = \frac{r_i \cdot R}{(R + K_i)} \\ R_i^* = \frac{m_i \cdot Y_i}{(r_i - m_i)} \end{cases}, \quad (9)$$

In this model, R and R_i^* are respectively the concentration of available light in $\text{J m}^{-2} \text{ day}^{-1}$ (i.e., resource that leaves are competing for), and the light needed to survive in a progressively more shaded canopy. r_i is the max photosynthetic rate ($\text{gC m}^{-2} \text{ day}^{-1}$), m_i is the maintenance respiration in $\text{gC m}^{-2} \text{ day}^{-1}$, Y_i carbon yield.

We assumed for hypothesis that:

- (1) Older leaves live in the shaded portions of the canopy, where light transmitted is reduced following Lambert Beer's exponential decay equation. For this reason, we expect an exponential reduction in absorbed photosynthetically active radiation (APAR);
- (2) An age dependent quasi-exponential decay in leaf quantum yield efficiency [53]. These decays impact on the reduction of r_i ;
- (3) Nitrogen content in older leaves is often lower than in young ones, because of its transfer from portions of the crown with low productivity to portions more exposed to light [53]. Since maintenance respiration is proportional to nitrogen content, we expect an exponential reduction in m_i ;
- (4) Y_i was assumed to be constant as in [17] because of the joint effect of reduction in respiration rate and quantum yield efficiency.

We assumed that the three components of the Equation (9) may have the following shapes:

$$\begin{cases} m_i = m_0 \cdot e^{-k_\eta \cdot t} + m_m \\ r_i = \left(\alpha_0 \cdot e^{-k_\alpha \cdot t} + \alpha_m \right) \cdot \left(\lambda_0 \cdot e^{-k_\lambda \cdot t} + \lambda_m \right) \\ k_i = \text{const} \end{cases}, \quad (10)$$

where t is time, the independent variable. m_0 , α_0 , λ_0 are MR, quantum yield and APAR at first day of budburst, the second year of life. m_m , α_m , λ_m are the theoretical minimum values of MR, quantum yield and APAR to let a proportion of leaves survive in a non-shading context. k_η , k_α , k_λ are the exponential parameters for the three functions.

Based on these hypotheses Equation (10) becomes:

$$R_i^* = \frac{\left(m_0 \cdot e^{-k_\eta \cdot t} + m_m \right) \cdot k_i}{\left(\alpha_0 \cdot e^{-k_\alpha \cdot t} + \alpha_m \right) \cdot \left(\lambda_0 \cdot e^{-k_\lambda \cdot t} + \lambda_m \right) - \left(m_0 \cdot e^{-k_\eta \cdot t} + m_m \right)}, \quad (11)$$

Assuming that $k_\eta = k_\alpha = k_\lambda$, and that the denominator of Equation (9) has to be greater than zero for the leaves in the i th generation to survive, R^* is a sigmoid positive function. Knowing R^* , we can calculate the amount of live leaf biomass of the i th generation ($_{(i)}$) as the inversion of R^* (i.e., $S_i^*(t)$). We simplified the theoretical model using the following function:

$$S_i^*(t) = \frac{1}{2}t^2 - \frac{2BF_{LS(i)} + 1}{2}t + BF_{LS(i)}, \quad (12)$$

where t is the number of days since leaves of the i th generation have emerged. According to this model the theoretical maximum age of each generation ($BF_{LS(i)}$) should correspond to the year in which the R^* almost reaches its asymptote. We used Equation (12) to quantify leaf turnover for each generation. About 60% of leaf turnover happened in early spring, when new leaves emerged: the amount of biomass lost every day was proportional to new leaf biomass production the same day. The rest of annual turnover happened in fall, when each leaf biomass generation was reduced of a constant value calculated from Equation (12). No foliar re-sprouting was simulated in fall, even though there are evidences of it for *Quercus ilex* [54]. Leaf biomass reduction was determined by linearly decreasing each B_i to the quantity predicted by the specific parabolic decay for the end of the year.

2.3.4. Production of Fresh Organic Matter

At a relatively fine temporal scale (i.e., daily time step) timing of litter formation and FOM production may be fundamental in correctly estimating R_h [55]. Littering for woody tissues followed the rationale of BIOME-BGM family [48]. Partitioning of leaves and fine root turnover followed [56]. FOM coming from any plant biomass pool to the soil was added to the litter pool. Litter pool was

consisting of three sub-pools: metabolic very labile C, structural labile C, and structural resistant C. FOM was divided into the three litter sub pools according to its original C:N ratio, as in [38]:

$$\left\{ \begin{array}{l} \frac{\partial}{\partial t} C_{i+1} = \frac{(CN_{lt}^{-1} - CN_i^{-1})}{(CN_i^{-1} - CN_{i+1}^{-1})} \cdot C_{lt} \\ \frac{\partial}{\partial t} C_i = \frac{\partial}{\partial t} C_{lt} - \frac{\partial}{\partial t} C_{i+1} \\ CN_i < CN_{lt} < CN_{i+1} \end{array} \right. , \quad (13)$$

where C_{lt} is the FOM entering litter from each structural C compartments of the plant. FOM C and N were distributed to the litter sub pools with closest C:N. C_{i+1} , represents the pool with higher recalcitrance. When CN_{lt} was higher than any litter sub pool, all the new C was added to the structural resistant pool; otherwise, if CN_{lt} was lower than the CN of the metabolic pool, all its C and N were added to the very labile sub pool. Litter C dynamically moved from a pool to another. Microbes absorbed and partially immobilized litter C in their biomass, and released it again in the soil, during the humification processes [57].

2.3.5. Optimization

We introduced a new calibration scheme to provide an optimized parameterization, and quantify uncertainty related to the equations used in 3D-CMCC-PSM to simulate NEE, and choice of a subset of 45 species-specific physiological parameters (Table S1). Calibration was performed on each site independently, using a training dataset composed by 3 years of EC daily NEE time-series. We chose 2000–2002 for DEHai, 2001–2003 for FIHyy, ITCpz, and FRPue; 2001–2004 for ITCol; 2006–2008 for ITRen. Because of the strong autocorrelation characterizing time-series, we didn't randomly split our dataset, and decided to use sub-time series of contiguous years [58]. We preferred to use the beginning of each simulation for training, before the global recession following the Financial Crisis of 2007–2008. We decided to do so to (1) facilitate comparison among different sites' simulations; (2) have the calibration start when vegetation structure was known from literature; and (3) reduce the risk for parameters to be fitted over a changing environment instead of eco-physiological properties.

To sample the parameterization space (the realistic values of each physiological parameter of the species simulated in each site), we randomly extracted 1000 parameter-set combinations from prior distributions. Prior distributions were assumed to be the same among individuals of the same species, different across species. We assumed each parameter to follow a truncated normal distribution, to avoid any possibility to have non-realistic negative values. Average and variance were estimated by using values found in literature, as in [37]. We used the same averaged value as in [37] for those parameters whose observations in literature were less than 3, because we didn't have enough information to calculate sample standard deviation. The optimization was performed by choosing the parameterization set maximizing the objective function QF through:

$$QF = \left(\frac{\sum_i^n Y_i^{obs} \cdot Y_i^{sim} - \frac{\sum Y_i^{obs} \cdot \sum Y_i^{sim}}{n}}{\sqrt{\sum (Y_i^{obs})^2 - \frac{(\sum Y_i^{obs})^2}{n}} \sqrt{\sum (Y_i^{sim})^2 - \frac{(\sum Y_i^{sim})^2}{n}}} \right)^2 \times \left[1 - \frac{\sum_i^n (Y_i^{obs} - Y_i^{sim})^2}{\sum_i^n (Y_i^{obs} - \bar{Y}^{obs})^2} \right], \quad (14)$$

where Y^{obs} represents EC daily NEE, Y^{sim} Modeled NEE for the same day, \bar{Y}^{obs} the average EC daily NEE over the train time series. The first part of the RHS of the equation represents the square of the Pearson Correlation coefficient (R), the second the Nash-Sutcliffe Efficiency index (NSE).

2.3.6. Validation Analysis

Results were compared to Eddy Covariance data on long-term daily, monthly and annual averages, over the full series of testing years (~5 years). For the validation, we used 2003–2007 for DEHai, 2004–2008 for FRPue and ITCpz, 2004–2011 for FIHy, 2005–2012 for ITCol, 2009–2011 for ITRen. Then we evaluated how the model performed in the different seasons aggregating values for months of the same season.

To evaluate the model efficiency, we calculated daily, monthly, and seasonal: (1) R; (2) NSE; (3) root mean square error (RMSE); and (4) mean absolute bias (MAB). Each statistic was considered differently informative [59] as summarized in Table 3. The model's ability in representing observed anomalies was determined by analyzing inter annual variability (IAVs) following [60] and [37].

