

LPJmL4 – a dynamic global vegetation model with managed land: Part I – Model description

Sibyll Schaphoff¹, Werner von Bloh¹, Anja Rammig², Kirsten Thonicke¹,
Hester Biemans³, Matthias Forkel⁴, Dieter Gerten^{1,5}, Jens Heinke¹,
Jonas Jägermeyr¹, Jürgen Knauer⁶, Fanny Langerwisch¹, Wolfgang Lucht^{1,5},
Christoph Müller¹, Susanne Rolinski¹, and Katharina Waha^{1,7}

¹Potsdam Institute for Climate Impact Research, PO Box 60 12 03, 14412 Potsdam, Germany

²Technical University of Munich, School of Life Sciences Weihenstephan, 85354 Freising, Germany

³Alterra, Wageningen University & Research, PO Box 47, 6700 AA Wageningen, the Netherlands

⁴TU Wien, Climate and Environmental Remote Sensing Group, Department of Geodesy and Geoinformation, Gusshausstraße 25-29, 1040 Wien, Austria

⁵Humboldt Universität zu Berlin, Department of Geography, Unter den Linden 6, 10099 Berlin, Germany

⁶Max Planck Institute for Biogeochemistry, Hans-Knöll-Str. 10, 07745 Jena, Germany

⁷CSIRO Agriculture & Food, 306 Carmody Rd, St Lucia QLD 4067, Australia

Correspondence to: Sibyll.Schaphoff@pik-potsdam.de

Abstract.

This paper provides a comprehensive description of the newest version of the Dynamic Global Vegetation Model with managed Land, LPJmL4. This model simulates - internally consistently - the growth and productivity of both natural and agricultural vegetation in coherently linked through their water, carbon and energy fluxes. These features render LPJmL4 suitable for assessing a broad range of feedbacks within, and impacts upon, the terrestrial biosphere as increasingly shaped by human activities such as climate change and land-use change. Here we describe the core model structure including recently developed modules now unified in LPJmL4. Thereby, we also review LPJmL model developments and evaluations in the field of permafrost, human and ecological water demand and improved representation of crop types. We summarize and discuss LPJmL model applications dealing with impacts of historical and future environmental change on the terrestrial biosphere at regional and global scale and provide a comprehensive overview over LPJmL publications since the first model description in 2007. To demonstrate the main features of the LPJmL4 model, we display reference simulation results for key processes such as the current global distribution of natural and managed ecosystems, their productivities, and associated water fluxes. A thorough evaluation of the model is provided in a companion paper. By making the model source code freely available at <https://gitlab.pik-potsdam.de/lpjml/LPJmL>, we hope to stimulate the application and further development of LPJmL4 across scientific communities, not least in support of major activities such as the IPCC and SDG process.

20 1 Introduction

The terrestrial biosphere, a highly dynamic key component of the Earth system, is undergoing significant and widespread transformations induced by human activities such as climate and land-use change. Humans have by now transformed about 40% of the terrestrial ice-free land surface into land used for agriculture and urban settlements (Ellis et al., 2010), thus pushing the planetary dynamics beyond boundaries that have been characteristic for the past ca. 12,000 years (Rockström et al., 2009). These interventions put at risk important functions of the biosphere such as the provisioning of floral and faunal biodiversity (Vörösmarty et al., 2010), the terrestrial carbon sink (Le Quéré et al., 2015) and the provisioning of accessible freshwater (Vörösmarty et al., 2010). Understanding and modelling the current and potential future dynamics of the Earth system thus renders it necessary to consider human activities as an integral part while representing the major dynamics of the biosphere in a spatio-temporally explicit and process-based manner, accounting for the feedbacks between vegetation, global carbon and water cycling, and the atmosphere. This would also allow numerical evaluation of potential implementation pathways for the United Nation's Sustainable Development Goals (SDGs – <https://sustainabledevelopment.un.org>) and their impacts on the terrestrial environment, complementing the important role that dynamic biosphere models have played in the United Nation's scientific assessment reports on climate change published by the United Nation's Intergovernmental Panel on Climate Change (IPCC, 2014).

By combining core features of global biogeographical and biogeochemical models developed in the 1990s, Dynamic Global Vegetation Models (DGVMs) emerged as the main tool to simulate processes underlying the dynamics of natural vegetation types (growth, mortality, resource competition, disturbances such as wildfires) and the associated carbon and water fluxes (Cramer et al., 2001; Prentice et al., 2007; Sitch et al., 2008; Friend et al., 2014). In light of the strengthening human interferences, DGVMs were further developed to integrate additional processes that are relevant to the original research quest of studying biogeography and biogeochemical cycles under climate change (Canadell et al., 2007). This includes the incorporation of human land-use and the simulation of agricultural production systems (Bondeau et al., 2007; Lindeskog et al., 2013), nutrient limitation (Zaehle et al., 2010; Smith et al., 2014), as well as hydrological modules and river routing schemes (Gerten et al., 2004; Rost et al., 2008). Knowledge derived from models that are designed to cover aspects of the earth system other than terrestrial vegetation and the carbon cycle, such as models of the global water balance, could evidentially improve the DGVMs' ability to also evaluate model performance for processes (e.g. river discharge) that are closely connected to the simulated vegetation and carbon cycle dynamics (Bondeau et al., 2007; Smith et al., 2014). The development towards more comprehensive models of Earth's land surface offers new possibilities for cross-disciplinary research.

DGVMs as land components of Earth system models still show large uncertainties about the terrestrial carbon (C) balance under future climate change (Friedlingstein et al., 2013). This uncertainty

partly results from differences in the simulation of soil and vegetation C residence times (Carvalhais et al., 2014; Friend et al., 2014). The time that C resides in an ecosystem is thereby strongly affected by the simulated processes of vegetation dynamics (Ahlström et al., 2015). These examples highlight the need to continuously improve process representations in DGVMs in order to reduce the uncertainty in projected ecosystem functioning and services under future climate change. This requires however, that model developments in specific fields or improvements for certain processes are synthesized and integrated into a unified, internally consistent model version.

The Lund-Potsdam-Jena DGVM for managed land (LPJmL, Bondeau et al. (2007)) originates from a former version of the model described by Sitch et al. (2003) and simulates growth and geographical distribution of natural "plant functional types" (PFTs), "crop functional types" (CFTs) and associated biogeochemical processes (mainly carbon cycling). Recent developments focused on an improved energy balance model able to estimate permafrost dynamics based on a vertical soil carbon distribution scheme and a new soil hydrological scheme (Schaphoff et al., 2013). Also, a new process-based fire module (SPITFIRE) was implemented that allows for detailed simulation of fire ignition, spread and effects to estimate fire impacts and emissions (Thonicke et al., 2010). An updated phenology scheme was developed, which now takes phenology limitations arising from low temperatures, limited light and drought into account (Forkel et al., 2014). Further model developments encompass the parallelization of the model to efficiently simulate river routing (Von Bloh et al., 2010) and the implementation of irrigation scheme (Rost et al., 2008), recently updated with a mechanistic representation of the three major irrigation systems (Jägermeyr et al., 2015). Biemans et al. (2011) implemented reservoir operations and irrigation extraction and evaluated the impact on river discharge. Other developments focused on a newly formulated implementation of different cropping systems in sub-Saharan Africa (Waha et al., 2013), Mediterranean agricultural plant types (Fader et al., 2015) and bioenergy crops such as sugarcane (Lapola et al., 2009), fast-growing grasses and bioenergy trees (Beringer et al., 2011). With these implementations, the potential of bioenergy production under future land-use, population and climate development could be extensively investigated (Haberl et al., 2011; Popp et al., 2011; Humpenöder et al., 2014). All developments, the core model structure and recently developed modules of DGVM LPJmL version 4.0 (in the following referred to as LPJmL4) will be described in section 2 in more detail. We show that the model in its present form allows for consistent and joint quantification of climate and land-use change impacts on the terrestrial biosphere, the water cycle, the carbon cycle, and on agricultural production (a systematic evaluation can be found in Part II of this paper). To give an overview of recent developments and applications of LPJmL4, we present:

1. A comprehensive description of the full model with all contributing developments since its original publication by Sitch et al. (2003) and Bondeau et al. (2007). We aim at consistently uniting all developments, including undocumented and already published developments, thus providing a comprehensive description of the full LPJmL4 model.

2. An overview over published LPJmL applications to review the improvement of process understanding.
3. A discussion of here presented standard LPJmL4 results that give an overview of simulated biogeochemical, hydrological and agricultural patterns at the global scale.

2 Model description

The original Lund-Potsdam-Jena (LPJ) DGVM was described in detail by Sitch et al. (2003). This description and the associated model evaluation focused on the modelling of growth and geographical distribution of natural "plant functional types" (PFTs) and associated biogeochemical processes (mainly carbon cycling). Building on the improved representation of the water balance (Gerten et al., 2004). Bondeau et al. (2007) introduced the representation of "crop functional types" (CFTs) and evaluated the role of agriculture for the terrestrial carbon balance in particular. This model is since then referred to as LPJmL (Lund-Potsdam-Jena managed Land) and provide the foundation for explicitly simulating agricultural production in a changing climate and for quantifying impacts of agricultural activities in assessments of the terrestrial carbon and water cycle.

Since, a number of further specific model developments and applications have been published, but a comprehensive model description of all developments and amendments is missing. The parts of LPJmL4 building on Bondeau et al. (2007) not only allow for quantifying changes in vegetation composition, the water cycle, the carbon cycle, and agricultural production, but also for explicitly simulating the dynamics and constraints within and among the modules, thereby providing a consistent and comprehensive representation of Earth's land surface processes. To demonstrate the interplay of all these model features in the new LPJmL4 version, the present paper documents the core model structure including equations and parameters from Sitch et al. (2003) and Bondeau et al. (2007) and all more recent code developments. SI-Fig. S1 provides a schematic overview of the model structure and Fig. 1 of the simulated carbon, water and energy fluxes. The following sections describe the model components: energy balance model and permafrost (2.1), plant physiology (2.2), plant functional (2.3) and crop functional types (2.4), soil litter and carbon pools (2.5), water balance (2.6) and land use (2.7).

2.1 Energy balance model and permafrost

The energy balance model includes the calculation of photosynthetic active radiation, daylength and potential evapotranspiration (2.1.1) and albedo (2.1.2). The permafrost module is based on a new calculation of the soil energy balance (2.1.3).

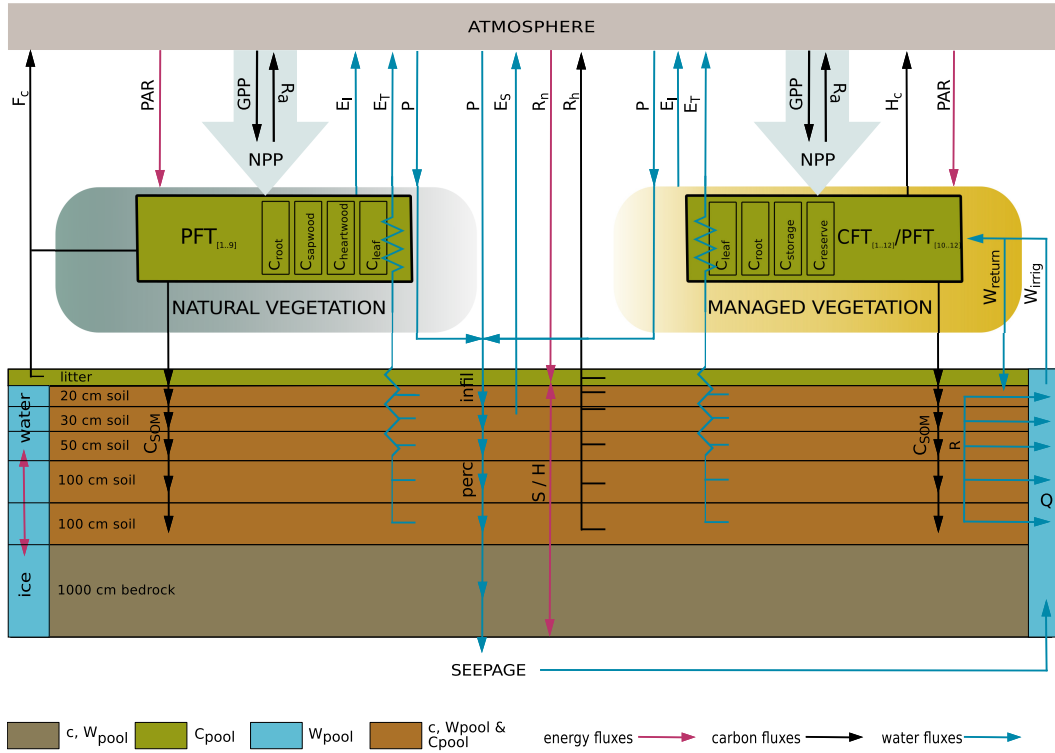


Figure 1. LPJmL4-scheme for Carbon, Water and Energy fluxes represented by the model.

C - carbon; W - water; S - sensible heat conduction; H - latent heat convection; c - energy conduction; R_n - net downward radiation (input); PAR - photosynthetic active radiation; E_i - interception; E_T - transpiration; E_S - evaporation; infil - infiltration; perc - percolation; P - precipitation (input); GPP - gross primary production; NPP - net primary production; R_a - autotrophic respiration; R_h - heterotrophic respiration; H_c - carbon harvested; F_c - carbon emitted by fire; SOM - soil organic matter; R - runoff; Q - discharge

2.1.1 Photosynthetic active radiation, daylength and potential evapotranspiration

Photosynthetic active radiation (PAR) is the primary energy force to photosynthesis (2.2.1) and, thus, to the whole carbon cycle. Total daily PAR in $\text{mol m}^{-2} \text{day}^{-1}$ is calculated as:

$$\text{PAR} = 0.5 \cdot c_q \cdot R_{s\text{day}}, \quad (1)$$

where $c_q = 4.6 \times 10^{-6}$ is the conversion factor from J to mol for solar radiation at 550 nm. Half of the daily incoming solar irradiance $R_{s\text{day}}$ is assumed to be PAR and atmospheric absorption to be the same for PAR and $R_{s\text{day}}$ (Prentice et al., 1993; Haxeltine and Prentice, 1996).

Similar to the role of PAR for the carbon cycle, potential evapotranspiration (PET) is the primary driver of the water cycle. The calculation of both PAR and PET follows the approach of Prentice et al. (1993), where the calculation of potential evapotranspiration [mm day^{-1}] is based on the theory of equilibrium evapotranspiration E_{eq} (Jarvis and McNaughton, 1986), given by:

$$E_{\text{eq}} = \frac{s}{s + \gamma} \cdot \frac{R_{n\text{day}}}{\lambda}, \quad (2)$$

where $R_{n_{\text{day}}}$ is daily surface net radiation [$\text{J m}^{-2} \text{day}^{-1}$] and λ is the latent heat of vaporization [J kg^{-1}] with a weak dependence on air temperature (T_{air} in $^{\circ}\text{C}$) derived from Monteith and Unsworth (1990, p. 376, Table A.3):

$$140 \quad \lambda = 2.495 \times 10^6 - 2380 \cdot T_{\text{air}} \quad (3)$$

s is the slope of the saturation vapour pressure curve [Pa K^{-1}], given by

$$s = 2.502 \times 10^6 \cdot \frac{\exp[17.269 \cdot T_{\text{air}} / (237.3 + T_{\text{air}})]}{(237.3 + T_{\text{air}})^2} \quad (4)$$

and γ is the psychrometric constant [Pa K^{-1}], given by

$$\gamma = 65.05 + 0.064 \cdot T_{\text{air}} \quad (5)$$

145 Following Priestley and Taylor (1972), PET (in mm) is subsequently calculated from E_{eq} as:

$$\text{PET} = \text{pt} \cdot E_{\text{eq}}, \quad (6)$$

where pt is the empirically derived Priestley-Taylor coefficient ($\text{pt} = 1.32$).