Table 3. Statistics used for Model's results validation against Eddy Covariance data.

| Statistics | Formulation | Use and Ranges |
|---------------------------|--|---|
| Pearson Coefficient | $r = \frac{\sum_i^n Y_i^{obs} \cdot Y_i^{sim} - \frac{\sum Y_i^{obs} \cdot \sum Y_i^{sim}}{n}}{\sqrt{\sum (Y_i^{obs})^2 - \frac{(\sum Y_i^{obs})^2}{n}} \sqrt{\sum (Y_i^{sim})^2 - \frac{(\sum Y_i^{sim})^2}{n}}}$ | Estimation of model's measure of correlation with EC data [0;1] |
| Nash Sutcliffe efficiency | $NSE = 1 - \frac{\sum_i^n (Y_i^{obs} - Y_i^{sim})^2}{\sum_i^n (Y_i^{obs} - \bar{Y}^{obs})^2}$ | Estimation of model's predictability $[-\infty;1]$ |
| Root Mean Square Error | $RMSE = \sqrt{\frac{\sum_i^n (Y_i^{obs} - Y_i^{sim})^2}{n}}$ | Estimation of model's accuracy $\text{gC m}^{-2} \text{ day}^{-1}$ [0; ∞] |
| Mean Absolute Bias | $MAB = \frac{1}{n} \sum_i^n \frac{ Y_i^{obs} - Y_i^{sim} }{\sigma(Y_i^{obs})}$ | Estimation of model's bias $\text{gC m}^{-2} \text{ day}^{-1}$ [0; ∞] |

3. Results

3.1. Evaluation of Daily, Seasonal, and Annual NEE Estimations

To evaluate 3D-CMCC-PSM NEE predictions, we compared predicted (MD) daily and monthly NEE time series to EC daily data. The analyses were performed only on the test data (i.e., portions of the series which have not been used for calibration) to avoid any effect of overfitting. The model showed high correlations with observed EC data at all sites for both daily and monthly fluxes, apart from ITCpz site (Table 4). Excluding ITCpz, R ranged at all sites from 0.65 to 0.84 for daily, and 0.59 to 0.97 for monthly scale. Beech-dominated Deciduous Forests (DBF) performed better than Conifer species (ENF) and evergreen Mediterranean broadleaved forests (EBF). ENF and EBF in FRPue performed similarly on daily scale, for all the statistics used. However, ENF predictability significantly increased on monthly scale (R ranging between 0.92 and 0.97), while EBF performed worse (R 0.42 in ITCpz, and 0.59 in FRPue). RMSE on average was $1.92 \text{ gC m}^{-2} \text{ day}^{-1}$. MAE ranged between 0.96 and $1.78 \text{ gC m}^{-2} \text{ day}^{-1}$, and on average it decreased almost twice on monthly timescale. MAB showed similar behavior for DBF and ENF. It ranged between 0.39 and $0.56 \text{ gC m}^{-2} \text{ day}^{-1}$ (0.50 on average) for daily time series. Mediterranean forests resulted as the ones with highest MAB, and showed no significant reduction when predictions were aggregated on monthly scale. Differently from the other simulations, even NSE just improved slightly for ITCpz, and even reduced for FRPue simulation.

Table 4. Daily and Monthly Validation statistics calculated on the test-set. As stated in Table 3, *R* and *NSE* are dimensionless; *RMSE* and *MAB* are $\text{gC m}^{-2} \text{day}^{-1}$.

| | DEHai | ITCol | FIHyy | ITRen | ITCpz | FRPue | Mean |
|-------------|-------|-------|-------|-------|-------|-------|------|
| Daily NEE | | | | | | | |
| <i>R</i> | 0.84 | 0.76 | 0.67 | 0.65 | 0.24 | 0.65 | 0.64 |
| <i>NSE</i> | 0.67 | 0.5 | 0.34 | 0.21 | -0.26 | 0.35 | 0.3 |
| <i>RMSE</i> | 1.84 | 2.7 | 1.48 | 2.32 | 1.8 | 1.39 | 1.92 |
| <i>MAB</i> | 0.39 | 0.5 | 0.53 | 0.56 | 1.15 | 0.76 | 0.65 |
| Monthly NEE | | | | | | | |
| <i>R</i> | 0.93 | 0.92 | 0.96 | 0.97 | 0.42 | 0.59 | 0.8 |
| <i>NSE</i> | 0.81 | 0.76 | 0.9 | 0.87 | 0.12 | 0.2 | 0.61 |
| <i>RMSE</i> | 1.15 | 1.58 | 0.45 | 0.72 | 1.24 | 1 | 1.02 |
| <i>MAB</i> | 0.28 | 0.32 | 0.21 | 0.24 | 1.25 | 0.86 | 0.53 |

Daily results were aggregated in seasonal series to evaluate seasonal predictability. Daily NEE were averaged to give a time series of mean seasonal NEE ($\text{gC m}^{-2} \text{day}^{-1}$), one value per season, for the duration of the test dataset. Seasonal aggregations showed that 3D-CMCC-PSM poorly performed in predicting seasonal fluxes. *NSE* was generally negative in summer, with the exclusion of DEHai and FRPue. 3D-CMCC-PSM generally best reproduced NEE dynamics in fall (*R* ranging between 0.22 and 0.89). ENF ecosystems showed consistently higher correlation in spring predictions, with *R* of 0.65 and *MAB* of $0.62 \text{ gC m}^{-2} \text{day}^{-1}$ on average. In the case of evergreen stands, 3D-CMCC-PSM consistently showed poor performance in summer. Expectedly, DBF performed the worst in winter (Table 5). *NSE* on average resulted positive only in fall for both DBF and EBF, and spring, for ENF stands.

Table 5. Seasonal validation statistics calculated on the test-sets and aggregated by ecosystem type. As stated in Tables 3 and 4, *R* and *NSE* are dimensionless; *RMSE* and *MAB* are $\text{gC m}^{-2} \text{day}^{-1}$. MAM stands for March, April, and May. JJA stands for June, July, and August. SON stands for September, October, and November. DJF stands for December, January, and February. DBF stands for Deciduous Broadleaf Forests, EBF for Evergreen Broadleaf Forests, ENF for Evergreen Needle leaf Forests.

| | <i>R</i> | <i>NSE</i> | <i>RMSE</i> | <i>MAB</i> |
|-----|----------|------------|-------------|------------|
| DBF | | | | |
| MAM | 0.43 | -0.36 | 2.77 | 0.72 |
| JJA | 0.36 | -0.02 | 2.96 | 1.17 |
| SON | 0.82 | 0.58 | 1.9 | 0.58 |
| DJF | 0.2 | -0.93 | 0.7 | 1.58 |
| EBF | | | | |
| MAM | 0.65 | 0.28 | 1.83 | 0.62 |
| JJA | 0.11 | -0.92 | 2.76 | 1.01 |
| SON | 0.51 | 0.03 | 1.56 | 0.74 |
| DJF | 0.38 | -1.59 | 0.51 | 0.82 |
| ENF | | | | |
| MAM | 0.18 | -0.41 | 1.95 | 1.2 |
| JJA | 0.32 | -0.34 | 1.58 | 1 |
| SON | 0.45 | -0.19 | 1.3 | 0.79 |
| DJF | 0.47 | -6.13 | 1.45 | 1.4 |

We used Taylor diagrams [61] to graphically summarize how closely Daily, Monthly and Annual NEE patterns matched EC observations (Figure 5). 3D-CMCC-PSM performance was generally satisfactory. Daily simulations resulted in all sites but ITCpz being within the ± 1 normalized standard deviation region. Monthly scale predictions were more consistent with EC data, especially for BDF and ENF sites. It resulted in all 4 simulations falling within ± 0.5 normalized standard deviation from the reference point, and *R* > 0.9. Again, 3D-CMCC-PSM performed worst in EBF, with FRPue still

inside ± 1 normalized standard deviation region, and ITCpz falling outside the ± 1.5 normalized SD region. The consistently worse predictability in ITCpz and FRPue confirm a systematic weakness in 3D-CMCC to represent fluxes for these sites as already described in [37]. Model performance on annual scale showed a different pattern, mostly because of some sites' consistent biases in seasonal NEE, and the difference in NEE magnitude. Delay in spring phenology, and the consistent underestimation of summer NEE, resulted in significant underestimation and scarce predictability of ITCol annual NEE ($R < 0.2$). ITCpz and FRPue resulted among the sites with higher annual predictability, partly because of the low seasonal variance in NEE, partly because winter and spring bias tends to compensate each other.