The terrestrial radiation balance is written as

$$R_n = (1 - \beta) \cdot R_s + R_l, \quad (7)$$

150 where R_n is net surface radiation; R_s is incoming solar irradiance (downward) at the surface and R_l the outgoing net long-wave radiation flux at the surface [all in W m^{-2}]; β is the short-wave reflection coefficient of the surface (albedo). The calculation of albedo depending on land surface conditions is described in section 2.1.2.

If not supplied directly as input variables to the model, the radiation terms R_s and R_l can be
155 computed for any day and latitude at given cloudiness levels (input), following Prentice et al. (1993).
 R_l can be approximated by a linear function of temperature and clear sky fraction:

$$R_l = (b + (1 - b) \cdot \text{ni}) \cdot (A - T_{\text{air}}), \quad (8)$$

where $b = 0.2$ and $A = 107$ are empirical constants. T_{air} is the mean daily air temperature in $^{\circ}\text{C}$, i.e. any effects of diurnal temperature variations are ignored. The proportion of bright sky (ni) is defined
160 by $\text{ni} = 1 - \text{cloudiness}$. The net outgoing daytime long-wave flux $R_{l_{\text{n}_{\text{day}}}}$ is obtained by multiplying with the length of the day in seconds:

$$R_{l_{\text{n}_{\text{day}}}} = R_l \cdot \text{daylength} \cdot 3600 \quad (9)$$

Instantaneous solar irradiance at the surface is computed from the solar constant, accounting for ni and the angular distance between the sun's rays and the local vertical (z):

$$165 \quad R_s = (c + d \cdot \text{ni}) \cdot Q_0 \cdot \cos(z) \quad (10)$$

where $c = 0.25$ and $d = 0.5$ are empirical constants that together represent the clear-sky transmittivity (0.75). Q_0 is the solar irradiance at day i , accounting for the variation of earth's distance to the sun:

$$Q_0 = Q_{00} \cdot (1 + 2 \cdot 0.01675 \cdot \cos(2 \cdot \pi \cdot i / 365)), \quad (11)$$

170 where Q_{00} is the solar constant with 1360 W m^{-2} . The solar zenith angle (z) correction of R_s is computed from the solar declination (δ , i.e. the angle between the orbital plane and the Earth's equatorial plane), which varies between $+23.4^\circ$ in northern hemisphere midsummer and -23.4° in northern hemisphere midwinter, the latitude (lat, in radians) and the hour angle h , i.e. the fraction of $2 \cdot \pi$ (in radians) which the earth has turned since the local solar noon.

$$175 \quad \cos(z) = \sin(\text{lat}) \cdot \sin(\delta) + \cos(\text{lat}) \cdot \cos(\delta) \cdot \cos(h) \quad (12)$$

with

$$\delta = -23.4 \cdot \pi / 180 \cdot \cos(2 \cdot \pi \cdot (i + 10) / 365) \quad (13)$$

To obtain the $R_{s\text{day}}$, eq. (10) needs to be integrated from sunrise to sunset, i.e. from $-h_{1/2}$ to $h_{1/2}$, where $h_{1/2}$ is the half-day length in angular units, computed as:

$$180 \quad h_{1/2} = \arccos\left(-\frac{\sin(\text{lat}) \cdot \sin(\delta)}{\cos(\text{lat}) \cdot \cos(\delta)}\right) \quad (14)$$

thus

$$\begin{aligned} R_{s\text{day}} &= (c + d \cdot \text{ni}) \cdot Q_0 \cdot (\sin(\text{lat}) \cdot \sin(\delta) \cdot h_{1/2} \\ &\quad + \cos(\text{lat}) \cdot \cos(\delta) \cdot h_{1/2}) \end{aligned} \quad (15)$$

The duration of sunshine of a single day (daylength in hours) is computed as:

$$185 \quad \text{daylength} = 24 \cdot \frac{h_{1/2}}{\pi} \quad (16)$$

2.1.2 Albedo

Albedo (β), the average reflectivity of the grid cell, was first implemented by Strengers et al. (2010) and later improved by considering several drivers of phenology as in Forkel et al. (2014).

$$\beta = \sum_{\text{PFT}=1}^{n_{\text{PFT}}} \beta_{\text{PFT}} \cdot \text{FPC}_{\text{PFT}} + F_{\text{bare}} \cdot (F_{\text{snow}} \cdot \beta_{\text{snow}} + (1 - F_{\text{snow}}) \cdot \beta_{\text{soil}}) \quad (17)$$

190 β depends on land surface condition and is based on a combination of defined albedo values for bare soil ($\beta_{\text{soil}} = 0.3$), snow ($\beta_{\text{snow}} = 0.7$ average value taken from Liang et al. (2005); Malik et al. (2012)) and plant compartments specific albedo values, where vegetation albedo (β_{PFT}) is simulated as the albedo of each existing PFT (β_{PFT}). FPC_{PFT} is the foliage projective cover of the respective PFT (see eq. 57). Parameters ($\beta_{\text{leaf,PFT}}$) were taken as suggested by Strugnelli et al. (2001) (see
195 SI-Table S5). Parameters $\beta_{\text{stem,PFT}}$ and $\beta_{\text{litter,PFT}}$ were obtained from Forkel et al. (2014) who optimized these parameters by using MODIS albedo time series. F_{snow} and F_{bare} are the snow coverage and the fraction of bare soil, respectively (Strengers et al., 2010).

2.1.3 Soil energy balance

The newly implemented calculation of the soil energy balance as described in Schaphoff et al. (2013) marks a new development and differs markedly from previous implementations of permafrost modules in LPJ (Beer et al., 2007). Soil water dynamics are computed daily (see section 2.6). The soil column is divided into five hydrological active layers of 0.2, 0.3, 0.5, 1 and 1 m depth (Δz) summing to 3 m (see section 2.6.1 and Fig. 1). Soil temperatures (T_{soil} in $^{\circ}\text{C}$) for these layers are computed with an energy balance model, including one-dimensional heat conduction and convection of latent heat. Freezing and thawing has been added to better account for soil ice dynamics. For a thermal buffer we assume an additional layer of 10 m thickness, which is only thermally and not hydrologically active. Soil parameters for thermal diffusivity ($\text{m}^2 \text{s}^{-1}$) at wilting point, at 15% of water holding capacity, and at field capacity and for thermal conductivity ($\text{W m}^{-1} \text{K}^{-1}$) at wilting point, and at saturation (for water and ice) are derived for each grid cell using soil texture from the Harmonized World Soil Database (HWSD) version (Nachtergaele et al., 2008). Relationships between texture and thermal properties are taken from Lawrence and Slater (2008). The one-dimensional heat conduction equation is:

$$\frac{\partial T_{\text{soil}}}{\partial t} = \alpha \cdot \frac{\partial^2 T_{\text{soil}}}{\partial z^2}, \quad (18)$$

where $\alpha = \lambda/c$ is thermal diffusivity, λ thermal conductivity, c heat capacity [in $\text{J m}^{-3} \text{K}^{-1}$]. T_{soil} at position z and time t is solved in its finite difference form following Bayazitoglu and Özışık (1988):

$$\frac{T_{\text{soil}}(t+1, l) - T_{\text{soil}}(t, l)}{\Delta t} = \alpha \cdot \frac{T_{\text{soil}}(t, l-1) + T_{\text{soil}}(t, l+1) - 2T_{\text{soil}}(t, l)}{(\Delta z)^2} \quad (19)$$

for soil layers l , including a snow layer, and time step t with the following boundary conditions:

$$T_{\text{soil}}(t=1, l=1) = T_{\text{air}}, \quad (20)$$

$$T_{\text{soil}}(t, l=n_{\text{soil}}+1) = T_{\text{soil}}(t, l=n_{\text{soil}}), \quad (21)$$

where $n_{\text{soil}} = 6$ is the number of soil layers. We assume a heatflux of zero below the lowest soil layer, i.e. below 13 m depth. The largest possible, numerically still stable time step Δt is calculated depending on Δz and soil thermal diffusivity α (Bayazitoglu and Özışık, 1988), which gives the stability criterion (r) for the finite-difference solution:

$$r = \frac{\alpha \Delta t}{(\Delta z)^2}. \quad (22)$$

For numerical stability ($1 - 2r$) needs to be > 0 , so that $r \leq 0.5$ as Δz is given from soil depth and α can be calculated from soil properties. The maximum stable Δt can be calculated:

$$\Delta t \leq \frac{(\Delta z)^2}{2 \cdot \alpha} \quad (23)$$

and therefore eq. (19) becomes:

$$T_{\text{soil}(t+1,l)} = r \cdot (T_{\text{soil}(t,l-1)} + T_{\text{soil}(t,l+1)} + (1 - 2r)T_{\text{soil}(t,l)}) \quad (24)$$

For the diurnal temperature range after Parton and Logan (1981), at least 4 time steps per day are calculated and the maximum number of time steps is set to 40 per day. Heat capacity (c) of the soil is calculated as the sum of the volumetric-specific heat capacities [in $\text{J m}^{-3} \text{K}^{-1}$] of soil minerals (c_{min}), soil water content (c_{water}) and soil ice content (c_{ice}) and their corresponding shares (m , in m^3) of the soil bucket:

$$c = c_{\text{min}} \cdot m_{\text{min}} + c_{\text{water}} \cdot m_{\text{water}} + c_{\text{ice}} \cdot m_{\text{ice}} \quad (25)$$

The heat capacity of air is neglected because of its comparatively low contribution to overall heat capacity. Thermal conductivity (λ) is calculated following Johansen (1977). Sensible and latent heat fluxes are calculated explicitly for the snow layer by assuming a constant snow density of 0.3 t m^{-3} and the resulting thermal diffusivity of $3.17 \times 10^{-7} \text{ m}^2 \text{ s}^{-1}$. Sublimation is assumed to be 0.1 mm day^{-1} , which corresponds to the lower end suggested by Gelfan et al. (2004).

The active layer thickness represents the depth of maximum thawing of the year. Freezing depth is calculated by assuming that the fraction of frozen water is congruent with the frozen soil bucket. The 0°C -isotherm within a layer is estimated by assuming a linear temperature gradient within the layer and this fraction of heat is assumed to be used for the thawing respectively freezing process. Temperature represents the amount of thermal energy available, whereas heat transport represents the movement of thermal energy into the soil by rain and melt water. Precipitation and percolation energy and the amount of energy which arises from the temperature difference between the temperature of the above layer (or the air temperature for the upper layer) and the temperature of the below layer, are assumed to be used for converting latent heat fluxes first. The residual energy is used to increase soil temperature. T_{soil} is initialized at the beginning of the spinup simulation by the mean annual air temperature.

2.2 Plant physiology

2.2.1 Photosynthesis

The LPJmL4 photosynthesis model is a 'big leaf' representation of the leaf-level photosynthesis model developed by Farquhar et al. (1980) and Farquhar and von Caemmerer (1982). These assumptions have been generalized by Collatz et al. (1991, 1992) for global modelling applications and for the stomatal response. The 'strong optimality' hypothesis (Haxeltine and Prentice, 1996; Prentice et al., 2000) is applied by assuming that Rubisco activity and the nitrogen content of leaves vary with canopy position and seasonally such as to maximize net assimilation at the leaf level. Most details are as in (Sitch et al., 2003) but a summary is provided in the following.

In LPJmL4, photosynthesis is simulated as a function of absorbed photosynthetically active radiation (APAR), temperature, daylength, and canopy conductance, for each PFT or CFT present in a grid cell and at daily time steps. APAR is calculated as the fraction of incoming net photosynthetically active radiation (PAR, see eq. (1)) that is absorbed by green vegetation (FAPAR):

$$\text{APAR}_{\text{PFT}} = \text{PAR} \cdot \text{FAPAR}_{\text{PFT}} \cdot \alpha_{\text{apPFT}} \quad (26)$$

α_{apPFT} is a scaling factor to scale leaf-level photosynthesis in LPJmL4 to stand level. The PFT-specific $\text{FAPAR}_{\text{PFT}}$ is calculated as follows:

$$\text{FAPAR}_{\text{PFT}} = \text{FPC}_{\text{PFT}} \cdot \left((\text{phen}_{\text{PFT}} - F_{\text{SnowGC}}) \cdot (1 - \beta_{\text{leaf,PFT}}) - (1 - \text{phen}_{\text{PFT}}) \cdot c_{\text{fstem}} \cdot \beta_{\text{stem,PFT}} \right), \quad (27)$$

where phen_{PFT} is the daily phenological status (ranging between 0 and 1) representing the fraction of full leaf coverage currently attained by the PFT, reduced by the green-leaf albedo $\beta_{\text{leaf,PFT}}$, the stem albedo $\beta_{\text{stem,PFT}}$ (for trees), and F_{SnowGC} is the fraction of snow in the green canopy. $c_{\text{fstem}} = 0.7$ is the masking of the ground by stems and branches without leaves (Strengers et al., 2010).

Based on this, gross photosynthesis rate A_{gd} is computed as the minimum of two functions (details in Haxeltine and Prentice (1996)):

1. The light-limited photosynthesis rate J_E ($\text{mol C m}^{-2} \text{ hour}^{-1}$)

$$J_E = C_1 \cdot \frac{\text{APAR}}{\text{daylength}}, \quad (28)$$

where for C3-Photosynthesis

$$C_1 = \alpha_{C3} \cdot T_{\text{stress}} \cdot \left(\frac{p_i - \Gamma_*}{p_i + 2 \cdot \Gamma_*} \right) \quad (29)$$

and for C4-Photosynthesis

$$C_1 = \alpha_{C4} \cdot T_{\text{stress}} \cdot \left(\frac{\lambda}{\lambda_{\text{maxC4}}} \right). \quad (30)$$

p_i is the leaf internal partial pressure of CO_2 given by $p_i = \lambda \cdot p_a$, where λ gives the ratio of the intercellular to ambient CO_2 concentration and p_a (in bar) is the ambient partial pressure of CO_2 , which reflects soil-plant water interaction (see eq. 40). T_{stress} is the PFT-specific temperature inhibition function, which limits photosynthesis at high and low temperatures. α_{C3} and α_{C4} are the intrinsic quantum efficiencies for CO_2 uptake in C3 and C4 plants respectively and Γ_* is the photorespiratory CO_2 compensation point.

$$\Gamma_* = \frac{[\text{O}_2]}{2 \cdot \tau}, \quad (31)$$

where $\tau = \tau_{25} \cdot q_{10,\tau}^{(T_{\text{air}} - 25) \cdot 0.1}$ is the specificity factor, it reflects the ability of Rubisco to discriminate between CO_2 and O_2 . $[\text{O}_2]$ is the partial pressure of O_2 (Pa) and τ_{25} is the τ value at 25°C and $q_{10,\tau}$ is the temperature sensitivity parameter.