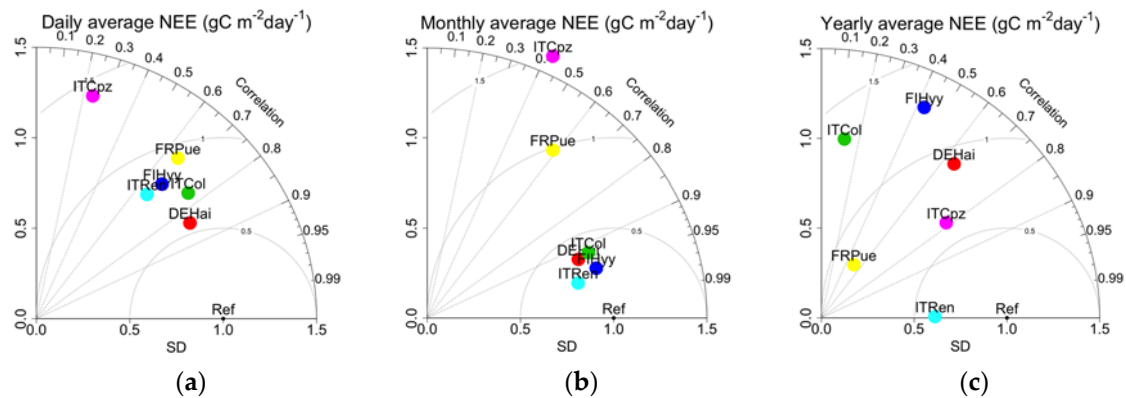


Figure 5. Taylor diagrams representing 3D-CMCC-PSM performance in (a) daily, (b) monthly and (c) annual NEE estimation for the test-set. ITCol and DEHai represent DBF (red and green dots); ITRen and FIHyy ENF (blue and turquoise). ITCpz and FRPue represent EBF (yellow and magenta dots). The closest a simulation lied to the “Ref” point, the better 3D-CMCC-PSM represented NEE patterns. X- and Y- axes represent NEE standard deviation (SD): the closest to 1, the better the performance. Simulations with $R \geq 0.9$, and difference in SD with EC NEE less than $0.5 \text{ gC m}^{-2} \text{ day}^{-1}$ showed very good performance. Simulations with $0.75 \leq R < 0.9$, and difference in SD with EC NEE between 0.5 and $1 \text{ gC m}^{-2} \text{ day}^{-1}$ showed good performance. Simulations with $0.35 \leq R < 0.75$, and difference in SD with EC NEE between 1 and $1.5 \text{ gC m}^{-2} \text{ day}^{-1}$ showed sufficiently good performance.

3.2. Anomalies and Parameters Related Uncertainty

Figure 6 shows uncertainty associated to random choice of parameters. Overall, uncertainty was expectedly higher in summer and fall. Such increase was particularly clear for deciduous forests, which not only showed wider NEE standard deviation, but also had optimal modeled NEE falling outside standard deviation area.

DBF sites showed also high uncertainty in estimating the first vegetative day, suggesting that a better representation of winter dormancy effects on budburst dates may significantly improve model's predictability. Uncertainty was generally lower in Mediterranean sites, probably because model's performance was generally lower, and fitting parameters to data would have little effect on performance. 3D-CMCC-PSM uncertainty was generally low for ENF for most of the year, but was generally high when temperatures were higher. Higher uncertainty for warmer days was generally found in DBF sites too, suggesting that 3D-CMCC-PSM was expectedly sensitive to high temperatures for both photosynthesis [62] and respiration [23]; because of such variability, calibrating parameters on data resulted in a significant boost in model's predictions.

NEE inter-annual variability was generally underestimated by the model. Nevertheless, 3D-CMCC-PSM correctly reproduced 81% of the sign of the anomalies, and residual difference in magnitude was usually less than $0.3 \text{ gC m}^{-2} \text{ day}^{-1}$. Highest difference in magnitude occurred in ITCol (difference in residuals higher than $0.5 \text{ gC m}^{-2} \text{ day}^{-1}$ in 5 years out of 12). Highest difference

was shown in ITCpz, where the sign was correctly reproduced only once out of 8 years, and having more than $1\text{gC m}^{-2}\text{ day}^{-1}$ of residual difference (Figure 7).

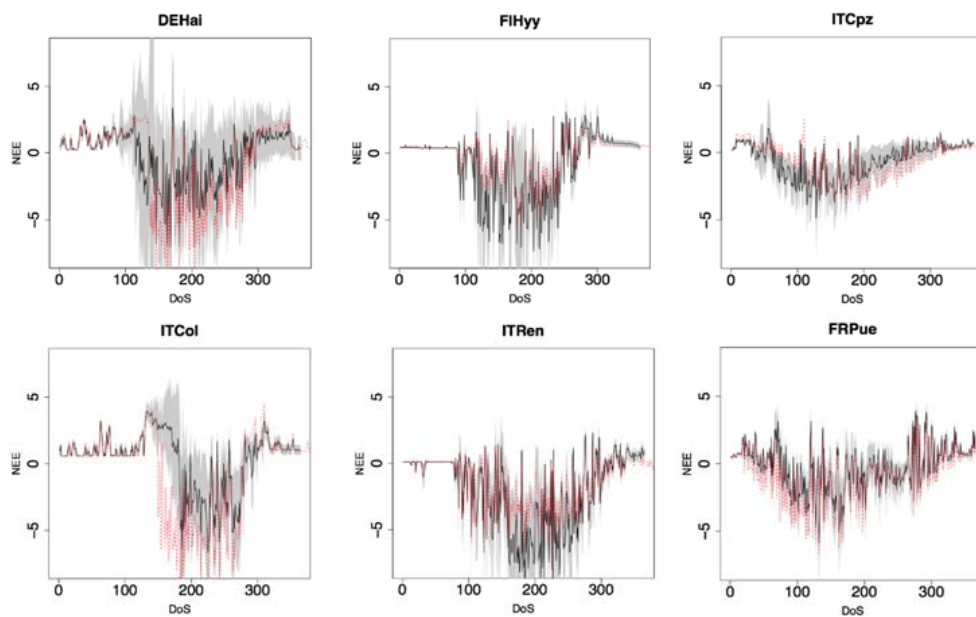


Figure 6. Model structure-related uncertainty in estimating NEE ($\text{gC m}^{-2}\text{ day}^{-1}$) per DoS (Day of Simulation) by a random choice of parameter values from prior distributions. Data represent 300 1-year simulations from randomly extracted parameterization-sets. Average daily simulations (black lines) and standard deviation (grey areas). Red dotted lines represent daily NEE simulation for the optimized parameterization set (Table S2). First column represents DBF sites, the second ENF, the third EBF.

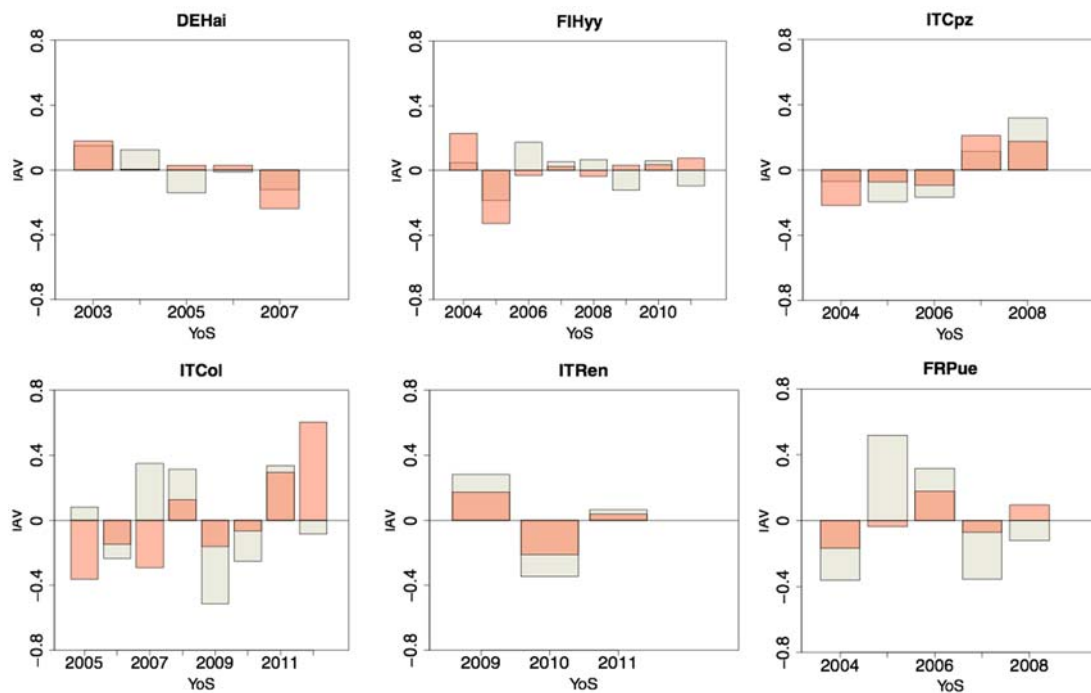


Figure 7. Inter-annual variability (IAV) for the test time-series ($\text{gC m}^{-2}\text{ month}^{-1}$). Observed IAV in grey boxes, simulated IAV in orange. First column represents DBF sites, the second ENF, the third EBF.

3.3. Comparison with the 5.1 Version of 3D-CMCC-FEM

We compared the results of 3D-CMCC-FEM 5.1 [37] and 3D-CMCC-PSM for daily and monthly GPP for ITRen, FRPue, DEHai and FIHy. We used the same non-optimized parameterization set for both versions. Except for FR-Pue, 3D-CMCC-PSM showed lower RMSE and higher R for daily GPP. Daily NSE too was generally higher in 3D-CMCC-PSM, except for FIHy. Monthly aggregated predictions were consistently outperforming those of 3D-CMCC-FEM 5.1 (Table 6).

Table 6. Comparisons between 3D-CMCC-FEM 5.1 (5.1) and 3D-CMCC-PSM (PSM) versions, using the same parameterization for 4 out of 6 sites (ITRen, FRPue, DEHai, FIHy). As stated in Table 3, R and NSE are dimensionless; RMSE is $\text{gC m}^{-2} \text{day}^{-1}$. Bold values represent best performing version.

| | Version | ITRen | FRPue | DEHai | FIHy | Avg |
|---------|---------|-------------|-------------|-------------|-------------|-------------|
| Daily | 5.1 | 0.81 | 0.82 | 0.92 | 0.91 | 0.87 |
| R | PSM | 0.88 | 0.64 | 0.93 | 0.91 | 0.85 |
| Monthly | 5.1 | 0.95 | 0.64 | 0.97 | 0.96 | 0.89 |
| R | PSM | 0.96 | 0.84 | 0.98 | 0.96 | 0.94 |
| Daily | 5.1 | 0.61 | −0.54 | 0.84 | 0.87 | 0.45 |
| NSE | PSM | 0.72 | 0.09 | 0.96 | 0.76 | 0.63 |
| Monthly | 5.1 | 0.91 | −0.11 | 0.94 | 0.91 | 0.66 |
| NSE | PSM | 0.91 | 0.56 | 0.98 | 0.92 | 0.84 |
| Daily | 5.1 | 2.09 | 1.52 | 1.85 | 1.56 | 1.76 |
| RMSE | PSM | 1.59 | 1.96 | 1.91 | 1.57 | 1.76 |
| Monthly | 5.1 | 0.97 | 1.01 | 1.07 | 0.91 | 0.99 |
| RMSE | PSM | 0.82 | 1.09 | 0.82 | 0.93 | 0.92 |

3.4. Daily and Monthly Reco

We evaluated Reco (ecosystem respiration) by comparing modeled (MD) and observed (EC) daily and monthly time series. Daily R ranged between 0.45 in ITCpz and 0.9 in FIHy (Table 7). RMSE was of $1.28 \text{ gC m}^{-2} \text{ day}^{-1}$ on average, and MEB ranged between 0.43 and $1 \text{ gC m}^{-2} \text{ day}^{-1}$. Most of the bias happened in summer, where Reco was generally overestimated, especially in ITCol and ITCpz. NSE was positive in any case but ITCpz. It was generally lower in DBF (0.32), and higher in FIHy (0.60) and FRPue (0.57). Reco at monthly timescale was strongly improved in predictability, especially for ITRen, whose R increased to 0.67 and NSE to 0.84. Monthly predictions showed improvements for the other simulations too, improving R and NSE of about 0.07. Since most of the bias occurred in summer, monthly predictions showed no dramatic improvements for neither RMSE nor MEB. The only exception was ITRen, whose RMSE reduced of about $0.4 \text{ gC m}^{-2} \text{ day}^{-1}$, suggesting that daily Reco may be noisy.

Table 7. Daily and Monthly Validation statistics for R_{eco} calculated on the test-set. As stated in Table 3, R and NSE are dimensionless; RMSE and MAB are $\text{gC m}^{-2} \text{ day}^{-1}$.

| | DEHai | ITCol | FIHy | ITRen | ITCpz | FRPue | Mean |
|--------------|-------|-------|------|-------|-------|-------|------|
| Daily Reco | | | | | | | |
| R | 0.79 | 0.71 | 0.90 | 0.67 | 0.45 | 0.86 | 0.73 |
| NSE | 0.32 | 0.32 | 0.60 | 0.34 | −0.43 | 0.57 | 0.29 |
| RMSE | 1.29 | 1.83 | 1.18 | 1.03 | 1.65 | 0.70 | 1.28 |
| MEB | 0.63 | 1.00 | 0.43 | 0.62 | 0.98 | 0.48 | 0.69 |
| Monthly Reco | | | | | | | |
| R | 0.86 | 0.79 | 0.96 | 0.84 | 0.54 | 0.93 | 0.82 |
| NSE | 0.40 | 0.37 | 0.66 | 0.69 | −0.40 | 0.67 | 0.40 |
| RMSE | 1.10 | 1.67 | 1.04 | 0.64 | 1.49 | 0.54 | 1.08 |
| MEB | 0.60 | 0.99 | 0.41 | 0.50 | 1.04 | 0.43 | 0.66 |

4. Discussion

4.1. 3D-CMCC-PSM Predictability in Estimating NEE

In general, the inclusion of a simplistic SOC routine resulted in a reliable estimation of daily and monthly NEE patterns. While daily and monthly patterns are consistent with EC data, seasonal patterns showed non-negligible misrepresentations, which resulted in negative NSE in most of the cases. This inconsistency may be driven by the strong seasonality in both R_{eco} and GPP [63], which positively affects correlation between EC data and MD results.

NEE patterns during summer and fall were much more consistent with measured ones, than winter and spring patterns. During these seasons the biases appeared mostly affected by estimation of R_{eco} . The scarcity of the model in representing EBF C fluxes was especially attributable to GPP predictions. 3D-CMCC-PSM and 3D-CMCC FEM inability in predicting GPP in ITCpz and FRPue sites, denoted the necessity to better represent the relations between Mediterranean forests and environmental factors [64,65]. In FRPue the model well reproduced spring, summer and fall NEE. On the other hand, it showed a bias of around $1 \text{ gC m}^{-2} \text{ day}^{-1}$ in winter, suggesting it was missing some particularly important seasonal processes. For example, evergreen phenology still didn't consider secondary or continuous growth. Thus, species like *Quercus ilex*, which exhibit secondary gem sprouts in fall [54], have fresh leaves and mild temperatures to guarantee photosynthetic activity in fall and winter, partly explaining 3D-CMCC FEM and 3D-CMCC-PSM systematic underestimation.

ITCpz showed the same pattern. However, differently from FRPue, it poorly performed also in early spring and summer, especially for GPP patterns. This poor performance was expected, because of the physical characteristics of the site. In fact, 3D-CMCC FEM soil water dynamics routine was still simplistic, and to date, such other similar models have not included any effect of water table dynamics. On the contrary, ITCpz is characterized with the presence of a shallow groundwater table, which seems to reduce water stress in early summer [37]. Moreover, we used average daily meteo data collected from the Eddy Covariance stations, and initialized simulations using average structural information found in literature. Uncertainty about those data was potentially high, and could have dramatically affected 3D-CMCC-PSM results.

Summer NEE misrepresentation in DBF was probably affected by the assumption that LAI and photosynthetic capacity reach their maximum in early summer, at the same time. On the contrary, maximum photosynthetic capacity may be reached in late summer, and varies across the canopy. Without taking this into account, GPP could be overestimated up to 40% [66]. Notwithstanding, comparing model outputs with published works [67,68], these defects are common also for other PBMs.

Seasonal patterns showed that the model consistently misrepresented NEE in winter, suggesting that R_{eco} still needs to be improved. Especially for DBF sites (e.g., DEHai), winter R_{eco} was mostly driven by RH. RH was exponentially affected by soil temperatures and especially moisture [69], which are calculated by the model, and could be over-fluctuating in winter. Moreover, EC data are prone to random noise [70], whose relative impact on performance metrics may be relatively larger.

Interestingly, annual predictions suggested reasonably high performance of 3D-CMCC-PSM, despite these seasonal inconsistencies. This suggests that biases are usually consistent within a season, but have different signs across seasons (Figure 8), resulting in a compensating effect at coarser time scale.

It was not always possible to individually validate the different components of respiration. Since EC Reco was not measured but inferred, evaluation metrics should be interpreted as a general ability of 3D-CMCC-PSM in predicting it. Despite daily Reco being noisy, 3D-CMCC-PSM could reproduce respiration processes well enough, at least at monthly timescale.

NPP:GPP ranged between 0.37 and 0.62, and was consistent with literature [71–73]. It was generally higher in DBF (0.49 in DEHai, 0.55 in ITCol), lower in EBF (0.38 in FRPue, 0.41 in ITCpz). These results matched with those of [71] who showed that the ratio between NPP and GPP (CUE) is

generally 0.53, ranging from 0.23 to 0.83. CUE was relatively low in FIHy, where Reco predictions were overestimated: therefore, poor predictability was probably ascribable to excesses in RA.