2. The Rubisco-limited photosynthesis rate J_C ($\text{mol C m}^{-2} \text{ hour}^{-1}$).

$$J_C = C_2 \cdot V_m, \quad (32)$$

where V_m is the maximum Rubisco capacity (see eq. 35) and

$$C_2 = \frac{p_i - \Gamma_*}{p_i + K_C \left(1 + \frac{[\text{O}_2]}{K_O}\right)} \quad (33)$$

K_C and K_O represent the Michaelis-Menten constants for CO_2 and O_2 , respectively. Daily gross photosynthesis A_{gd} is then given by:

$$A_{\text{gd}} = \frac{\left(J_E + J_C - \sqrt{(J_E + J_C)^2 - 4 \cdot \theta \cdot J_E \cdot J_C}\right)}{2 \cdot \theta \cdot \text{daylength}} \quad (34)$$

The shape parameter θ describes the co-limitation of light and Rubisco activity (Haxeltine and Prentice, 1996). Subtracting leaf respiration (R_{leaf} , given in eq. 46), gives the daily net photosynthesis (A_{nd}), thus V_m is included in J_C and R_{leaf} . To calculate optimal A_{nd} , the zero point of the first derivative is calculated (i.e. $\partial A_{\text{nd}} / \partial V_m \equiv 0$). The thus derived maximum Rubisco capacity V_m is:

$$V_m = \frac{1}{b} \cdot \frac{C_1}{C_2} ((2 \cdot \theta - 1) \cdot s - (2 \cdot \theta \cdot s - C_2) \cdot \sigma) \cdot \text{APAR} \quad (35)$$

with

$$\sigma = \sqrt{1 - \frac{C_2 - 2}{C_2 - \theta s}} \text{ and } s = 24 / \text{daylength} \cdot b \quad (36)$$

The daily net daytime photosynthesis (A_{dt}) is given by subtracting dark respiration:

$$R_d = (1 - \text{daylength}/24) \cdot R_{\text{leaf}} \quad (37)$$

see eq. (46) for R_{leaf} and A_{dt} is

$$A_{\text{dt}} = A_{\text{nd}} - R_d \quad (38)$$

The photosynthesis rate can be related to canopy conductance (g_c in mm s^{-1}) through the CO_2 diffusion gradient between the intercellular air spaces and the atmosphere:

$$g_c = \frac{1.6 A_{\text{dt}}}{p_a \cdot (1 - \lambda)} + g_{\text{min}}, \quad (39)$$

where g_{min} [mm s^{-1}] is a PFT-specific minimum canopy conductance scaled by FPC that occurs due to processes other than photosynthesis. Combining both methods determining A_{dt} (eqs. 38, 39) gives:

$$0 = A_{\text{dt}} - A_{\text{dt}} = A_{\text{nd}} + (1 - \text{daylength}/24) \cdot R_{\text{leaf}} - p_a \cdot (g_c - g_{\text{min}}) \cdot (1 - \lambda) / 1.6 \quad (40)$$

This equation has to be solved for λ which is not possible analytically because of the occurrence of λ in A_{nd} in the second term of the equation. Therefore, a numerical bisection algorithm is used to solve the equation and obtain λ solving. The actual canopy conductance is calculated as a function of water stress depending on the soil moisture status (section 2.6.2) and thus the photosynthesis rate is related to actual canopy conductance. All parameter values are given in SI-Table S6.

2.2.2 Phenology

The phenology module of tree and grass PFTs is based on the growing season index (GSI) approach (Jolly et al., 2005). Thereby the continuous development of canopy greenness is modelled based on empirical relations to temperature, day length and drought conditions. The GSI approach was modified for its use in LPJmL (Forkel et al., 2014) so that it accounts for the limiting effects of cold temperature, light, water availability and heat stress on the daily phenology status phen_{PFT} :

$$\text{phen}_{\text{PFT}} = f_{\text{cold}} \cdot f_{\text{light}} \cdot f_{\text{water}} \cdot f_{\text{heat}} \quad (41)$$

Each limiting function can range between 0 (full limitation of leaf development) and 1 (no limitation of leaf development). The limiting functions are defined as logistic functions and depend also on the previous day's value:

$$f(x)_t = f(x)_{t-1} + (1/(1 + \exp(\text{sl}_x \cdot (x - b_x))) - f(x)_{t-1}) \cdot \tau_x, \quad (42)$$

where x is daily air temperature for the cold and heat stress-limiting functions f_{cold} and f_{heat} , respectively, and stands for, short-wave downward radiation in the light-limiting function f_{light} , and water availability for the water-limiting function f_{water} . The parameters b_x and sl_x are the inflection point and slope of the respective logistic function; τ_x is a change rate parameter that introduces a time-lagged response of the canopy development to the daily meteorological conditions. The empirical parameters were estimated by optimizing LPJmL simulations of FAPAR against 30 years of satellite-derived time series of FAPAR (Forkel et al., 2014).

2.2.3 Productivity

Autotrophic respiration

Autotrophic respiration is separated into carbon costs for maintenance and growth and is calculated as in Sitch et al. (2003). Maintenance respiration (R_x in $\text{gC m}^{-2} \text{ day}^{-1}$) depends on tissue-specific C:N ratios (for above- $\text{CN}_{\text{sapwood}}$ and below-ground tissues CN_{root}). It further depends on temperature (T), either air temperature (T_{air}) for above- and soil temperatures (T_{soil}) for below-ground tissues, on tissue biomass ($C_{\text{sapwood,ind}}$ resp. $C_{\text{root,ind}}$) and phenology (phen_{PFT} , see eq. (41)) and is calculated at a daily timestep as follows:

$$R_{\text{sapwood}} = P \cdot r_{\text{PFT}} \cdot k \cdot \frac{C_{\text{sapwood,ind}}}{\text{CN}_{\text{sapwood}}} \cdot g(T_{\text{air}}) \quad (43)$$

$$R_{\text{root}} = P \cdot r_{\text{PFT}} \cdot k \cdot \frac{C_{\text{root,ind}}}{\text{CN}_{\text{root}}} \cdot g(T_{\text{soil}}) \cdot \text{phen}_{\text{PFT}} \quad (44)$$

The respiration rate (r_{PFT} , in $\text{gC gN}^{-1} \text{ day}^{-1}$) is a PFT-specific parameter on a 10°C base to represent acclimation of respiration rates to average conditions (Ryan, 1991): k refers to the value proposed by Sprugel et al. (1995) and P is the mean number of individuals per unit area.

The temperature function $g(T)$, describing the influence of temperature on maintenance respiration, is defined as:

$$g(T) = \exp \left[308.56 \cdot \left(\frac{1}{56.02} - \frac{1}{(T + 46.02)} \right) \right] \quad (45)$$

Eq. (45) is a modified Arrhenius equation (Lloyd and Taylor, 1994), where T is either air or soil temperature $[\text{C}]$. This relationship is described by Tjoelker et al. (1999) for a consistent decline of autotrophic respiration with temperature.

While leaf respiration (R_{leaf}) depends on V_m (see eq. (35)) with a static parameter b depending on photosynthetic pathway:

$$R_{\text{leaf}} = V_m \cdot b \quad (46)$$

Gross primary production (GPP, calculated by eq. (34) and converted to $\text{gC m}^{-2} \text{ day}^{-1}$) is reduced by maintenance respiration. Growth respiration, the carbon costs for producing new tissue, is assumed to be 25% of the remainder. The residual is the annual net primary production (NPP):

$$\text{NPP} = (1 - r_{\text{gr}}) \cdot (\text{GPP} - R_{\text{leaf}} - R_{\text{sapwood}} - R_{\text{root}}), \quad (47)$$

where $r_{\text{gr}} = 0.25$ is the share of growth respiration (Thornley, 1970).

370 **Reproduction cost**

As in Sitch et al. (2003), a fixed fraction of 10% of annual NPP is assumed to be carbon costs for producing reproductive organs and propagules in LPJmL4. Since only a very small part of the carbon allocated to reproduction finally enters the next generation, the reproductive carbon allocation is added to the above-ground litter pool to preserve a closed carbon balance in the model.

375 **Tissue turnover**

As in Sitch et al. (2003), a PFT-specific tissue turnover rate is assigned to the living tissue pools (SI-Table S8 and Fig. 1). Leaves and fine roots are transferred to litter and living sapwood to heartwood. Root turnover rates are calculated on a monthly basis, and the conversion of sapwood to heartwood annually. Leaf turnover rates depend on the phenology of the PFT: it is calculated at leaf fall for deciduous and daily for evergreen PFTs.

2.3 Plant functional types

Vegetation composition is determined by the fractional coverage of populations of different plant functional types (PFTs). PFTs are defined to account for the variety of structure and function among plants (Smith et al., 1993). In LPJmL4 11 PFTs are defined, of which eight are woody (two tropical, three temperate, three boreal) and three are herbaceous (Table 1). PFTs are simulated in LPJmL4 as average individuals. Woody PFTs are characterised by the population density and the state variables: crown area (CA) and the size of four tissue compartments: leaf mass (C_{leaf}), fine root mass (C_{root}), sapwood mass (C_{sapwood}), and heartwood mass ($C_{\text{heartwood}}$). The size of all state variables is averaged across the modelled area. The state variables of grasses are represented only by the leaf and root compartments. The physiological attributes and bioclimatic limits control the dynamics of the PFT (see SI-Table S4). PFTs are located in one stand per grid cell and as such compete for light and soil water. That means their crown area and leaf area index determines their capacity to absorb photosynthetic active radiation for photosynthesis (see section 2.2.1) and their rooting profiles determine the access to soil water influencing their productivity (see section 2.6.2). In the following, we describe how carbon is allocated to the different tissue compartments of a PFT (2.3.1) and vegetation dynamics (2.3.2), i.e. how the different PFTs interact. The vegetation dynamics component of LPJmL4 includes the simulation of establishment and different mortality processes.

2.3.1 Allocation

The allocation of carbon is simulated as described in Sitch et al. (2003) and all parameter values are given in SI-Table S6. The assimilated amount of carbon (the remaining NPP) constitutes the annual woody carbon increment which is allocated to leaves, fine roots and sapwood such that four basic allometric relationships (eq. 48 - 51) are satisfied. The pipe model from Shinozaki et al. (1964) and Waring et al. (1982) prescribes that each unit of leaf area must be accompanied by a corresponding area of transport tissue (described by the parameter $k_{\text{la:sa}}$) and, the sapwood cross-sectional area (SA_{ind}):

$$\text{LA}_{\text{ind}} = k_{\text{la:sa}} \cdot \text{SA}_{\text{ind}}, \quad (48)$$

where LA_{ind} is the average individual leaf area and ind gives the index for the average individual.

A functional balance exists between investment in fine root biomass and investment in leaf biomass, respectively. Carbon allocation to $C_{\text{leaf,ind}}$ is determined by the maximum leaf-to-root mass ratio lr_{max} (SI-Table S8), which is a constant and by a water stress index ω (Sitch et al., 2003), which stands for that under water-limited conditions, plants are modelled to allocate relatively more carbon to fine root biomass, which ensures the allocation of relatively more carbon to fine roots under water-limited conditions.

$$C_{\text{leaf,ind}} = \text{lr}_{\text{max}} \cdot \omega \cdot C_{\text{root,ind}} \quad (49)$$

415 The relation between tree height (H) and stem diameter (D) is given as in Huang et al. (1992).

$$H = k_{\text{allom2}} \cdot D^{k_{\text{allom3}}} \quad (50)$$

The crown area (CA_{ind}) to stem diameter (D) relation is based on inverting Reinecke's rule (Zeide, 1993) with k_{rp} as the Reinecke parameter:

$$CA_{\text{ind}} = \min(k_{\text{allom1}} \cdot D^{k_{\text{rp}}}, CA_{\text{max}}), \quad (51)$$

420 which relates tree density to stem diameter under self-thinning conditions. CA_{max} is the maximum crown area allowed. The reversal used in LPJmL4 gives the expected relation between stem diameter and crown area. The assumption here is a closed canopy, but no crown overlap.

By combining the allometric relations of eq. (48) - (51) it follows that the relative contribution of sapwood respiration increases with height, which restricts the possible height of trees.

425 Assuming cylindrical stems and constant wood density (WD), H can be computed and is inversely related to SA_{ind} :

$$SA_{\text{ind}} = \frac{C_{\text{leaf,ind}} \cdot SLA}{k_{\text{la:sa}}} \quad (52)$$

From this follows:

$$H = \frac{C_{\text{sapwood,ind}} \cdot k_{\text{la:sa}}}{WD \cdot C_{\text{leaf,ind}} \cdot SLA} \quad (53)$$

430 Stem diameter can then be calculated by inverting eq. (50). Leaf area is related to leaf biomass $C_{\text{leaf,ind}}$ by PFT-specific SLA, thus, the individual leaf area index (LAI_{ind}) is given by:

$$LAI_{\text{ind}} = \frac{C_{\text{leaf,ind}} \cdot SLA}{CA_{\text{ind}}} \quad (54)$$

SLA is related to leaf longevity (α_{leaf}) in month (see SI-Table S6), which determines whether deciduous or evergreen phenology suits a given climate suggested by Reich et al. (1997). The equation
435 is based on the form suggested by Smith et al. (2014) for needleleaved and broadleaved PFTs as follows:

$$SLA = \frac{2 \times 10^{-4}}{DM_C} \cdot 10^{\beta_0 - \beta_1 \cdot \log(\alpha_{\text{leaf}}) / \log(10)} \quad (55)$$

The parameter β_0 is adapted for broadleaved ($\beta_0 = 2.2$) and needleleaved trees ($\beta_0 = 2.08$) and for grass ($\beta_0 = 2.25$) and β_1 is set to 0.4. Both parameters were derived from data given in Kattge
440 et al. (2011). The dry matter carbon content of leaves DM_C is set to 0.4763 obtained from Kattge et al. (2011). LAI_{ind} can be converted into foliar projective cover (FPC_{ind} , which is the proportion of ground area covered by leaves) using the canopy light-absorption model (Lambert-Beer law, Monsi (1953)):

$$FPC_{\text{ind}} = 1 - \exp(-k \cdot LAI_{\text{ind}}), \quad (56)$$

445 where k is the PFT specific light extinction coefficient (see SI-Table S5). The overall FPC of a PFT in a grid cell is obtained by the product LAI_{ind} , mean individual CA_{ind} , and mean number of individuals per unit area (P), which is determined by the vegetation dynamics (see section 2.3.2).

$$FPC_{PFT} = CA_{ind} \cdot P \cdot FPC_{ind} \quad (57)$$

FPC_{PFT} directly measures the ability of the canopy to intercept radiation (Haxeltine and Prentice, 450 1996).

2.3.2 Vegetation dynamics

Establishment

For PFTs within their bioclimatic limits ($T_{c,min}$, see SI-Table S4), each year, new woody PFT individuals and herbaceous PFTs can establish depending on available space. Woody PFTs have a 455 maximum establishment rate k_{est} of 0.12 (saplings $m^{-2} a^{-1}$), which is a medium value of tree density for all biomes (Luyssaert et al., 2007). New saplings can establish on bare ground in the grid cell that is not occupied by woody PFTs. Establishment rate of tree individuals is calculated:

$$EST_{TREE} = k_{est} \cdot (1 - \exp(-5 \cdot (1 - FPC_{TREE}))) \cdot \frac{(1 - FPC_{TREE})}{n_{estTREE}} \quad (58)$$

The number of new saplings per unit area (EST_{TREE} in $ind m^{-2} a^{-1}$) is proportional to k_{est} and to 460 the FPC of each PFT present in the grid cell (FPC_{TREE} resp. FPC_{GRASS}). It declines in proportion to canopy light attenuation when the sum of woody FPCs exceeds 0.95, thus simulating a decline in establishment success with canopy closure (Prentice et al., 1993). $n_{estTREE}$ gives the number of tree PFTs present in the grid cell. Establishment increases the population density P . Herbaceous PFTs can establish if the sum of all FPCs is less than 1. If the accumulated growing degree days (GDD) 465 reach a PFT-specific threshold GDD_{min} the respective PFT is established (SI-Table S5).