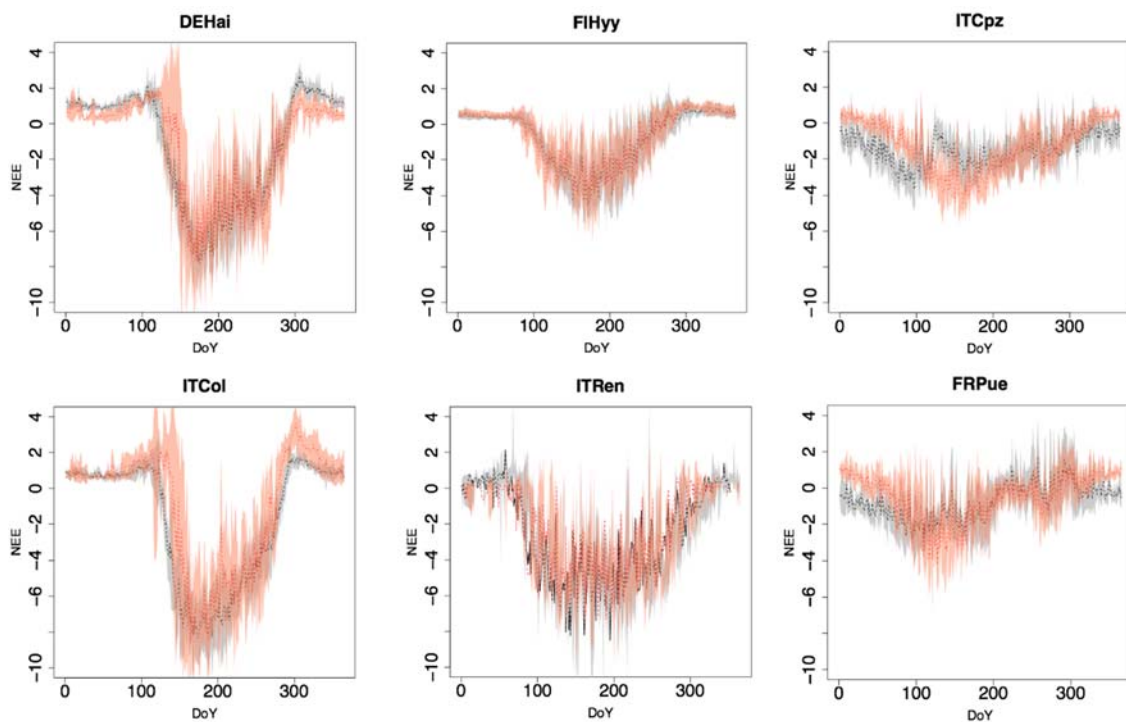


Figure 8. Patterns in daily NEE ($\text{gC m}^{-2} \text{day}^{-1}$) per DoY (Day of Year) calculated from test-sets on a site level. Observed EC average patterns (black dotted line) and standard deviation (gray area). Simulated average patterns (red dotted line) and standard deviation (orange area). First column represents DBF sites, the second ENF, the third EBF.

Ecosystem respiration was overestimated during summer, causing NEE systematic overestimation at each simulation, especially in ITCol. Except for FRPue and ITCol, RA grew about $1 \text{ gC m}^{-2} \text{day}^{-1}$ higher than RH, suggesting that it may be the principal driver of biased summer Reco (Figure 9). This misbehavior may be related to the method used to estimate maintenance respiration, an exponential relationship between respiration, moisture, and temperature [61].

Lack of data to validate the SOC dynamics reduced the spectrum of speculations, which could be statistically analyzed. SOC didn't change its quantity in ten years; this result was consistent with the theoretical stability of the SOC, an indicator which rarely changes within 10 years if no strong disturbance events (e.g., land use changes) have occurred [74]. Litter C was highly fluctuating within a year, but its quantity was stable if compared at the end of each year. This suggested that the model realistically represented litter turnover and decomposition, since residues were degraded into humus labile substances about within a year [75]. Microbial biomass was highly variable, as expected. However, the magnitude of change was too broad throughout the simulations. These results may be related to the use of 5% as the initial active microbial biomass for each site, a value that may be far from the equilibrium for different soils. Moreover, tradeoffs within microbial growth and between the environmental conditions may be scarcely represented. As a matter of fact, 3D-CMCC-PSM simulated the soil as having the same physical-chemical structure throughout the profile. This implied that microbes could find the same amount of C, O_2 , and living space, with no depth limitation.

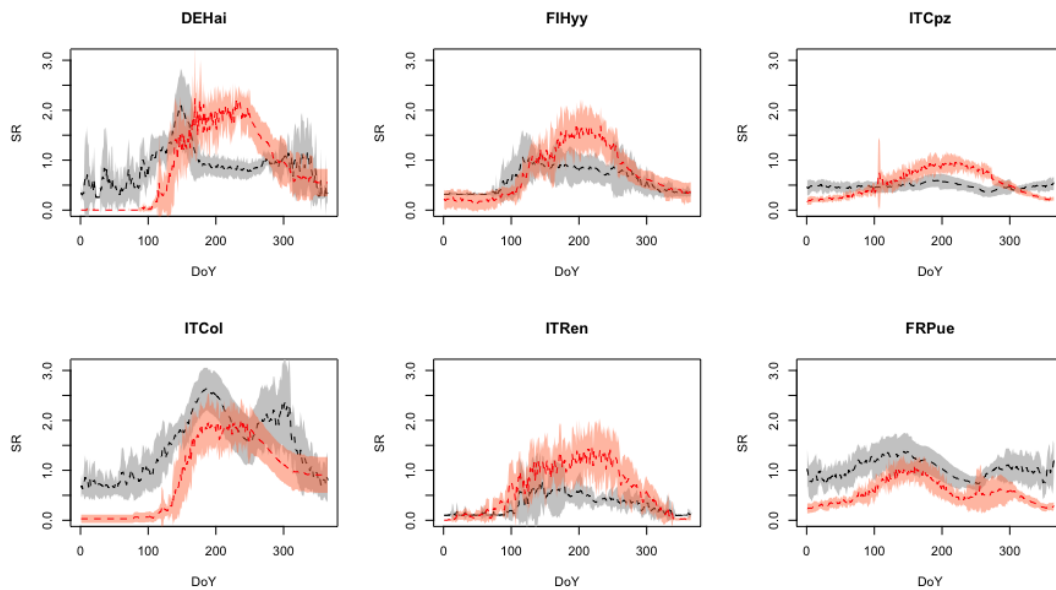


Figure 9. Patterns in daily soil respiration (SR) heterotrophic and autotrophic components ($\text{gC m}^{-2} \text{day}^{-1}$). Patterns per DoY (Day of Year) calculated from test-sets on a site level. Microbial respiration average patterns (black-dotted lines) and standard deviation (gray areas). Autotrophic root respiration average patterns (red-dotted lines) and standard deviation (orange areas). First column represents DBF sites, the second ENF, the third EBF.

4.2. 3D-CMCC-PSM Uncertainty in Estimating NEE

We analyzed 300 random parameterization-sets per site to quantify model assumptions and uncertainty. The model showed different behaviors in different sites, but expectedly consistent across species behaving in a similar functional way. These results may suggest that using functional traits combinations to provide physiological parameters, instead of fixed species-specific ones, may produce still reliable and more general predictions, particularly useful in case of larger spatial/temporal simulations [76–78]. Using a species-level parameterization, in fact, may result in a too-fine “resolution” because: (1) it would require excessive computational resources and a finely-detailed parameterization, usually inaccessible on a broad scale [79]; and (2) the model’s rationale in predicting forest structure is mainly driven by competition for resources. However, there are not explicit tradeoffs, positive interactions between different tree cohorts, or intra-specific trait variability, which are fundamental to forecast forest ecosystem structure on long-run simulations [80]. Having fixed species-specific parameters throughout a century would potentially result that only a very reduced amount of species would dominate the different cohorts on landscape to regional scale.

According to Figures 6–8, strong uncertainties still reside in timing for the different phenological phases. The biggest source of uncertainty in deciduous stands was driven by the amount of degree days needed to begin the vegetative period. The use of a site-specific thermal sum (GDD) to activate vegetation period is widely used, but proven to be very site sensitive, and not very effective for a regional generalization [81]. On the other hand, the processes triggering bud-burst timing are still partly unknown. Moreover, those models proposing a process-oriented promoter-inhibition rationale are generally over-complex and not prone to spatial generalization [82]. A possible solution in this context is to use remotely-sensed data to train a latitudinal-explicit regression, constraining GDD estimation.

3D-CMCC-PSM showed high uncertainty also in catching the beginning of the senescence phase. The new phenological scheme didn’t reduce such uncertainty, since it was still using a photoperiod threshold as the senescence phase trigger [37]. Another strong source of uncertainty in summer GPP may be held by the over-simplicity of soil structure and thus of the soil water routine.

As shown in [37], EC data are prone to high uncertainty. We focused on NEE fluxes to reduce the uncertainty cascade related to NEE partitioning. The next natural step will be reframing the model with a hierarchical Bayesian fashion, to quantify error propagation and parameter uncertainty from the posterior distribution [83].

Daily R_{eco} estimation was affected by the cascade of uncertainties related to the calculation of R_A and heterotrophic respiration, calculated independently. R_A routine may strongly be influenced by uncertainties in R_M estimation, which often resulted in R_A overestimation. The R_M was in fact simulated by a set of empirical relations, which involve the use of a fixed non-acclimating Q10 factor, whose generality is known to be inaccurate [84]. Moreover, the rationale of Ryan's R_M calculation [85] is affected by uncertainty in estimating daily increment of N pools, generally estimated by forest ecosystem models as a fixed proportion of daily C increment.

5. Conclusions

Soil respiration has a key role in determining NEE in a deterministic fashion [86]. In general, this work showed how the inclusion of a simplistic soil carbon routine allowed prediction of trends and variability of NEE across the most diffuse European forest ecosystems. Modifications in the phenology scheme produced slight improvements in predicting GPP. However, they were still limited by the correct estimation of bud burst timing, leaf senescence starting point and duration. The use of an optimized parameter-set improved the model's performance only for those sites where the bio-geophysical processes were correctly reproduced. As a matter of fact, we showed how Mediterranean terrestrial forests, which showed lacks in representing some biological and/or physical processes, performed significantly worse than the other modeled sites, regardless of the use of optimized parameters.

In conclusion, we think that 3D-CMCC-PSM can reliably estimate NEE and R_{eco} dynamics in a forest ecosystem, especially in scaling up daily results to monthly NEE averages. We think that 3D-CMCC-PSM is a solid basis to further explore the effects of soil structure on carbon and Water dynamics, especially in Mediterranean systems, and be used as a tool for predicting forest growth and ecosystem services, and address questions related to future scenario forecasting.

Supplementary Materials: The following are available online at www.mdpi.com/1999-4907/8/6/220/s1, Table S1: Mean and standard deviation of the priors used to sample parameters used for the optimization, Table S2: Values of the parameters used for the 6 simulations, produced by the optimization scheme.

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References

- Baldocchi, D.; Ryu, Y.; Keenan, T. Terrestrial Carbon Cycle Variability. *F1000Res* **2016**, *5*. [[CrossRef](#)] [[PubMed](#)]
- Göckede, M.; Foken, T.; Aubinet, M.; Aurela, M.; Banza, J.; Bernhofer, C.; Bonnefond, J.M.; Brunet, Y.; Carrara, A.; Clement, R.; et al. Quality control of CarboEurope flux data & ndash; Part 1: Coupling footprint analyses with flux data quality assessment to evaluate sites in forest ecosystems. *Biogeosciences* **2008**, *5*, 433–450. [[CrossRef](#)]
- Xiao, J.; Zhuang, Q.; Law, B.E.; Chen, J.; Baldocchi, D.D.; Cook, D.R.; Oren, R.; Richardson, A.D.; Wharton, S.; Ma, S.; et al. A continuous measure of gross primary production for the conterminous U.S. derived from MODIS and AmeriFlux data. *Remote Sens. Environ.* **2010**, *114*, 576–591. [[CrossRef](#)]
- Tang, X.G.; Liu, D.W.; Song, K.S.; Munger, J.W.; Zhang, B.; Wang, Z.M. A new model of net ecosystem carbon exchange for the deciduous-dominated forest by integrating MODIS and flux data. *Ecol. Eng.* **2011**, *37*, 1567–1571. [[CrossRef](#)]
- Reichstein, M.; Falge, E.; Baldocchi, D.D.; Papale, D.; Valentini, R.; Aubinet, M.; Berbigier, P.; Bernhofer, C.; Buchmann, N.; Gilmanov, T.; et al. On the separation of net ecosystem exchange into assimilation and ecosystem respiration: Review and improved algorithm. *Glob. Chang. Biol.* **2005**, *11*, 1–16. [[CrossRef](#)]
- Lasslop, G.; Migliavacca, M.; Bohrer, G.; Reichstein, M.; Bahn, M.; Ibrom, A.; Jacobs, C.; Kolari, P.; Papale, D.; Vesala, T.; et al. On the choice of the driving temperature for eddy-covariance carbon dioxide flux partitioning. *Biogeosciences* **2012**, *9*, 5243–5259. [[CrossRef](#)]
- Potter, C.S.; Randerson, J.T.; Field, C.B.; Matson, P.A.; Vitousek, P.M.; Mooney, H.A.; Klooster, S.A. Terrestrial ecosystem production: A process model based on global satellite and surface data. *Glob. Biogeochem. Cycles* **1993**, *74*, 811–841. [[CrossRef](#)]
- Prince, S.D.; Goward, S.N. Global primary production: A remote sensing approach. *J. Biogeogr.* **1995**, *22*, 815–835. [[CrossRef](#)]
- Mäkelä, A.; Landsberg, J.; Ek, A.R.; Burk, T.E.; Ter-Mikaelian, M.; Agren, G.I.; Oliver, C.D.; Puttonen, P. Process-based models for forest ecosystem management: Current state of the art and challenges for practical implementation. *Tree Physiol.* **2000**, *20*, 289–298. [[CrossRef](#)] [[PubMed](#)]
- Yuan, W.P.; Liu, S.G.; Zhou, G.S.; Zhou, G.Y.; Tieszen, L.L.; Baldocchi, D.; Bernhofer, C.; Gholz, H.; Goldstein, A.H.; Goulden, M.L.; et al. Deriving a light use efficiency model from eddy covariance flux data for predicting daily gross primary production across biomes. *Agric. For. Meteorol.* **2007**, *1433*, 189–207. [[CrossRef](#)]
- Yuan, W.; Cai, W.; Xia, J.; Chen, J.; Liu, S.; Dong, W.; Merbold, L.; Law, B.; Arain, A.; Beringer, J.; et al. Global comparison of light use efficiency models for simulating terrestrial vegetation gross primary production based on the LaThuile database. *Agric. For. Meteorol.* **2014**, *192–193*, 108–120. [[CrossRef](#)]
- Cai, W.W.; Yuan, W.P.; Liang, S.L.; Zhang, X.T.; Dong, W.J.; Xia, J.Z.; Fu, Y.; Chen, Y.; Liu, D.; Zhang, Q. Improved estimations of gross primary production using satellite-derived photosynthetically active radiation. *J. Geophys. Res. Biosci.* **2014**, *119*. [[CrossRef](#)]
- Lacointe, A. Carbon allocation among tree organs. A review of basic processes and representation in functional-structural tree models. *Ann. For. Sci.* **2000**, *57*, 521–533. [[CrossRef](#)]
- Sievänen, R.; Nikinmaa, E.; Nygren, P.; Ozier-Lafontaine, H.; Perttunen, J.; Hakula, H. Components of functional-structural tree models. *Ann. For. Sci.* **2000**, *57*, 399–412. [[CrossRef](#)]
- Lu, M.; Nygren, P.; Perttunen, J.; Pallardy, S.G.; Larsen, D.R. Application of the functional-structural tree model LIGNUM to growth simulation of short-rotation eastern cottonwood. *Silva Fenn.* **2011**, *45*, 431–474. [[CrossRef](#)]
- Nikinmaa, E.; Sievänen, R.; Hölttä, T. Dynamics of leaf gas exchange, xylem and phloem transport, water potential and carbohydrate concentration in a realistic 3-D model tree crown. *Ann. Bot.* **2014**, *114*, 653–666. [[CrossRef](#)] [[PubMed](#)]
- Monteith, J.; Moss, C.J. Climate and the efficiency of crop production in Britain. *Philos. Trans. R. Soc. Lond.* **1977**, *281*, 277–294. [[CrossRef](#)]
- Trumbore, S. Carbon respired by terrestrial ecosystems—recent progress and challenges. *Glob. Chang. Biol.* **2006**, *12*, 141–153. [[CrossRef](#)]
- Mitchell, S.; Beven, K.; Freer, J. Multiple sources of predictive uncertainty in modeled estimates of net ecosystem CO₂ exchange. *Ecol. Model.* **2009**, *220*, 3259–3270. [[CrossRef](#)]