Background mortality

Mortality is modelled by a fractional reduction of P . Mortality always leads to a reduction in biomass per unit area. Similar as in Sitch et al. 2003, a background mortality rate ($mort_{greff}$ in 470 $ind m^{-2} a^{-1}$), the inverse of mean PFT longevity, is applied from the yearly growth efficiency ($greff = bm_{inc} / (C_{leaf,ind} \cdot SLA)$) (Waring, 1983) expressed as the ratio of net biomass increment (bm_{inc}) to leaf area:

$$mort_{greff} = P \cdot \frac{k_{mort1}}{1 + k_{mort2} \cdot greff}, \quad (59)$$

where k_{mort1} is an asymptotic maximum mortality rate, and k_{mort2} is a parameter governing the slope of the relationship between growth efficiency and mortality (SI-Table S6).

475 Stress mortality

Mortality from competition occurs when tree growth leads to too high tree densities (FPC of all trees exceeds > 95%). In this case, all tree PFTs are reduced proportionally to their expansion. Herbaceous PFTs are outcompeted by expanding trees until these reach their maximum FPC of 95% or by light competition between herbaceous PFTs. Dead biomass is transferred to the litter pools.

480 Boreal trees can die from heat stress ($\text{mort}_{\text{heat}}$ in $\text{ind m}^{-2} \text{a}^{-1}$) (Allen et al., 2010). It occurs in LPJmL4 when a temperature threshold ($T_{\text{mort,min}}$ in °C, SI-Table S4) is exceeded, but only for boreal trees (Sitch et al., 2003). Temperatures above this threshold are accumulated over the year (gdd_{tw}) and this is related to a parameter value of the heat damage function (tw_{PFT}), which is set to 400:

$$485 \text{ mort}_{\text{heat}} = P \cdot \min \left(\frac{\text{gdd}_{\text{tw}}}{\text{tw}_{\text{PFT}}}, 1 \right) \quad (60)$$

P is reduced for both $\text{mort}_{\text{heat}}$ and $\text{mort}_{\text{greff}}$.

Fire disturbance and mortality

Two different fire modules can be applied in the LPJmL4 model: the simple Glob-FIRM model (Thonicke et al., 2001) and the process-based SPITFIRE model (Thonicke et al., 2010). In Glob-
490 FIRM, fire disturbances are calculated as an exponential probability function dependent on soil moisture in the top 50 cm and a fuel load threshold. The sum of the daily probability determines the length of the fire season. Burnt area is assumed to increase nonlinearly with increasing length of fire season. The fraction of trees killed within the burnt area depends on a PFT-specific fire resistance parameter for woody plants, while all litter and live grasses are consumed by fire. Glob-FIRM does
495 not specify fire ignition sources and assumes a constant relationship between fire season length and resulting burnt area. The PFT-specific fire resistance parameter implies that fire severity is always the same, an approach suitable for model applications to multi-century time scales or paleo-climate conditions.

In SPITFIRE, fire disturbances are simulated as the fire processes risk, ignition, spread and ef-
500 fects separately. The climatic fire danger is based on the Nesterov index $\text{NI}(N_d)$, which describes atmospheric conditions critical to fire risk for day N_d :

$$\text{NI}(N_d) = \sum_{d=1}^{N_d} T_{\text{max}}(d) \cdot (T_{\text{max}}(d) - T_{\text{dew}}(d)), \quad (61)$$

if $\text{Pr}(d) \leq 3 \text{ mm}$

where T_{max} and T_{dew} are the daily maximum and dew-point temperature, and d is a positive temperature day with a precipitation of less than 3 mm. The probability of fire spread P_{spread} decreases
505 linearly as litter moisture ω_0 increases towards its moisture of extinction m_e :

$$P_{\text{spread}} = \begin{cases} 1 - \omega_0/m_e, & \omega_0 \leq m_e \\ 0, & \omega_0 > m_e \end{cases} \quad (62)$$

Combining NI and P_{spread} we can calculate the fire danger index FDI:

$$\text{FDI} = \max \left\{ 0, 1 - \frac{1}{m_e} \cdot \exp \left(-\text{NI} \cdot \sum_{p=1}^n \frac{\alpha_p}{n} \right) \right\} \quad (63)$$

to interpret the qualitative fire risk in quantitative terms. The value of α_p defines the slope of the probability risk function given as the average PFT parameter (see SI-Table S9) for all existing PFTs (n). SPITFIRE considers human-caused and lightning-caused fires as sources for fire ignition. Lightning-caused ignition rates are prescribed from the OTD/LIS satellite product (Christian et al., 2003). Since it quantifies total flash rate, we assume that 20% of these are cloud-to-ground flashes (Latham and Williams, 2001) and that, under favourable burning conditions, their effectiveness to start fires is 0.04 (Latham and Williams, 2001; Latham and Schlieter, 1989). Human-caused ignitions are modelled as a function of human population density assuming that ignition rates are higher in remote regions and declines with increasing level of urbanisation and associated effects of landscape fragmentation, infrastructure and improved fire monitoring. The function is:

$$n_{h,\text{ig}} = P_D \cdot k(P_D) \cdot a(N_D)/100, \quad (64)$$

where

$$k(P_D) = 30.0 \cdot \exp(-0.5 \cdot \sqrt{P_D}). \quad (65)$$

P_D is the population density [individuals km^{-2}], and $a(N_D)$ [ignitions individual $^{-1}$ day $^{-1}$] is a parameter describing the inclination of humans to use fire and cause fire ignitions. In absence of further information $a(N_D)$ can be calculated from fire statistics using the following approach

$$a(N_D) = \frac{N_{h,\text{obs}}}{t_{\text{obs}} \cdot \text{LFS} \cdot P_D}, \quad (66)$$

where $N_{h,\text{obs}}$ is the average number of human-caused fires observed during the observation years t_{obs} in a region with the average length of fire season (LFS) and the mean human population density. Assuming that all fires ignited in one day have the same burning conditions in a 0.5° grid cell with the grid cell size A , we combine fire danger, potential ignitions and the mean fire area A_f to obtain daily total burnt area with:

$$A_b = \min(E(n_{\text{ig}}) \cdot \text{FDI} \cdot A_f, A) \quad (67)$$

We calculate $E(n_{\text{ig}})$ with the sum of independent estimates of numbers of lightning ($n_{l,\text{ig}}$) and human-caused ignition events ($n_{h,\text{ig}}$), disregarding stochastic variations. A_f is calculated from forward and backward rate of spread which depends on the dead fuel characteristics, fuel load in the respective dead fuel classes and wind speed. Dead plant material entering the litter pool is subdivided into 1-, 10-, 100- and 1000 hour fuel classes, describing the amount of time to dry a fuel particle of a specific size (1-hour fuel refers to leaves and twigs and 1000 hour fuel to tree boles). As described

by Thonicke et al. (2010): "The forward rate of spread $ROS_{f,surface}$ [$m \min^{-1}$] is given by:

$$ROS_{f,surface} = \frac{I_R \cdot \zeta \cdot (1 + \Phi_w)}{\rho_b \cdot \epsilon \cdot Q_{ig}}, \quad (68)$$

540 where I_R is the reaction intensity, i.e. the energy release rate per unit area of fire front [$kJ m^{-2} \min^{-1}$]; ζ is the propagating flux ratio, i.e. the proportion of I_R that heats adjacent fuel particles to ignition; Φ_w is a multiplier that accounts for the effect of wind in increasing the effective value of ζ ; ρ_b is the fuel bulk density [$kg m^{-3}$], assigned by PFT and weighted over the 1-, 10- and 100-hour dead fuel classes; ϵ is the effective heating number, i.e. the proportion of a fuel particle that is heated to
545 ignition temperature at the time flaming combustion starts; and Q_{ig} is the heat of pre-ignition, i.e. the amount of heat required to ignite a given mass of fuel [$kJ kg^{-1}$]. With fuel bulk density ρ_b defined as a PFT parameter, surface-area-to-volume ratios change with fuel load." Assuming that fires burn longer under high fire danger, we define fire duration (t_{fire}) [min] as

$$t_{fire} = \frac{241}{1 + 240 \cdot \exp(-11.06 \cdot FDI)} \quad (69)$$

550 In the absence of topographic influence and changing wind directions during one fire event or discontinuities of the fuel bed, fires burn an elliptical shape. Thus, the mean fire area [in ha] is defined as follows:

$$\overline{A_f} = \frac{\pi}{4 \cdot L_B} \cdot D_T^2 \cdot 10^{-4} \quad (70)$$

with L_B is length to breadth ratio of elliptical fire, and D_T is the length of major axis with:

$$555 \quad D_T = ROS_{f,surface} \cdot t_{fire} + ROS_{b,surface} \cdot t_{fire} \quad (71)$$

with $ROS_{b,surface}$, surface as the backward rate of spread. L_B for grass and trees, respectively, is weighted depending on the foliage projective cover of grasses relative to woody PFTs in each grid cell.

SPITFIRE differentiates fire effects depending on burning conditions (intra- and interannual). If
560 fires have developed insufficient surface fire intensity ($< 50 \text{ kW m}^{-1}$), ignitions are extinguished (and not counted in the model output). If the surface fire intensity has supported high enough scorch height of the flames, resulting scorching of the crown is simulated. Here, the tree architecture through the crown length, height of the tree determines fire effects and describes an important feedback between vegetation and fire in the model. PFT-specific parameters describe the trees sensitivity to or influence
565 on scorch height and crown scorch. Surface fires consume dead fuel and live grass as a function of their fuel moisture content. The amount of biomass burnt results from crown scorch and surface fuel consumption.

Post-fire mortality is modelled as a result of two fire mortality causes: crown and cambial damage. The latter occurs when insufficient bark thickness allows the heat of the fire to damage the cambium.

570 It is defined as the ratio of the residence time of the fire to the critical time for cambial damage. The probability of mortality due to crown damage (CK) is:

$$P_m(\text{CK}) = r_{\text{CK}} \cdot \text{CK}^p, \quad (72)$$

where r_{CK} is a resistance factor between 0 and 1, and p is in the range of 3 to 4 (see SI-Table S9). The biomass of trees which die from either mortality cause is added to the respective dead fuel classes.

575 In summary, the PFT composition and productivity strongly influences fire risk through the moisture of extinction, fire spread through composition of fuel classes (fine vs. coarse fuel), openness of the canopy and fuel moisture, fire effects through stem diameter, crown length and bark thickness of the average tree individual. The higher the proportion of grasses in a grid cell the faster fires can spread, the smaller the trees and/or the thinner their bark the higher the proportion of the crown
580 scorched and the higher their mortality.

2.4 Crop functional types

In LPJmL4, twelve different annual crop functional types (CFTs) are simulated (SI-Table S10), similar to Bondeau et al. (2007) with the addition of sugarcane. The basic idea of CFTs is that these are parameterized as one specific representative crop (e.g. wheat, *Triticum aestivum* L.) to represent a
585 broader group of similar crops (e.g. temperate cereals). In addition to the crops represented by the twelve CFTs, other annual and perennial crops (other crops) are typically represented as managed grassland. Bioenergy crops are simulated to account for woody (willow trees in temperate regions, eucalyptus for tropical regions) and herbaceous types (*Miscanthus*) (Beringer et al., 2011)). The physical cropping area (i.e. proportion per grid cell) of each CFT, the group of other crops, managed
590 grasslands and bioenergy crops can be prescribed for each year and grid cell by using gridded land use data described in Fader et al. (2010) and Jägermeyr et al. (2015), see section 2.7. In principle, any land use dataset (including future scenarios) can be implemented in LPJmL4 at any resolution.

Crop varieties and phenology

Phenological development of crops in LPJmL4 is driven by temperature through accumulation of
595 growing degree days, and can be modified by vernalization requirements and sensitivity to daylength (photoperiod) for some CFTs and some varieties. Phenology is represented as a single phase from planting to physiological maturity. Different varieties of a single crop species are represented by different phenological heat unit requirements to reach maturity (phu), but also different harvest indices (hi_{opt}), i.e. the fraction of the above-ground biomass that is harvested, is typically CFT-specific, but
600 can be specified to represent specific varieties (Asseng et al., 2013; Bassu et al., 2014; Kollas et al., 2015; Fader et al., 2010).

Heat units (hu_t , growing degree days) are accumulated (hu_{sum}) daily (see eq. (73)). The daily heat unit increment (hu_t) is the difference between the daily mean temperature of day t and the CFT-

specific base temperature (see SI-Table S10). The increment hu_t cannot be less than zero at any given
605 day. The phenological stage of the crop development (fphu) is expressed as the ratio of accumulated
(hu_{sum}) and required phenological heat units (phu) (see eq. (74)). Physiological maturity is reached
as soon as the sufficient growing degree days have been accumulated (fphu= 1.0). Both unfulfilled
vernalization requirements as well as unsuitable photoperiod affect the phenological development
of the CFTs (see eq. (78) and eq. (79)). Therefore, the daily increment hu_t at day t is scaled by
610 reduction factors v_{rf} for vernalization and p_{rf} for photoperiod:

$$hu_{sum} = \sum_{t'=sdate}^t hu_{t'} \cdot v_{rf} \cdot p_{rf} \quad (73)$$

and

$$fphu = hu_{sum}/phu. \quad (74)$$

Wheat and rapeseed are implemented as spring and winter varieties. The model endogenously de-
615 termines which variety to grow based on the average climate of past decades. If internally computed
sowing dates for winter varieties (see below Section 2.7.1) indicate that the winter is too long to al-
low for growing winter varieties, which is prior to day 258 (90) for wheat and 241 (61) for rapeseed on
the northern (southern hemisphere), spring varieties are grown instead. These are computed on the
basis of the sowing dates (sdate) as an indication for the length of the cropping season, constrained
620 by crop-specific limits. For winter varieties of wheat and rapeseed, phu is computed as:

$$phu = -0.1081 \cdot (sdate - keyday)^2 + 3.1633 \\ \cdot (sdate - keyday) + phu_{w_{high}}, \quad phu \leq phu_{w_{low}}, \quad (75)$$

where $phu_{w_{low}}$ and $phu_{w_{high}}$ are minimum and maximum phu requirements for winter varieties, re-
spectively. The sowing date sdate can either be internally computed (see section 2.7.1) or prescribed
625 for a crop and pixel. The parameter keyday is day 365 on the northern and day 181 on the south-
ern hemisphere. For spring varieties of wheat and rapeseed, as well as for all other crops, phu is
computed as:

$$phu = \max(T_{base_{low}}, atemp_{20}) \cdot pf_{CFT}, \\ phu_{s_{high}} \geq phu \geq phu_{s_{low}}, \quad (76)$$

630 where $phu_{s_{low}}$ and $phu_{s_{high}}$ are minimum and maximum phu requirements for spring varieties, re-
spectively, $T_{base_{low}}$ is the minimum base temperature for the accumulation of heat units, $atemp_{20}$ is
the 20-year moving average annual temperature and pf_{CFT} is a CFT-specific scaling factor.

Vernalization requirements pvd are zero for spring varieties and are computed for winter varieties:

$$635 \quad pvd = vern_{date20} - sdate - ppvd_{CFT}, \quad 0 \leq pvd \leq 60 \quad (77)$$

with $ppvd_{CFT}$ as a CFT-specific vernalization factor, $sdate$ as the Julian day of the year of sowing and $vern_{date20}$ as the multi-annual average of the first day of the year when temperatures rise above a CFT-specific vernalization threshold (T_{vern} , see SI-Table S10). The effectiveness of vernalization is dependent on the daily mean temperature, being ineffective below $-4^{\circ}C$ and above $17^{\circ}C$, being fully effective between $3^{\circ}C$ and $10^{\circ}C$ and the effectiveness scales linear between -4 and 3 and between 10 and $17^{\circ}C$. The effective number of vernalizing days vd_{sum} is accumulated until the requirements (pvd) as computed in eq. (77) are met or until phenology has progressed over 20% of its phenological development (i.e. $fphu \geq 0.2$). Crop varieties can be parameterized as sensitive to photoperiod (i.e. daylength), but here are assumed to be insensitive. Parameter settings can be adjusted for specific applications, such as in model intercomparisons (Asseng et al., 2013; Bassu et al., 2014; Kollas et al., 2015). Photoperiod restrictions are active until the crop reaches senescence.

The reduction factors are computed as:

$$v_{rf} = (vd_{sum} - 10.0) / (pvd - 10.0), \quad (78)$$

forcing v_{rf} to be between 0 and 1, and

$$p_{rf} = (1 - p_{sens}) \cdot \min(1, \max(0, (daylength - p_b) / (p_s - p_b))) + p_{sens}, \quad (79)$$

where p_{sens} is the parametrized sensitivity to photoperiod ($0 \dots 1$), daylength is the duration of daylight (sunrise to sunset) in hours (see section 2.1.1), p_b is the base photoperiod in hours and p_s is the saturation photoperiod in hours.

Crop growth and allocation

Photosynthesis and autotrophic respiration of crops are computed as for the herbaceous natural PFTs (see section 2.2.1 and 2.2.3). Light absorption for photosynthesis is computed based on the Lambert-Beer law (Monzi, 1953), except for maize. For maize, LPJmL4 employs a linear -FPAR model (Zhou et al., 2002) and a maximum leaf area index (LAI_{max}) of 5 instead of 7 as for all other CFTs (Fader et al., 2010). Daily NPP accumulates to total biomass and is allocated daily to crop organs in a hierarchical order: roots, leaves, storage organ, mobile reserves/stem (pool). The fraction of biomass that is allocated to each compartment depends on the phenological development stage ($fphu$). The fraction of total biomass that is allocated to the roots (f_{root}) ranges between 40% at planting and 10% at maturity, modified by water stress:

$$f_{root} = \frac{0.4 - (0.3 \cdot fphu) \cdot wdf}{wdf + \exp(6.13 - 0.0883 \cdot wdf)}, \quad (80)$$

where wdf is the ratio between accumulated daily transpiration and accumulated daily water demand since planting, representing a measure of the average water stress. After allocation to the roots, biomass is allocated to the leaves. Leaf area development follows a CFT-specific shape that is

controlled by phenological development (fphu), the onset of senescence (ssn) and the shape of green
 670 LAI decline after onset of senescence. The ideal CFT-specific development of the canopy (equation
 81) is thus described as a function of the maximum LAI (lai_{max}) and the phenological develop-
 ment (fphu) with two turning points in the phenological development (fphu_c and fphu_k) and the
 corresponding fraction of the maximum green LAI reached at these stages (flai_{max_c} and flai_{max_k}):

$$\text{flai}_{\max} = \frac{\text{fphu}}{\text{fphu} + c \cdot (c/k)^{\frac{\text{fphu}_c - \text{fphu}}{\text{fphu}_k - \text{fphu}_c}}} \quad (81)$$

675 with

$$c = \frac{\text{fphu}_c}{\text{flai}_{\max_c} - \text{fphu}_c} \quad (82)$$

$$k = \frac{\text{fphu}_k}{\text{flai}_{\max_k} - \text{fphu}_k} \quad (83)$$

The onset of senescence is defined as a point in the phenological development fphu_{sen}. After
 the onset of senescence, i.e. fphu ≤ fphu_{sen}, no more biomass is allocated to the leaves and the

680 maximum green LAI is computed as:

$$\text{flai}_{\max} = \left(\frac{1 - \text{fphu}}{1 - \text{fphu}_{\text{sen}}} \right)^{\text{ssn}} \cdot (1 - \text{flai}_{\max_h}) + \text{flai}_{\max_h} \quad (84)$$

with flai_{max_h} as the green LAI fraction at which harvest occurs. This optimal development of LAI
 is modified by acute water stress. For this, the daily increment lai_{inc}, which is optimal for day *t* is
 computed as:

$$685 \text{ lai}_{\text{inc}_t} = (\text{flai}_{\max_t} - \text{flai}_{\max_{t-1}}) \cdot \text{lai}_{\max} \quad (85)$$

with flai_{max_t} as the maximum green LAI of day *t* and flai_{max_{t-1}} as the maximum green LAI of
 the previous day. The daily increment lai_{inc} is additionally scaled with the daily water stress (ω),
 which is calculated as the ratio of actual transpiration and demand (see section 2.6.2) on that day.
 The calculation of lai_{inc} applies to daily LAI increments which are independent of each other. The

690 LAI on day *t* is accumulated from daily LAI increments,

$$\text{LAI}_t = \sum_{t'=\text{sdate}}^t \text{lai}_{\text{inc}_{t'}} \cdot \omega \quad (86)$$

and implies that the LAI development cannot recover from water-limitation induced reductions in
 LAI. Until the onset of senescence, the daily LAI determines the biomass allocated to the leaves
 by dividing LAI by specific leaf area (SLA). SLA is computed as in eq. (55) using the β₀ value
 695 for grasses (2.25) and CFT-specific α_{leaf} values (SI-Table S11). Its calculation was adjusted for SLA
 values given in Xu et al. (2010). Biomass in the storage organ is computed by phenological stage and
 the harvest index (HI), which describes the fraction of the above ground biomass that is allocated to
 the storage organ:

$$\text{HI} = \begin{cases} \text{fhi}_{\text{opt}} \cdot \text{hi}_{\text{opt}}, & \text{if } \text{hi}_{\text{opt}} \geq 1 \\ \text{fhi}_{\text{opt}} \cdot (\text{hi}_{\text{opt}} - 1) + 1, & \text{otherwise} \end{cases} \quad (87)$$

700 with

$$fhi_{opt} = 100 \cdot fphu / (100 \cdot fphu + \exp(11.1 - 10.0 \cdot fphu)) \quad (88)$$

As the harvest index HI is defined relative to above-ground biomass, roots and tubers have HI values larger than 1.0 which needs to be accounted for in the allocation of biomass to the storage organ (see eq. (87)). If biomass is limiting (low NPP), biomass is allocated in hierarchical order, starting with roots (which can always be satisfied, as it is 40% of total biomass maximum), followed by leaves (C_{leaf}) (where eventually the LAI is temporarily reduced, impacting APAR and thus NPP) and the storage organ (C_{so}). If biomass is not limiting, the allocation to the storage organ, this is computed from the harvest index (HI) and total above ground biomass:

$$C_{so} = HI \cdot (C_{leaf} + C_{so} + C_{pool}) \quad (89)$$

710 Excess biomass after allocating to roots, leaves and storage organ is allocated to a pool (C_{pool}) that represents mobile reserves and the stem. At harvest, storage organs are collected from the field and crop residues can be left on the field or removed (for scenario setting, see Bondeau et al. (2007, e.g.)). If removed, a fraction of 10% of the above-ground biomass (leaves and pool) is assumed to remain on the field as stubbles. Stubbles and root biomass enter the litter pools after harvest.

715 2.5 Soil and litter carbon pools

Important for the global carbon balance are the biogeochemical processes in soil and litter. The LPJmL4 litter pool consists of CFT- resp. PFT-dependent pools for leaf, root and wood. The soil consists of a fast and a slow organic matter pool. Decomposition fluxes transferring litter carbon into soil carbon and losses for heterotrophic respiration (R_h) are described in the following section.

720 2.5.1 Decomposition

Decomposition of organic matter pools is represented by first-order kinetics (Sitch et al., 2003)

$$\frac{dC_{(l)}}{dt} = -k_{(l)} \cdot C_{(l)}, \quad (90)$$

where $C_{(l)}$ is the carbon pool size of soil or litter and $k_{(l)}$ is the annual decomposition rate per layer (l) in day^{-1} . Integrating for a time interval Δt (here 1 day) yields:

$$725 \quad C_{(t+1,l)} = C_{(t,l)} \cdot \exp(-k_{(l)} \cdot \Delta t), \quad (91)$$

where $C_{(t,l)}$ and $C_{(t+1,l)}$ are the carbon pool sizes at the beginning and the end of the day. The amount of carbon decomposed per layer is:

$$C_{(t,l)} \cdot (1 - \exp(-k_{(l)} \cdot \Delta t)) \quad (92)$$

at which 70% of decomposed litter goes directly into the atmosphere $R_{h,\text{litter}}$, the remaining is transferred to the soil carbon pools, 98.5% to the fast soil carbon pool and 1.5% to the slow carbon pool (Sitch et al., 2003).

$$R_h = R_{h,\text{litter}} + R_{h,\text{fastSoil}} + R_{h,\text{slowSoil}} \quad (93)$$

The decomposition rates for root litter and soil ($k_{(l,\text{PFT})}$) is a function of soil temperature and soil moisture:

$$k_{(l,\text{PFT})} = \frac{1}{\tau_{10\text{PFT}}} \cdot g(T_{\text{soil}(l)}) \cdot f(\theta_{(l)}), \quad (94)$$

which is reciprocal to the mean residence time ($\tau_{10\text{PFT}}$). Root litter decomposition is defined for all PFTs (0.3 a^{-1}) and for fast and slow soil carbon (0.03 and 0.001 a^{-1} resp.) as in Sitch et al. (2003), p represents the different pools. The decomposition rate of leaf and wood litter is defined as PFT-specific decomposition rates at 10°C for leaf, wood and root, which has been analysed and proposed by Brovkin et al. (2012) for leaf and wood. The temperature dependence function for the fast and slow soil carbon and the leaf and root litter pool $g(T_{\text{soil}})$ was already described in eq. (45). For wood litter decomposition it is calculated as follows:

$$k_{\text{wood,PFT}} = (Q_{10\text{wood,litter}})^{\frac{(T_{\text{soil}}-10)}{10.0}} \quad (95)$$

SI-Table S7 presents $(1/\tau_{10\text{PFT}})$ used for leaves and wood and the $Q_{10\text{wood,litter}}$ parameter for temperature-dependent wood decomposition in the litter pool. The soil moisture function follows Schaphoff et al. (2013):

$$f(\theta_{(l)}) = 0.0402 - 5.005 \cdot \theta_{(l)}^3 + 4.269 \cdot \theta_{(l)}^2 + 0.719 \cdot \theta_{(l)} \quad (96)$$

$\theta_{(l)}$ is the soil volume fraction of the layer l . Parameters are chosen based on the assumption that rates are maximal at field capacity and decline for higher $\theta_{(l)}$ to 0.2. $f(\theta_{(l)})$ is very small (0.0402) when $\theta_{(l)}$ equals 1 due to oxygen limitation and when $\theta_{(l)}$ is 0.

To account for different decomposition rates in the different soil layers, a vertical soil carbon distribution is now implemented in LPJmL4 following Schaphoff et al. (2013). Jobbagy and Jackson (2000) suggested a cumulative log-log distribution of the fraction of soil organic carbon (Cf_l) as a function of depth with:

$$\text{Cf}_l = 10^{k_{\text{soc}} \cdot \log_{10}(d_{(l)})}, \quad (97)$$

where $d_{(l)}$ is the relative share of the layer l in the entire soil bucket and the parameters k_{soc} was adjusted for the soil layer depth now used in LPJmL4 (see SI-Table S7). The total amount of soil carbon C_{stotal} is estimated from the mean annual decomposition rate $k_{\text{mean}(l)}$ and the mean litter

input into the soil as in (Sitch et al., 2003), but is distributed to all root layers separately (eq. (98)).
The envisaged vertical soil distribution $C_{(l)}$:

$$C_{(l)} = \sum_{PFT=1}^{n_{PFT}} d_{(l)}^{k_{socPFT}} \cdot C_{s_{total}} \quad (98)$$

is estimated after a carbon equilibrium phase of 2310 years. The mean decomposition rate for each PFT $k_{meanPFT}$ can be derived from the mean annual decomposition rate $k_{mean(l)}$ of the spinup years
as a layer-weighted value derived from eq. (97):

$$k_{meanPFT} = \sum_{l=1}^{n_{soil}} k_{mean(l)} \cdot Cf_{(l,PFT)} \quad (99)$$

The annual carbon shift rates $C_{shift(l,p)}$ describe the organic matter input from the different PFTs into the respective layer due to cryoturbation and bioturbation and are designed for global applications:

$$C_{shift(l,PFT)} = \frac{Cf_{(l,PFT)} \cdot k_{mean(l)}}{k_{meanPFT}} \quad (100)$$

2.6 Water balance

The terrestrial water balance is a pivotal element in LPJmL4 as water and vegetation are linked in multiple ways:

1. the coupling of plant transpiration and carbon uptake from the atmosphere through stomatal conductance in the process of photosynthesis;
2. the down-regulation of photosynthesis, plant growth and productivity in response to soil water limitation (relative to atmospheric moisture demand), in case actual canopy conductance is below potential canopy conductance (in the demand function that describes transpiration);
3. the effect of changes in vegetation type, distribution, phenology and production on evaporation, transpiration, interception, runoff and soil moisture;
4. the anthropogenic stimulation of crop growth through irrigation with water taken from rivers, dams, lakes and assumed renewable groundwater.

These couplings of water and vegetation dynamics enable simulations of the interacting mutual feedbacks between freshwater cycling in and above the earth surface and terrestrial vegetation dynamics.

2.6.1 Soil water balance

Advancing the former two-layer approach (Sitch et al., 2003), LPJmL4 divides the soil column into five hydrological active layers of 0.2, 0.3, 0.5, 1 and 1 m thickness (Schaphoff et al., 2013). Water holding capacity (water content at permanent wilting point, at field capacity and at saturation) and

hydraulic conductivity are derived for each grid cell using soil texture from the Harmonized World
 790 Soil Database (HWSD) version 1 (Nachtergaele et al., 2008) and relationships between texture and
 hydraulic properties from Cosby et al. (1984), see also section 2.1.3.

Water content in soil layers is altered by infiltrating rainfall and vertical movement of gravitational
 water (percolation). Since the accuracy needed for a global model does not justify the computational
 costs of an exact solution of the governing differential equation, a simplified storage approach is
 795 implemented in LPJmL4. Rather than calculating infiltration and percolation of precipitation at once,
 total precipitation is divided in portions of 4 mm that are routed through the soil one after another.
 This effectively emulates a time discretization, which leads to a higher proportion of runoff being
 generated for higher amounts precipitation.

Infiltration

800 The infiltration rate of rain and irrigation water into the soil (infil, in mm) depends on current soil
 water content of the first layer as follows:

$$\text{infil} = P \cdot \sqrt{1 - \frac{SW_{(1)} - W_{\text{pwp}(1)}}{W_{\text{sat}(1)} - W_{\text{pwp}(1)}}}, \quad (101)$$

where $W_{\text{sat}(1)}$ is the soil water content at saturation and $W_{\text{pwp}(1)}$ the soil water content at wilting
 point, and $SW_{(1)}$ the total actual soil water content of the first layer, all in mm. P is the amount of
 805 water in the current portion of daily precipitation or applied irrigation water (maximum 4 mm). The
 surplus water that does not infiltrate is assumed to generate surface runoff.