20. Yuan, F.; Arain, M.A.; Barr, A.G.; Black, T.A.; Bourque, C.P.A.; Coursolle, C.; Margolis, H.A.; Mccaughey, J.H.; Wofsy, S.C. Modeling analysis of primary controls on net ecosystem productivity of seven boreal and temperate coniferous forests across a continental transect. *Glob. Chang. Biol.* **2008**, *14*, 1765–1784. [[CrossRef](#)]
21. Xu, B.; Yang, Y.; Li, P.; Shen, H.; Fang, J. Global patterns of ecosystem carbon flux in forests: A biometric databased synthesis. *Glob. Biogeochem. Cycles* **2014**, *28*, 962–973. [[CrossRef](#)]
22. Zobitz, J.M.; Moore, D.J.P.; Sacks, W.J.; Monson, R.K.; Bowling, D.R.; Schimel, D.S. Integration of process-based soil respiration models with whole-ecosystem CO₂ measurements. *Ecosystems* **2008**, *11*, 250–269. [[CrossRef](#)]
23. Lloyd, J.; Taylor, J.A. On the temperature dependence of soil respiration. *Funct. Ecol.* **1994**, *8*, 315–323. [[CrossRef](#)]
24. Kirschbaum, M.U.F. Soil respiration under prolonged soil warming: Are rate reductions caused by acclimation or substrate loss? *Glob. Chang. Biol.* **2004**, *10*, 1870–1877. [[CrossRef](#)]
25. Xu, J.; Chen, J.; Brosofske, K.; Li, Q.; Weintraub, M.; Henderson, R.; Wilske, B.; John, R.; Jensen, R.; Li, H.; et al. Influence of timber harvesting alternatives on forest soil respiration and its biophysical regulatory factors over a 5-year period in the Missouri Ozarks. *Ecosystems* **2011**, *14*, 1310–1327. [[CrossRef](#)]
26. Schimel, J.P.; Weintraub, M.N. Soil organic matter does not break itself down the implications of exoenzyme activity on microbial carbon and nitrogen limitation in soil: A theoretical model. *Soil Biol. Biochem.* **2003**, *35*, 549–563. [[CrossRef](#)]
27. Tian, H.; Lu, C.; Yang, J.; Banger, K.; Huntzinger, D.N.; Schwalm, C.R.; Michalak, A.M.; Cook, R.; Ciais, P.; Hayes, D.; et al. Global patterns and controls of soil organic carbon dynamics as simulated by multiple terrestrial biosphere models: Current status and future directions. *Glob. Biogeochem. Cycles* **2015**, *29*, 775–792. [[CrossRef](#)] [[PubMed](#)]
28. Chapin, S., III; McFarland, J.; David McGuire, A.; Euskirchen, E.S.; Ruess, R.W.; Kielland, K. The changing global carbon cycle: Linking plant-soil carbon dynamics to global consequences. *J. Ecol.* **2009**, *97*, 840–850.
29. Hibbard, K.A.; Law, B.E.; Reichstein, M.; Sulzman, J. An analysis of soil respiration across northern hemisphere temperate ecosystems. *Biogeochemistry* **2005**, *73*, 29–70. [[CrossRef](#)]
30. Knohl, A.; Schulze, E.D.; Kolle, O.; Buchmann, N. Large carbon uptake by an unmanaged 250-year-old deciduous forest in Central Germany. *Agric. For. Meteorol.* **2003**, *118*, 151–167. [[CrossRef](#)]
31. Facelli, J.M.; Pickett, S.T.A. Plant litter: Its dynamics and effects on plant community structure. *Bot. Rev.* **1991**, *57*, 1–32. [[CrossRef](#)]
32. Richardson, A.D.; Anderson, R.S.; Arain, M.A.; Barr, A.G.; Bohrer, G.; Chen, G.; Chen, J.M.; Ciais, P.; Davis, K.J.; Desai, A.R.; et al. Terrestrial biosphere models need better representation of vegetation phenology: Results from the North American Carbon Program Site Synthesis. *Glob. Chang. Biol.* **2012**, *18*, 566–584. [[CrossRef](#)]
33. Kucharik, C.J.; Barford, C.C.; El Maayar, M.; Wofsy, S.C.; Monson, R.K.; Baldocchi, D.D. A multiyear evaluation of a Dynamic Global Vegetation Model at three AmeriFlux forest sites: Vegetation structure, phenology, soil temperature, and CO₂ and H₂O vapor exchange. *Ecol. Model.* **2006**, *196*, 1–31. [[CrossRef](#)]
34. Ryu, S.R.; Chen, J.; Noormets, A.; Bresee, M.K.; Ollinger, S.V. Comparisons between PnET-Day and eddy covariance based gross ecosystem production in two Northern Wisconsin forests. *Agric. For. Meteorol.* **2008**, *148*, 247–256. [[CrossRef](#)]
35. Richardson, A.D.; Keenan, T.F.; Migliavacca, M.; Ryu, Y.; Sonnentag, O.; Toomey, M. Climate change, phenology, and phenological control of vegetation feedbacks to the climate system. *Agric. For. Meteorol.* **2013**, *169*, 156–173. [[CrossRef](#)]
36. Collalti, A.; Perugini, L.; Santini, M.; Chiti, T.; Nolè, A.; Matteucci, G.; Valentini, R. A process-based model to simulate growth in forests with complex structure: Evaluation and use of 3D-CMCC Forest Ecosystem Model in a deciduous forest in Central Italy. *Ecol. Model.* **2014**, *272*, 362–378. [[CrossRef](#)]
37. Collalti, A.; Marconi, S.; Ibrom, A.; Trotta, C.; Anav, A.; D'andrea, E.; Grünwald, T. Validation of 3D-CMCC Forest Ecosystem Model (v. 5.1) against eddy covariance data for 10 European forest sites. *Geosci. Model Dev.* **2016**, *9*, 479–504. [[CrossRef](#)]
38. Zhang, Y.; Li, C.; Zhou, X.; Moore, B. A simulation model linking crop growth and soil biogeochemistry for sustainable agriculture. *Ecol. Model.* **2002**, *151*, 75–108. [[CrossRef](#)]
39. FLUXNET-Fluxdata. Available online: <http://fluxnet.fluxdata.org> (accessed on 3 June 2014).

40. Papale, D.; Reichstein, M.; Aubinet, M.; Canfora, E.; Bernhofer, C.; Longdoz, B.; Kutsch, W.; Rambal, S.; Valentini, R.; Vesala, T.; et al. Towards a standardized processing of Net Ecosystem Exchange measured with eddy covariance technique: Algorithms and uncertainty estimation. *Biogeosciences* **2006**, *3*, 571–583. [[CrossRef](#)]
41. Mencuccini, M.; Bonosi, L. Leaf/sapwood area ratios in Scots pine show acclimation across Europe. *Can. J. For. Res.* **2001**, *31*, 442–456. [[CrossRef](#)]
42. Pilegaard, K.; Mikkelsen, T.N.; Beier, C.; Jensen, N.O.; Ambus, P.; Ro-Poulsen, H. Field Measurements of Atmosphere—Biosphere Interactions in a Danish Beech Forest. *Boreal Environ. Res.* **2003**, *8*, 315–333.
43. Landsberg, J.J.; Waring, R.H. A generalised model of forest productivity using simplified concepts of radiation-use efficiency, carbon balance and partitioning. *For. Ecol. Manag.* **1997**, *95*, 209–228. [[CrossRef](#)]
44. Molina, J.A.E.; Clapp, C.E.; Shaffer, M.J.; Chichester, F.W.; Larson, W.E. NCSOIL, A Model of Nitrogen and Carbon Transformations in Soil: Description, Calibration, and Behavior1. *Soil Sci. Soc. Am. J.* **1983**, *47*, 85. [[CrossRef](#)]
45. Sollins, P.; Homann, P.; Caldwell, B.A. Stabilization and destabilization of soil organic matter: Mechanisms and controls. *Geoderma* **1996**, *74*, 65–105. [[CrossRef](#)]
46. Tamon, F.; Kobayashi, K.; Li, C.; Yagi, K.; Hasegawa, T. Revising a Process-based Biogeochemistry Model (DNDC) to Simulate Methane Emission from Rice Paddy Fields under Various Residue Management and Fertilizer. *Glob. Chang.* **2008**, *14*, 382–402. [[CrossRef](#)]
47. Coleman, K.; Jenkinson, D.S. “ROTHC-26.3.” A Model for the Turnover of Carbon in Soils, No. November 1999. Available online: http://www.uni-kassel.de/~w_dec/Modellierung/wdec-rothc_manual.pdf (accessed on 1 October 2014).
48. Running, S.W.; Raymond Hunt, E., Jr. *Generalization of a Forest Ecosystem Process Model for Other Biomes, BIOME-BGC, and an Application for Global-Scale Models*; Academic Press, Inc.: London, UK, 1993; pp. 141–158.
49. Delpierre, N.; Dufréne, E.; Soudani, K.; Ulrich, E.; Cecchini, S.; Boé, J.; François, C. Modelling Interannual and Spatial Variability of Leaf Senescence for Three Deciduous Tree Species in France. *Agric. For. Meteorol.* **2009**, *149*, 938–948. [[CrossRef](#)]
50. Keel, S.G.; Schädel, C. Expanding leaves of mature deciduous forest trees rapidly become autotrophic. *Tree Physiol.* **2010**, *30*, 1253–1259. [[CrossRef](#)] [[PubMed](#)]
51. Chabot, B.F.; Hicks, D.J. The Ecology of Leaf Life Spans. *Ann. Rev. Ecol. Syst.* **1982**, *13*, 229–259. Available online: <http://www.jstor.org/stable/2097068> (accessed on 6 October 2015). [[CrossRef](#)]
52. Tilman, D. Constraints and tradeoffs: Toward a predictive theory of competition and succession. *Oikos* **1990**, *58*, 3–15. [[CrossRef](#)]
53. Waring, R.H.; Landsberg, J.J.; Williams, M. Net Primary Production of Forests: A Constant Fraction of Gross primary production? *Tree Physiol.* **1998**, *18*, 129–134. [[CrossRef](#)] [[PubMed](#)]
54. Praciak, A.; Pasiecznik, N.; Sheil, D.; van Heist, M.; Sassen, M.; Correia, C.S.; Dixon, C.; Fyson, G.; Rushford, K.; Teeling, C. *The CABI Encyclopedia of Forest Trees*; CABI: Oxfordshire, UK, 2013.
55. Kuzyakov, Y. Priming effects: Interactions between living and dead organic matter. *Soil Biol. Biochem.* **2010**, *42*, 1363–1371. [[CrossRef](#)]
56. Thornton, P. *Biome BGC Version 4.2: Theoretical Framework of Biome-BGC*; Technical Documentation: Missoula, MT, USA, 2010.
57. Liang, B.; Lehmann, J.; Sohi, S.P.; Thies, J.E.; O’Neill, B.; Trujillo, L.; Luizão, F.J. Black carbon affects the cycling of non-black carbon in soil. *Org. Geochem.* **2010**, *41*, 206–213. [[CrossRef](#)]
58. James, G.; Witten, D.; Hastie, T.; Tibshirani, R. *An Introduction to Statistical Learning*; Springer: New York, NY, USA, 2013; Volume 6.
59. Walther, B.A.; Moore, J.L. The concepts of bias, precision and accuracy, and their use in testing the performance of species richness estimators, with a literature review of estimator performance. *Ecography* **2005**, *28*, 815–829. [[CrossRef](#)]
60. Keenan, T.F.; Baker, I.; Barr, A.; Ciais, P.; Davis, K.; Dietze, M.; Dragoni, D.; Gough, C.M.; Grant, R.; Hollinger, D.; et al. Terrestrial Biosphere Model Performance for Inter-Annual Variability of Land-Atmosphere CO₂ Exchange. *Glob. Chang. Biol.* **2012**, *18*, 1971–1987. [[CrossRef](#)]
61. Taylor, K. Summarizing multiple aspects of model performance in a single diagram. *J. Geophys. Res.* **2001**, *106*, 7183–7192. [[CrossRef](#)]