Percolation

Subsequent percolation through the soil layers is calculated by the storage routine technique (Arnold
 et al., 1990) as used in regional hydrological models such as SWIM (Krysanova et al., 1998).

$$810 \quad FW_{(t+1, l)} = FW_{(t, l)} \cdot \exp\left(-\frac{\Delta t}{TT_{(l)}}\right), \quad (102)$$

where $FW_{(t, l)}$ and $FW_{(t+1, l)}$ are the soil water content between field capacity and saturation at
 the beginning and the end of the day for all soil layers l , respectively. Δt is the time interval (here,
 24 hours) and TT_l determines the travel time through the soil layer in hours:

$$TT_{(l)} = \frac{FW_{(l)}}{HC_{(l)}} \quad (103)$$

815 $HC_{(l)}$ is the hydraulic conductivity of the layer in mm hour^{-1} :

$$HC_{(l)} = K_{s(l)} \cdot \left(\frac{SW_{(l)}}{W_{\text{sat}(l)}}\right)^{\beta_{(l)}}, \quad (104)$$

where $W_{\text{sat}(l)}$ is the soil water content at saturation, $K_{s(l)}$ is the saturated conductivity [in mm hour^{-1}]
 and $SW_{(l)}$ the total soil water content of the layer [in mm]. Thus, percolation can be calculated by

subtracting $FW_{(t,l)}$ from $FW_{(t+1,l)}$ for all soil layers:

$$820 \quad \text{perc}'_{(l)} = FW_{(t,l)} \cdot \left[1 - \exp\left(\frac{-\Delta t}{TT_{(l)}}\right) \right] \quad (105)$$

The percolation $\text{perc}_{(l)}$ [in mm day^{-1}] is limited by soil moisture of the lower layer, similar to the infiltration approach.

$$\text{perc}_{(l)} = \text{perc}'_{(l)} \cdot \sqrt{1 - \frac{SW_{(l)} - W_{\text{pwp}(l)}}{W_{\text{sat}(l)} - W_{\text{pwp}(l)}}} \quad (106)$$

Excess water over the saturation levels forms lateral runoff in each layer and contributes to sub-
825 surface runoff. The formation of groundwater, which is the seepage from the bottom soil layer, has been recently introduced into LPJmL4 (Schaphoff et al., 2013). Both surface and subsurface runoff are simulated to accumulate to river discharge (see section 2.6.3).

2.6.2 Evapotranspiration

Similar to Gerten et al. (2004), evapotranspiration (ET) is the sum of vapor flow from the earth
830 surface to the atmosphere. It consists of three major components: evaporation from bare soils, evaporation of intercepted rainfall from the canopy, and plant transpiration through leaf stomata. The calculation of these different components in LPJmL4 is based on equilibrium evapotranspiration (E_{eq}) resp. the PET as described in section 2.1.1 and eq. (2) and (6).

Canopy evaporation

835 Canopy evaporation is the evaporation of rainfall that has been intercepted by the canopy, limited either by PET or the amount of intercepted rainfall I (both in mm day^{-1}):

$$E_{\text{canopy}} = \min(\text{PET}, I) \quad (107)$$

The amount of intercepted rainfall is given as:

$$I = \sum_{\text{PFT}=1}^{n_{\text{PFT}}} I_{\text{PFT}} \cdot \text{LAI}_{\text{PFT}} \cdot \text{Pr}, \quad (108)$$

840 where I_{PFT} is the interception storage parameter for each PFT (Gerten et al., 2004), LAI_{PFT} the PFT-specific leaf area per unit of grid cell area and Pr is daily precipitation in mm day^{-1} .

Soil evaporation

Soil evaporation (E_s in mm day^{-1}) only occurs from bare soil, where the vegetation cover (f_v) is less than 100%. The f_v is the sum of all present PFT's FPC (see eq. (57)) taking daily phenology
845 into account. The evaporation flux depends on available energy for the vaporization of water (see eq. (6)) and the available water in the soil. LPJmL4 assumes that water for evaporation is available

from the upper 0.3 m of the soil, implicitly accounting for some capillary rise. Evaporation-available soil water (w_{evap}) is thus all water above wilting point of the upper layer (0.2 m) and one third of the second layer (0.3 m). Actual evaporation is then computed according to eq. (110), with w being
850 the evaporation-available water relative to the water holding capacity in that layer whc_{evap}

$$w = \min(1, w_{\text{evap}} / \text{whc}_{\text{evap}}) \quad (109)$$

thus:

$$E_s = \text{PET} \cdot w^2 \cdot (1 - f_v) \quad (110)$$

This potential evaporation flux is reduced if a portion of the water is frozen or if the energy for the
855 vaporization has already been used to vaporize water that was intercepted by the canopy or for plant transpiration (see section 2.6.2).

Plant transpiration coupled with photosynthesis

Plant transpiration (E_T in mm day^{-1}) is modelled as the lesser of plant-available soil water supply function (S) and atmospheric demand function (D), following Federer (1982):

$$860 \quad E_T = \min(S, D) \cdot f_v \quad (111)$$

S depends on a PFT-specific maximum water transport capacity (E_{max} in mm day^{-1}) and the relative water content (w_r) and phenology (phen_{PFT}):

$$S = E_{\text{max}} \cdot w_r \cdot \text{phen}_{\text{PFT}} \quad (112)$$

The water accessible for plants (w_r) is computed from the relative water content at field capacity
865 (w_l) and the fraction of roots (rootdist_l) within each soil layer (l) as

$$w_r = \sum_{l=1}^{n_{\text{soil}}-1} w_l \cdot \text{rootdist}_l \quad (113)$$

rootdist_l can be calculated from the proportion of roots from surface to soil depth z , rootdist_z , as in Jackson et al. (1996):

$$\text{rootdist}_z = \frac{\int_0^z (\beta_{\text{root}})^{z'} dz'}{\int_0^{z_{\text{bottom}}} (\beta_{\text{root}})^{z'} dz'} = \frac{1 - (\beta_{\text{root}})^z}{1 - (\beta_{\text{root}})^{z_{\text{bottom}}}}, \quad (114)$$

870 where β_{root} represents a numerical index for root distribution (for parameter values see SI-Table S8). rootdist_l is given by the difference $\text{rootdist}_{z(l)} - \text{rootdist}_{z(l-1)}$. If the soil depth of layer l is greater than the thawing depth then rootdist_l is set to zero. The non-zero rootdist_l are rescaled in such a way that their sum is normalized to 1 considering the reallocation of the root distribution under freezing conditions.

875 Plants in natural vegetation compete for water resources and thus only have access to the fraction of water that corresponds to their foliage projected cover (FPC_{PFT})

$$S_{PFT} = S \cdot FPC_{PFT} \quad (115)$$

For agricultural crops, water supply is also dependent on their root biomass bm_{root}

$$S = E_{max} \cdot w_r \cdot (1 - \exp(-0.0411 \cdot bm_{root})) \quad (116)$$

880 Atmospheric demand (D) is a hyperbolic function of g_c (see section 2.2.1 and eq. (39)), following Monteith (1995), and employs a maximum Priestley-Taylor coefficient $\alpha_m = 1.391$, describing the asymptotic transpiration rate, and a conductance scaling factor $g_m = 3.26$:

$$D = (1 - wet) \cdot E_{eq} \cdot \alpha_m / (1 + g_m / g_c), \quad (117)$$

where wet is the fraction of E_{eq} that was used to vaporize intercepted water from the canopy (see 885 section 2.6.2) and g_c is the potential canopy conductance. If S is not sufficient to fulfill transpiration demand g_c is recalculated for $D = S$ and photosynthesis rate might be adjusted (see section 2.2.1).

2.6.3 River routing

Description of the river routing module

The river routing module computes the lateral exchange of discharge (see section 3.1.2 for input) 890 between grid cells through the river network (Rost et al., 2008). The transport of water in the river channel is approximated by a cascade of linear reservoirs. River sections are divided into n homogeneous segments of length L , each behaving like a linear reservoir. Following the unit hydrograph method (Nash, 1957), the outflow $Q_{out}(t)$ of a linear reservoir cascade for an instantaneous inflow Q_{in} is given as:

$$895 \quad Q_{out}(t) = Q_{in} \cdot \frac{1}{K \cdot \Gamma(n)} \left(\frac{t}{K} \right)^{n-1} \cdot \exp(-t/K), \quad (118)$$

where $\Gamma(n)$ is the gamma function that replaces $(n-1)!$ to allow for non-integer values of n . K is the storage parameter, defined as the hydraulic retention time of a single linear reservoir segment of length L . It can be calculated as the average travel time of water through a single river segment:

$$K = \frac{L}{v}, \quad (119)$$

900 where v is the average flow velocity.

The river routing in LPJmL4 is calculated at a time step of $\Delta t = 3$ hour. We assume a globally constant flow velocity v of 1 m s^{-1} and a segment length L of 10 km to calculate the parameters n and K for each route between grid-cell mid points. At the start of the simulation, for each route the unit hydrograph for a rectangular input impulse of length Δt is determined from eq. (118).

905 Because eq. (118) assumes an instantaneous input impulse, we numerically determine the response to a rectangular input impulse by adding up the responses of a series of 100 consecutive instantaneous input impulses. From the obtained unit hydrograph, the sum of outflow during each subsequent time step Δt is recorded until 99% of the total input impulse has been released (maximum 24 time steps). During simulation thus determined response function is then used to calculate the convolution
 910 integral for the flow packages routed through the network. An efficient parallelisation of the river-routing scheme using global communicators of the MPI message passing library is described in Von Bloh et al. (2010).

2.6.4 Irrigation and dams

LPJmL4 explicitly accounts for human influences on the hydrological cycle by accounting for irriga-
 915 tion water abstraction, consumption and return flows, and non-agricultural water consumption from households, industry and livestock (HIL), as well as an implementation of reservoirs and dams.

Irrigation

LPJmL4 features a mechanistic representation of the world's most important irrigation systems (sur-
 face, sprinkler, drip), which is key to refined global simulations of agricultural water use as con-
 920 strained by biophysical processes and water tradeoffs along the river network. HIL water use in each grid cell is based on Flörke et al. (2013) (accounting for 201 km³ in the year 2000). We assume HIL water to be withdrawn prior to irrigation water. LPJmL4 comes with the first input dataset that details the global distribution of irrigation types for each cell and crop type (Jägermeyr et al., 2015). Irrigation water partitioning is dynamically calculated in coupling to the modelled water balance,
 925 and climate, soil, and vegetation properties. The spatial pattern of improved irrigation efficiencies are presented in Jägermeyr et al. (2015).

Irrigation water demand is withdrawn from available surface water, i.e. river discharge (see sec-
 tion 2.6.3), lakes, and reservoirs (section 2.6.5), and if not sufficient in the respective grid cell,
 requested from neighboring upstream cells. The amount of daily irrigation water requirements is
 930 based on the soil water deficit, resulting crop water demand (net irrigation requirements, NIR), and irrigation-system-specific application requirements (specified below). If soil moisture goes below the CFT-specific irrigation threshold (it), the total amount (daily gross irrigation requirements) is requested for abstraction. NIR is defined as the water needs of the top 50 cm soil layer to avoid water limitation to the crop. It is calculated to meet field capacity (W_{fc}) if the water supply (root-available
 935 soil water) falls below the atmospheric demand (potential evapotranspiration, see section 2.6.2) as:

$$\text{NIR} = W_{fc} - w_a - w_{ice}, \quad \text{NIR} \geq 0, \quad (120)$$

where w_a is actually available soil water and w_{ice} the frozen soil content in mm. Due to inefficien-
 cies in any irrigation system, excess water is required to meet the water demand of the crop. To this

end, we calculate system-specific conveyance efficiencies (E_c) and application requirements (AR),
 940 which lead to gross irrigation requirements (GIR, in mm):

$$\text{GIR} = \frac{\text{NIR} + \text{AR} - \text{Store}}{E_c}, \quad (121)$$

where Store stands in as a storage buffer (see also SI-Fig.S2 for conceptual description). For
 pressurized systems (sprinkler and drip), E_c is set to 0.95. For surface irrigation we link E_c to soil
 saturated hydraulic conductivity (K_s , see section 2.6.1), adopting E_c estimates from Brouwer et al.
 945 (1989). Half of conveyance losses are assumed to occur due to evaporation from open water bodies
 and the remainder is added to the return flow as drainage.

Indicative of application losses, AR represents the excess water needed to uniformly distribute
 irrigation across the field. AR is calculated as a system-specific scalar of the free water capacity:

$$\text{AR} = (W_{\text{sat}} - W_{\text{fc}}) \cdot d_u - \text{FW}, \quad \text{AR} \geq 0, \quad (122)$$

950 where d_u is the water distribution uniformity scalar, as a function of the irrigation system and FW
 represents the available free water (see sections 2.6 and 2.6.2) (see Jägermeyr et al. (2015) for de-
 tails).

Irrigation scheduling is controlled by Pr and the irrigation threshold (IT) that defines tolerable
 soil water depletion prior to irrigation (see SI-Table S14). Accessible irrigation water is subtracted
 955 by the precipitation amount. Irrigation water volumes that are not released (if $S > \text{IT}$) are added to
 Store and are available for the next irrigation event. Withdrawn irrigation volumes are subsequently
 reduced by conveyance losses.

Irrigation water application is assumed to occur below the canopy for surface and drip systems,
 and sprinkler systems above-canopy, which leads to interception losses (calculated as described
 960 above). Drip system are assumed to apply irrigation water localized to the plant root zone below
 the surface and thereby reduce soil evaporation by 60% (section 2.6.2). Note that drip systems are
 parameterized to represent a modest form of deficit irrigation, i.e. to save water and not to maximize
 yields. For detailed parameterization of the three irrigation systems implemented, see SI-Table 14
 and Jägermeyr et al. (2015).

965 **2.6.5 Dams, lakes and reservoirs**

The operation of large reservoirs affects the seasonal discharge patterns downstream of the dam, as
 well as the amount of water that is locally available for irrigation.

In LPJmL4, reservoirs are considered starting from the prescribed year they were built (Biemans
 et al., 2011). The reservoir is filled daily with discharge from upstream locations and with local
 970 precipitation. At the beginning of an operational year, which is defined as the first month when mean
 monthly inflow is lower than mean annual inflow, the actual storage in the reservoir is compared
 with the maximum storage capacity of the reservoir. The reservoir outflow factor of the following
 year is adjusted accordingly to compensate for interannual flow fluctuations.

Subsequently, a target release is defined based on the main purpose of the reservoir. Dams built primarily for irrigation are assumed to release their water proportionally to gross irrigation water demand downstream. Dams built primarily for other purposes (hydropower, flood control, etc.) are assumed to be designed for releasing a constant water volume throughout the year. The actual release from a reservoir is simulated to depend on its storage capacity relative to its inflow. If an irrigation purpose is defined for the reservoir, part of the outflow is diverted to irrigated lands downstream. Cells receive water from the reservoirs when the following conditions are met: the cells have a lower altitude than the cell containing the reservoir, and they are situated along the main river downstream or at maximum five cells upstream. Thus, a cell can receive water from multiple reservoirs.

As irrigation demands vary daily, water released from reservoirs can be stored in the conveyance system for up to five days. If the total irrigation water demand to a reservoir cannot be fulfilled, all requesting fields are supplied with the same fraction of their demand (see Biemans et al. (2011) for details).

2.7 Land use

Human land use is represented in LPJmL4 by dividing grid cells, which have the same climate and soil-texture input, into separate sub-units, referred to as stands. Stands are driven by the same input data, but changes in soil water and soil carbon are computed separately. When new stands are created, their soils are direct copies of the stand from which they are generated. If stands are merged, soil properties are averaged according to the two stands' size to maintain mass and energy balance. Natural vegetation (i.e. PFTs), agricultural crops (i.e. CFTs), managed grasslands and bioenergy plantations are represented on separate stands that can partly or fully cover any grid cell. The size of each stand is determined by the extent of land-use, defined by the input data prescribing fractions of each land-use type (crops, managed grassland, bioenergy plantations; all as rainfed and/or irrigated cultivation). All natural vegetation grows on a single natural vegetation stand on which all present PFTs compete for water and light. Agricultural crops are implemented as monocultures where only one single crop is cultivated and where there is no competition for resources with other stands (fields or natural vegetation) within that cell. For each crop and irrigation system (irrigated or rainfed) there can always only be one stand within one grid cell. For irrigated crops only one irrigation system (sprinkler, surface or drip, see section 2.6.4) can be selected.

At the beginning of each simulation year, present total agricultural land (all crops, all managed grasslands) is compared with land requirements for the year according to the input data. If there is too little agricultural land available, the needed fraction is cleared (liberated) from the natural vegetation stand, if there is too much, the excess agricultural land is abandoned and merged into the natural vegetation stand, leaving new space there for establishment of natural vegetation. Uncultivated cropland (i.e. outside the cropping period) is merged in set-aside stands, separated in irrigated and rainfed to prevent that irrigation water from irrigated stands is transferred to rainfed stands in

off-seasons. Set-aside land from irrigated agriculture is not irrigated during fallow periods, but is kept separate because of the soil water content that is enhanced through irrigation during the growing period. Land can be transferred between the two set-aside stands if the ratio between irrigated and rainfed cropland changes.

Depending on the scenario setting, if inter-cropping is assumed, a simple intercrop (grass) can be grown on the set-aside stand during the fallow period. Once a sowing date for a crop is reached (see section 2.7.1), the prescribed fraction of that crop and irrigation system is removed from the set-aside stand by copying the soil properties of the set-aside stand to the newly created stand and reducing the set-aside stand's size accordingly. The crop is then cultivated on that newly created stand and returned to the set-aside upon harvest of the crop.

2.7.1 Sowing dates

Sowing dates are simulated based on a set of rules depending on climate and crop specific thresholds as described in Waha et al. (2012). The start of the growing period is assumed to depend either on the onset of the wet season in tropical and subtropical regions or on the exceeding of a crop-specific temperature threshold for emergence in temperate regions.

We describe the intra-annual variability of precipitation and temperature in each location using variation coefficients for temperature (CV_{temp}) and precipitation (CV_{prec}), calculated from past monthly climate data. We assume temperature seasonality if CV_{temp} exceeds 0.01 and precipitation seasonality if CV_{prec} exceeds 0.4. Hence, four seasonality types can be differentiated (SI-Fig. S3):

1. temperature seasonality
2. precipitation seasonality, and
3. temperature and precipitation seasonality
4. no temperature and no precipitation seasonality

For locations with a combined temperature and precipitation seasonality, we additionally consider the mean temperature of the coldest month. If it exceeds 10°C, we assume absence of a cold season, i.e. the risk of frost occurrence is negligible, assuming temperatures are high enough to sow all year-round. Accordingly, precipitation seasonality defines the timing of sowing. If the mean temperature of the coldest month is equal to or below 10°C, temperature seasonality determines the timing of sowing. In regions with precipitation seasonality only, sowing date is at the onset of the main wet season. The precipitation-to-potential-evapotranspiration ratio is used to find moist and dry months in a year, as suggested by Thornthwaite (1948). The main wet season is identified by the largest sum of monthly precipitation-to-potential-evapotranspiration ratios of 4 consecutive months, because the length of that period aligns well with the length of the growing period of the majority of the simulated crops. In regions with bimodal rainfall patterns, the wet season starts with the first month of the

longest wet season. Crops are sown at the first wet day in the main wet season of the year i.e. when
1045 daily precipitation exceeds 0.1 mm. The onset of the growing period depends on temperature, if
temperature seasonality is detectable. Accordingly, crop emergence is related to temperature, and
thus sowing starts when daily average temperature exceeds a certain threshold (T_{fall} resp. T_{spring}
SI-Table S10). Locations without any temperature or precipitation seasonality e.g. in the wet tropics
crops are sown on the 1st of January. These assumptions lead to a possible adaptation of projected
1050 sowing dates.

2.7.2 Management and cropping intensity

Agricultural management is represented as a distinct set of options and a calibration of cropping
intensity. Explicit management options include:

1. cultivar choices, see section 2.4
- 1055 2. sowing dates, see section 2.7.1
3. irrigation shares and type section 2.6.4
4. residue removal section 2.4
5. intercropping section 2.7

Different irrigation systems can be represented as follows. For drip irrigation systems assuming lo-
1060 calised sub-surface water application, soil evaporation is reduced, so that only 40% of the applied
irrigation water are available for evaporation. Also, for rainwater management (see section 2.6.2),
soil evaporation can be reduced, mimicking agricultural management systems like mulching tech-
niques or conservation tillage (Jägermeyr et al., 2016). Secondly, to simulate improved rainwater
management (see section 2.6.1), the infiltration capacity can be increased, mimicking agricultural
1065 management practices such as different tillage systems or organic mulching (Jägermeyr et al., 2016).

Other than that, management options are not treated explicitly in the LPJmL4 model, that is, it
assumes no nutrient limitation to crop growth. Current management patterns, which is desirable for,
e.g., studies of the carbon or water cycle, can be represented by calibrating national cropping in-
tensity to FAO statistics as described in Fader et al. (2010). For this the maximum leaf area index
1070 (LAI_{max}), the harvest index parameter (hi_{opt}), and a scaling factor for scaling leaf-level photosyn-
thesis to stand level (α_a , Haxeltine and Prentice (1996)) are scaled in combination. LAI_{max} can
range between 1 (lowest intensity) and 5 for maize or 7 for all other crops (highest intensity), and α_a
ranges from 0.4 to 1.0. The Parameter hi_{opt} is crop-specific (see SI-Table 11), which can be reduced
by up to 20%, assuming there are more robust in return less productive varieties (Gosme et al., 2010).

1075 2.7.3 Managed grassland

On managed grassland stands, only herbaceous PFTs (TrH, TeH, PoH, see Table 1 for definition) can establish conditional to their bio-climatic limits (see SI-Table S4). If more than one herbaceous PFT establishes, these compete for light and water resources, but do not interact with other stands in that grid cell. In contrast to annual C allocation described above (see section 2.3.1), LPJmL4 simulates
 1080 managed grasslands and herbaceous biomass plantations (see section 2.7.4) with a daily allocation and turnover scheme where it dynamically computes leaf biomass per day as described below. It therefore enables to better represent the current phenological state and suitable times for harvest.

Daily allocation of managed grasslands

The allocation scheme is designed to distributes daily biomass increment to leaf and root biomass
 1085 in a way that best fulfils the predetermined ratio of leaf to root mass, l_r , for the whole plant. This allows for short-term deviations from allowed leaf-to-root-mass ratios l_r after harvest events, when much of the leaf biomass is removed. After a harvest event, NPP is first allocated to leaves until l_r is restored. If more CO_2 is assimilated than needed for maintenance respiration (i.e. NPP is positive), assimilated carbon B_I is allocated to the root (R) and the leaf carbon pool (L) by calculating the
 1090 respective increments (L_I, R_I) (eqs. (123) and (124)).

$$L_I = \min \left(B_I, \max \left(\frac{B_I + R - L/l_r}{1 + 1/l_r}, 0 \right) \right), \quad (123)$$

$$R_I = B_I - L_I. \quad (124)$$

In case of negative NPP (i.e. maintenance and growth respiration are larger than the GPP of that day), both compartments (leaves and roots) are reduced proportionally. l_r is scaled with a measure
 1095 of average growing-season water stress (mean of daily ratios of plant water supply to atmospheric demand, see section 2.6.2) to account for the functional relationship that plants allocate more carbon to roots under dry conditions.

$$l_r = l_{r_p} \cdot W_{\text{supply}}/W_{\text{demand}} \quad (125)$$

Grassland harvest routine

1100 LPJmL4 employs a default harvest scheme that attempts to approximate the actual global grassland production of 2.3 Gt DM (Herrero et al., 2013) while avoiding degradation. A harvest event of grass biomass occurs when leaf biomass increases over the previous month. Prior to the harvest event, grass leaf biomass (C_{leaf}) and the biomass after the last harvest event (MC_{leaf}) is summed up for all grass species at the managed grassland stand:

$$1105 \quad C_{\text{leaf}} = \sum_{\text{PFT}} B_{\text{mlPFT}}, \quad MC_{\text{leaf}} = \sum_{\text{PFT}} M_{\text{mlPFT}}. \quad (126)$$

On the last day of each month harvest occurs and the harvest index H_{frac} is determined depending on the leaf biomass C_{leaf} .

$$\text{If day } (C_{\text{leaf}} > MC_{\text{leaf}}) \quad H_{\text{frac}} = 1 - \frac{1000}{1000 + C_{\text{leaf}}} \quad (127)$$

Harvested biomass is taken from the leaf biomass of each herbaceous PFT. Depending on the amount of carbon in the leaves the harvested fraction is increasing (SI-Fig. S4) and biomass harvested depends on the present leaf carbon. In the absence of any detailed information about actual grassland management systems, this generic harvest routine does not represent specific management systems but allows for simulating regular harvest events (be it grazing or mowing) during productive periods of the year and the harvest amount is automatically adjusting to productivity.

2.7.4 Biomass plantations

Three biomass functional types (BFTs) were implemented in LPJmL4 (one fast-growing C4 grass, a temperate and a tropical tree) to allow for the simulation of dedicated biomass plantations (Beringer et al., 2011). These BFTs are generic representations of some of the most promising types of crops for the production of 2nd generation biofuels, biomaterials, or energy (possibly in combination with carbon capture and storage mechanisms). Their parametrization is partly identical to their natural PFT equivalents tropical C4 perennial grass, temperate broadleaved summergreen tree and tropical broadleaved raingreen tree yet with some important modifications to characterize the enhanced growth characteristics of these managed vegetation types (see SI-Table S12).

Woody energy crops are represented as short rotation coppice systems (SRC). In short intervals young tree stems are cut down to near ground stumps, implemented as regularly cycles (see SI-Table S13). A grown root system and nutrient storage in roots and stumps enables high yielding varieties of poplar, willow and Eucalyptus used for SRC to regrow forcefully in renewal years. Until plantations need to replanted after 40 years, several harvest cycles are possible. Under an effective pest and fire control on modern biomass plantations mortality and fire occurrence are reduced (to zero emissions) compared to natural vegetation.

3 Modelling protocol

The objective of this publication is to provide a comprehensive description of the LPJmL4 model. Here we also provide some outputs from a standard simulation of the historic period 1901 to 2011, which is also the basis for the actual, more comprehensive model evaluation described in the companion paper (Schaphoff et al., under Revision).

3.1 Model setup and inputs

For this simulation, all carbon and water pools in the model are initialized to zero and a spinup simulation for 5000 years is conducted in which plants dynamically establish, grow and die following

the model dynamics described above. After the soil carbon equilibrium phase of 2310 simulation
 1140 years (see section 2.5), equilibrium soil carbon pool sizes are estimated and corrected, depending on
 organic matter input and mean decomposition rate in each grid cell, and after another 2690 spinup
 simulation years, all carbon pools have reached a dynamic equilibrium. In this phase LPJmL4 sim-
 ulates only natural vegetation. For the spinup simulation, we cyclically repeat the first 30 years of
 climate data input and prescribe atmospheric carbon dioxide concentrations at 278 ppm. During
 1145 the second phase of the spinup simulation, land use is introduced in the year 1700, from where it
 is updated annually according to the historic land-use data set (see Fader et al. (2010) and section
 3.1.2).

3.1.1 Climate, river routing and soil inputs

We use monthly climate data inputs on precipitation provided by the Global Precipitation Clima-
 1150 tology Centre (GPCC Full Data Reanalysis Version 7.0, (Becker et al., 2013)), daily mean tem-
 perature from Climatic Research Unit (CRU TS version 3.23 University of East Anglia Climatic
 Research Unit; Harris (2015); Harris et al. (2014)), shortwave downward radiation and net down-
 ward longwave radiation are reanalysis data from ERA-Interim (Dee et al., 2011), and the num-
 ber of wet days per months are derived synthetically as suggested by New et al. (2000), which
 1155 is used to allocate monthly precipitation to individual days. Precipitation are stochastically dis-
 aggregated while preserving monthly sum, temperature linearly interpolated (Gerten et al., 2004).
 Besides climate information, the model is forced with invariant information on the soil texture
 (FAO/IIASA/ISRIC/ISSCAS/JRC, 2012; Nachtergaele et al., 2008) and annual information on land-
 use from Fader et al. (2010), but now also explicitly describing sugar cane areas (see sectio 3.1.2).
 1160 For the SPITFIRE module LPJmL4 uses additional input. Dew point temperature is approximated
 from daily minimum temperature (Thonicke et al., 2010). Monthly average wind speeds are based on
 NCEP re-analysis data, which were regridded to CRU (NOAA-CIRES Climate Diagnostics Center,
 Boulder, Colorado, USA, Kalnay et al. (1996)).

For the transport directions we use the global ($0.5^\circ \times 0.5^\circ$). Atmospheric CO_2 concentrations
 1165 are used from the Mauna Loa station (NOAA/ESRL, <http://www.esrl.noaa.gov/gmd/ccgg/trends/>).
 Simulated Topological Network (STN-30) drainage direction map (Vorosmarty and Fekete, 2011).
 STN-30 organizes the Earth’s land area into drainage basins and provides the river network topology
 under the assumption that each grid cell can drain into one of the eight next-neighbor cells, as well
 as detailed information on water reservoirs obtained from GRanD database (Lehner et al., 2011),
 1170 including informations on storage capacity, total area and main purpose. Natural lakes are obtained
 from Lehner and Döll (2004). A complete overview of all inputs used here are given in SI-Table S2.

3.1.2 Land use input

In principle, LPJmL4 can be driven by any land-use data information. As the default land-use input file, the cropping areas for each of the CFTs are taken from MIRCA2000 (Portmann et al., 2010) which is a combination of crop-specific areas from Monfreda et al. (2008) and areas equipped for irrigation from 1900-2005 from (Siebert et al., 2015). Monfreda et al. (2008) defined 175 crops and this number was reduced to 26 in MIRCA2000, (therefore, e. g., the group pulses consist of 12 individual crops). These land-use patterns that have been derived from maximum monthly growing areas per crop and grid cell have been combined and if these areas add up to more than 1, i.e. when sequential cropping systems are present, total cropland fraction was reduced to not exceed physical land area in each pixel. A more detailed description of the procedure can be found in Fader et al. (2010). After the implementation of sugar cane as a 12th annual crop that is explicitly represented in LPJmL4 (Lapola et al., 2009), the standard land-use input data set was amended by subtracting the sugar cane areas from the "others" band and implementing it as a separate input data band. All 16 input data bands (CFTs 1-12, others, managed grassland, bioenergy grass and bioenergy trees) are included four times in the data set, the first 16 bands representing purely rainfed agricultural areas, the second, third and fourth set representing irrigated areas of these land-use types for surface, sprinkler and drip irrigation respectively.