62. Booth, B.B.B.; Jones, C.D.; Collins, M.; Totterdell, I.J.; Cox, P.M.; Sitch, S.; Huntingford, C.; Betts, R.A.; Harris, G.R.; Lloyd, J. High sensitivity of future global warming to land carbon cycle processes. *Environ. Res. Lett.* **2012**, *7*, 024002. [[CrossRef](#)]
63. Zhao, Y.; Ciais, P.; Peylin, P.; Viovy, N.; Longdoz, B.; Bonnefond, J.M.; Rambal, S.; Klumpp, K.; Olliso, A.; Cellier, P.; et al. How errors on meteorological variables impact simulated ecosystem fluxes: A case study for six French sites. *Biogeosciences* **2012**, *9*, 2537–2564. [[CrossRef](#)]
64. Doblas-Miranda, E.; Martínez-Vilalta, J.; Lloret, F.; Álvarez, A.; Ávila, A.; Bonet, F.J.; Ferrandis, P. Reassessing global change research priorities in mediterranean terrestrial ecosystems: How far have we come and where do we go from here? *Glob. Ecol. Biogeogr.* **2015**, *24*, 25–43. [[CrossRef](#)]
65. Huber-Sanwald, E.; Huenneke, L.F.; Jackson, R.B.; Kinzig, A.; Leemans, R.; Lodge, D.M.; Walker, B.H. Biodiversity global biodiversity scenarios for the year 2100. *Science* **2000**, *287*, 1770–1774.
66. Muraoka, H.; Koizumi, H. Photosynthetic and structural characteristics of canopy and shrub trees in a cool-temperate deciduous broadleaved forest: Implication to the ecosystem carbon gain. *Agric. For. Meteorol.* **2005**, *134*, 39–59. [[CrossRef](#)]
67. Kramer, K.; Leinonen, I.; Bartelink, H.H.; Berbigier, P.; Borghetti, M.; Cienciala, E.; Dolman, A.J.; Froer, O.; Gracia, C.; Granier, A.; et al. Evaluation of Six Process-Based Forest Growth Models Using Eddy-Covariance Measurements of CO₂ and H₂O Fluxes at Six Forest Sites in Europe. *Glob. Chang. Biol.* **2002**, 213–230. [[CrossRef](#)]
68. Keenan, T.; Garcia, R.; Sabate, S.; Gracia, C.A. Process Based Forest Modelling: A Thorough Validation and Future Prospects for Mediterranean Forests in a Changing World. *Cuad. Soc. Esp. Cienc. For.* **2007**, *92*, 81–92.
69. Zhou, W.; Hui, D.; Shen, W. Effects of soil moisture on the temperature sensitivity of soil heterotrophic respiration: A laboratory incubation study. *PLoS ONE* **2014**, *9*, e92531. [[CrossRef](#)] [[PubMed](#)]
70. Hollinger, D.Y.; Richardson, A.D. Uncertainty in eddy covariance measurements and its application to physiological models. *Tree Physiol.* **2005**, *25*, 873–885. [[CrossRef](#)] [[PubMed](#)]
71. DeLucia, E.H.; Drake, J.E.; Thomas, R.B.; Gonzalez-Meler, M.A. Forest carbon use efficiency: Is respiration a constant fraction of gross primary production? *Glob. Chang. Biol.* **2007**, *13*, 1157–1167. [[CrossRef](#)]
72. Ito, A.; Oikawa, T. Global mapping of terrestrial primary productivity and light-use efficiency with a process-based model. In *Global Environmental Change in the Ocean and on Land*; TERRAPUB: Tokyo, Japan, 2004; pp. 343–358.
73. Sabaté, S.; Gracia, C.A.; Sánchez, A. Likely effects of climate change on growth of *Quercus ilex*, *Pinus halepensis*, *Pinus pinaster*, *Pinus sylvestris* and *Fagus sylvatica* forests in the Mediterranean region. *For. Ecol. Manag.* **2002**, *162*, 23–37. [[CrossRef](#)]
74. Chiti, T.; Papale, D.; Smith, P.; Dalmonech, D.; Matteucci, G.; Yeluripati, J.; Valentini, R. Predicting changes in soil organic carbon in mediterranean and alpine forests during the Kyoto Protocol commitment periods using the CENTURY model. *Soil Use Manag.* **2010**, *26*, 475–484. [[CrossRef](#)]
75. Dickinson, C.H. *Biology of Plant Litter Decomposition*; Elsevier: Amsterdam, The Netherlands, 2012; Volume 2.
76. Yang, Y.; Zhu, Q.; Peng, C.; Wang, H.; Xue, W.; Lin, G.; Li, S. A novel approach for modelling vegetation distributions and analysing vegetation sensitivity through trait-climate relationships in China. *Sci. Rep.* **2016**, *6*. [[CrossRef](#)] [[PubMed](#)]
77. Yang, Y.; Zhu, Q.; Peng, C.; Wang, H.; Chen, H. From plant functional types to plant functional traits: A new paradigm in modelling global vegetation dynamics. *Prog. Phys. Geogr.* **2015**, *39*, 514–535. [[CrossRef](#)]
78. Messier, J.; McGill, B.J.; Lechowicz, M.J. How do traits vary across ecological scales? A case for trait-based ecology. *Ecol. Lett.* **2010**, *13*, 838–848. [[CrossRef](#)] [[PubMed](#)]
79. Claussen, M.; Mysak, L.A.; Weaver, A.J.; Crucifix, M.; Fichet, T.; Loutre, M.-F.; Weber, S.L.; Alcamo, J.; Alexeev, V.A.; Berger, A.; et al. Earth System Models of Intermediate Complexity: Closing the Gap in the Spectrum of Climate System Models. *Clim. Dyn.* **2002**, *18*, 579–586.
80. Lavorel, S.; Díaz, S.; Cornelissen, J.H.C.; Garnier, E.; Harrison, S.P.; McIntyre, S.; Urcelay, C. Plant functional types: Are we getting any closer to the Holy Grail? In *Terrestrial Ecosystems in a Changing World*; Springer: Berlin/Heidelberg, Germany, 2007; pp. 149–164.
81. Schaber, J.; Badeck, F.W. Physiology-Based Phenology Models for Forest Tree Species in Germany. *Int. J. Biometeorol.* **2003**, *47*, 193–201. [[CrossRef](#)] [[PubMed](#)]

82. Linkosalo, T.; Lappalainen, H.K.; Hari, P. A comparison of phenological models of leaf bud burst and flowering of boreal trees using independent observations. *Tree Physiol.* **2008**, *28*, 1873–1882. [[CrossRef](#)] [[PubMed](#)]
83. Uusitalo, L.; Lehikoinen, A.; Helle, I.; Myrberg, K. An overview of methods to evaluate uncertainty of deterministic models in decision support. *Environ. Model. Softw.* **2015**, *63*, 24–31. [[CrossRef](#)]
84. Wythers, K.R.; Reich, P.B.; Bradford, J.B. Incorporating temperature-sensitive Q10 and foliar respiration acclimation algorithms modifies modeled ecosystem responses to global change. *J. Geophys. Res. Biogeosci.* **2013**, *118*, 77–90. [[CrossRef](#)]
85. Ryan, M.G.; Gower, S.T.; Hubbard, R.M. Woody Tissue Maintenance Respiration of Four Conifers in Contrasting Climates. *Oecologia* **1995**, *101*, 133–140. [[CrossRef](#)] [[PubMed](#)]
86. Valentini, R.; Matteucci, G.; Dolman, A.J.; Schulze, E.D.; Rebmann, C.; Moors, E.J.; Granier, A.; Lindroth, A. Respiration as the main determinant of Carbon balance in European forests. *Nature* **2000**, *404*, 851–865. [[CrossRef](#)] [[PubMed](#)]



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