3.2 Standard outputs

The multiple aspects of the terrestrial biosphere and hydrosphere that are implemented in LPJmL4 allow for assessing multiple processes from natural and managed land which span ecological, hydrological and agricultural components. The consistent single modelling framework allows for analyzing interactions among these multiple sectors from local to global scale spanning seasons to centuries. A list of the key parameters calculated by LPJmL4 are shown in SI-Table S3.

Being driven by climate and land-use data LPJmL4 can be applied to quantify both climatic and anthropogenic impacts on the terrestrial biosphere. Computed dynamics of biogeochemical and hydrological processes thus arise from vegetation dynamics in natural ecosystems under climate change and elevated atmospheric CO₂ concentrations, land-use change as well as climate and management driven changes in managed ecosystems. Each grid cell can be dominated by managed land (croplands and pastures) but still contain fractions of natural vegetation, and vice versa. We here apply the anthromes concept (cf. Ellis et al. (2010)) to illustrate the global distribution of natural vegetation and managed land as simulated by LPJmL4 (Fig. 2). We use simulated vegetation carbon, potential evapotranspiration, foliar projective cover for each PFT, managed grassland and CFT, and combine it with climate input data to map natural biomes and anthromes at the global scale (see Boit et al. (2016) for the algorithm description). The composition of natural ecosystems is dynamically computed by LPJmL4, as the different PFTs compete with each other. Bounded by the bioclimatic limits,

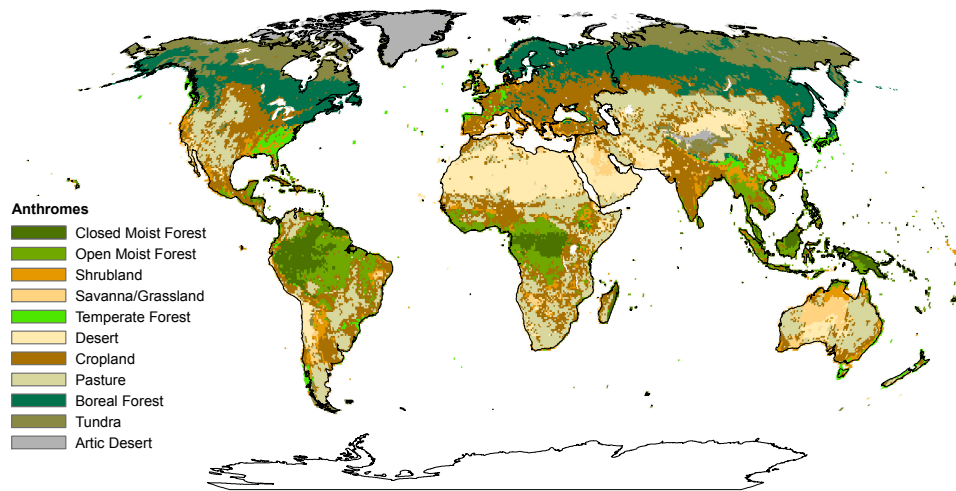
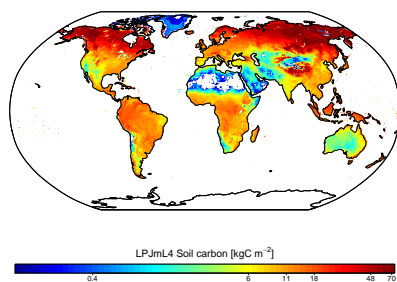


Figure 2. Global anthrome classes in the potential natural vegetation and the agricultural areas. Anthrome classes are defined using coverage with (a) types of dominating natural vegetation (e.g. tropical forest), (b) dominant agricultural usage (e.g. cropland) and external drivers (e.g. temperature)..

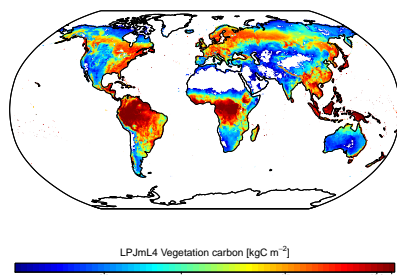
the modelled global distribution of forests, shrubland and natural grasslands as well as the spatial extent of polar and alpine ecosystems and deserts are in broad agreement with the biomes identified by Olson et al. (2001). The integrated mapping of biomes and anthromes underlines the extent of anthropogenic impacts on the terrestrial biosphere and how much of the potential vegetation coverage is left. By applying the land-use input (see section 3.1.2); LPJmL4 simulates cropland in 27% and pasture in 16% of the ice-free global land area. Simulating biophysical and biochemical processes in densely populated or urban areas could be considered in future model developments of LPJmL4 given their large spatial extent (Ellis et al., 2010).

Carbon and water fluxes, productivity and harvest of crops and managed grasslands have also been quantified. Results show that soils are the largest carbon pool of the terrestrial biosphere, with highest amounts of more than 60 kgC m^{-2} in the boreal zone, most notably in permafrost soils (see Fig. 3a and 3b). Vegetation carbon pools are largest in the tropics with almost 20 kgC m^{-2} and in the temperate zone with about 8 kgC m^{-2} . The large vegetation carbon pools are a result of high net primary productivity (NPP) in tropical and subtropical ecosystems, which process about 1000 to $1200 \text{ gC m}^{-2} \text{ a}^{-1}$, respectively (see Fig. 4b). Fire carbon emissions are highest in the tropics as

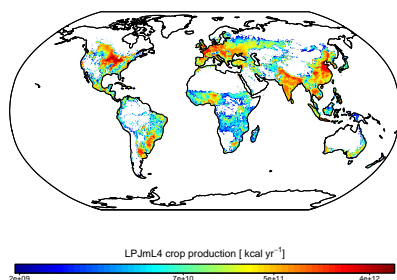
a result of high ignition probability and high biomass values simulated (Fig. 4c). Crop productivity is determined by climatic conditions and management strategies and is currently highest in the temperate zone of North America and Europe, but also in regions in eastern China, the irrigated Ganges Valley in India and temperate South America (see Fig. 3c).



(a)

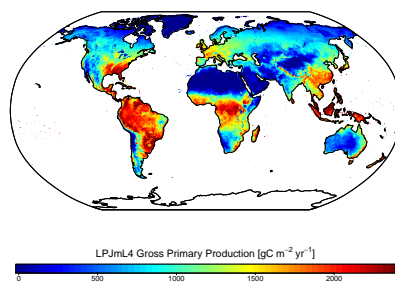


(b)

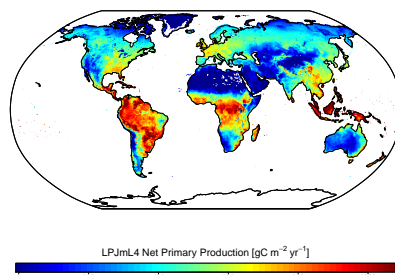


(c)

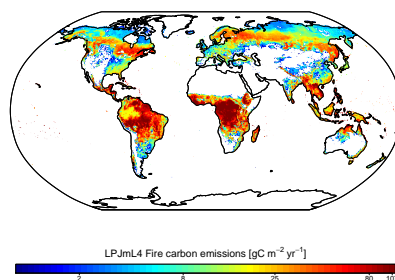
Figure 3. Soil (a) and vegetation (b) carbon pool and cumulative crop production (c) computed by LPJmL4 as an average of the time period 1996-2005. Note: Values plotted on a logarithmic scale.



(a)



(b)



(c)

Figure 4. Annual GPP (a), NPP (b) and fire carbon emissions (c) computed by LPJmL4 as an average of the time period 1996-2005. Note: Values for fire carbon emissions are plotted on a logarithmic scale.

Over the 20th century, changes in climate, land use and atmospheric CO₂ concentrations had distinct effects on the terrestrial carbon stocks and fluxes. Global vegetation carbon declined by 20 PgC after 1940 and is rising again since 2005, whereas carbon stored in soil and litter increased constantly over the simulation period (see SI-Fig. S5). GPP, NPP and heterotrophic respiration also follow this trend and show considerable interannual variability, while fire-related carbon emissions declined in the 1970s and remained relatively stable thereafter. Interception and runoff also show a positive trend, while evaporation from bare soil decreased (see SI-Fig. S5).

4 Discussion

Previous versions of LPJmL4 were used in a large number of applications to evaluate vegetation, water and carbon dynamics under current and future climate and land use change. In total almost a hundred papers were published since 2007 which cover a wide range of model developments and process analyses (see references in SI-Table S1) including 18 studies that describe significant model developments. The majority of the studies deal with modelling human land-use, with a focus on different crop types ($N = 54$ studies), managed grasslands ($N = 21$) and agricultural water use ($N = 18$). Most were conducted at global scale ($N = 58$) but also at regional scales, mainly for Europe, Africa and Amazonia. Many studies ($N = 43$) investigate potential future impacts of climate change, while the remaining studies evaluate effects of current or historic climates. In the following, we will highlight some of the most important previous publications using LPJmL, describing the most prominent fields of model application.

An important field of LPJmL model application, testing and subsequent development is the analysis of historic events. LPJmL simulation results contributed to the analysis of extreme event impacts on the biosphere globally (Zscheischler et al., 2014a, b) and at pan-European scale (Rammig et al., 2015; Rolinski et al., 2015). In combination with remote sensing and eddy flux data, LPJmL has been applied to estimate ecosystem respiration (Jägermeyr et al., 2014) and productivity (e.g., Jung et al., 2008). The increasing trend in atmospheric CO₂ amplitude could be explained by increasing productivity in subarctic and boreal forest ecosystems and less so by increases in agricultural land and productivity (Forkel et al., 2016) as simulated building on the improved phenology scheme by Forkel et al. (2014). The coupling of LPJmL to the climate model SPEEDY allowed for investigating climate-vegetation feedbacks from land-use change (e.g., Strengers et al., 2010; Boisier et al., 2012).

Studies on agricultural water consumption (Rost et al., 2008) and virtual water contents and water footprints for crops (Fader et al., 2010, 2011) have contributed to illuminate the role of agriculture in human water consumption.

The majority of LPJmL applications addresses the impacts of climate and land-use change on various biogeochemical and ecosystem properties of the terrestrial biosphere. To evaluate impacts of climate change on ecosystem processes and the carbon cycle at the global scale, Heyder et al. (2011)

performed a risk analysis of terrestrial ecosystems based on an integrated metric that considered joint changes in carbon and water fluxes, carbon stocks and vegetation structure. Applying CMIP3 and CMIP5 climate change scenarios to LPJmL, severe impacts for the terrestrial biosphere were found when global warming levels exceed 3K local temperature in cold and tropical biomes and 4K in the temperate biome (Heyder et al., 2011; Ostberg et al., 2013). The coupling of LPJmL to the integrated assessment model IMAGE allowed to evaluate feedbacks between land-use change, the carbon balance and climate change, so that the dynamics of a potential reversal of the terrestrial carbon balance could be assessed (Müller et al., 2016). Also economic feedbacks of agricultural production were evaluated by coupling LPJmL to the agro-economic model MAgPIE (Lotze-Campen et al., 2008), for example measures of land-use protection for climate change mitigation (Popp et al., 2014).

Regional climate-change applications investigated the role of CO₂ fertilization on Amazon rain-forest stability (Rammig et al., 2010), and analyzed additional threats arising from tropical deforestation (Gumpenberger et al., 2010; Poulter et al., 2010). Boit et al. (2016) applied the anthromes concept to LPJmL simulation results to differentiate the relative importance of future climate vs. land-use change in Latin America. In that study, land-use change was identified as the main driver of biome shifts and biome degradation early in the 21st century while climate change impacts would dominate the second half. LPJmL simulations also showed that in the boreal forest and Arctic ecosystems, 100 years of future climate change might destabilize carbon stored in permafrost soils over several centuries due to feedbacks between permafrost and dynamic vegetation (Schaphoff et al., 2013).

Several studies used LPJmL to investigate water limitations of natural ecosystems Gerten et al. (2013) and of food production in particular (e.g., Gerten et al., 2008; Biemans et al., 2011, 2013). Gerten et al. (2011) applied LPJmL to quantify "green" and "blue" water requirements for future food security, finding that water scarcity will increase in many countries, as confirmed by other studies prepared in the context of multi-model intercomparisons such as ISIMIP (e.g., Schewe et al., 2014). In the context of planetary boundaries, Gerten et al. (2013) and Steffen et al. (2015) have proposed sub-global modifications to the planetary boundary for freshwater use and Jägermeyr et al. (2017) quantified therefore needed environmental flow requirements in view of the Sustainable Development Goals. Future climate change effects on irrigation requirements were investigated by Konzmann et al. (2013); Jägermeyr et al. (2015); Fader et al. (2016).

LPJmL simulations have also shown (Asseng et al., 2015; Müller and Robertson, 2014; Rosenzweig et al., 2014) that climate change constitutes a major threat to agricultural productivity, especially in the tropics, but with large uncertainties regarding the benefits from elevated atmospheric CO₂ on crop water use and productivity (Müller et al., 2015; Deryng et al., 2016). It was also shown that the potential for major cereal crop production may decline in a future climate (Pugh et al., 2016b).

Building on the development of bioenergy plantations (Beringer et al., 2011), LPJmL was applied to analyse synergies and trade-offs of biomass plantations finding that demands for future bioenergy potentials implicit in climate mitigation and climate engineering portfolios could only be met at substantial environmental costs (Boysen et al., 2016; Heck et al., 2016).

The present publication describing the LPJmL4 model code and the companion paper (Schaphoff et al., under Revision) providing a thorough model evaluation are intended to serve as a comprehensive description of the current LPJmL4 model. The model code will be published under an open source license on a <https://gitlab.pik-potsdam.de/lpjml/LPJmL>. We hope that this will help to promote further development and improvement of LPJmL4 and to foster high-profile research in areas such as multi-sectoral climate change impacts, earth system dynamics, planetary boundaries, and SDGs. Besides the ongoing implementation of the dynamics of major nutrients, such as nitrogen, there are new plant physiological insights that have not yet found their way into LPJmL4 or related models (Pugh et al., 2016a; Rogers et al., 2017). The consistent coverage of natural and managed ecosystems as well as the full carbon and water dynamics linked by vegetation dynamics and land-use management is central to further development of LPJmL4.

5 Code and data availability

The model code of LPJmL4 is publicly available through PIK's gitlab server at <https://gitlab.pik-potsdam.de/lpjml/LPJmL> and an exact version of the code described here is archived under doi "xyz". The output data from the model simulations described here is available at the research data repository <http://dataservices.gfz-potsdam.de/portal/> under doi "ABC".

Acknowledgements. This study was supported by the German Federal Ministry of Education and Research's (BMBF's) "PalMod 2.3 Methankreislauf, Teilprojekt 2 Modellierung der Methanemissionen von Feucht- und Permafrostgebieten mit Hilfe von LPJmL", (FKZ 01LP1507C). A.R. and J.H. acknowledge funding from the Helmholtz Alliance "Remote Sensing and Earth System Dynamics. We thank the Climatic Research Unit for providing global gridded temperature input, the Global Precipitation Climatology Centre for providing precipitation input and the coordinators of ERA-Interim for providing shortwave downward radiation and net downward longwave radiation. Furthermore, we thank the authors of MIRCA2000 for providing land use input. We thank Jannes Breier and Nele Steinmetz for editorial help. Finally, the many people involved in the development, testing and application of LPJmL4 and its predecessors are gratefully acknowledged.

References

- Ahlström, A., Xia, J., Arneth, A., Luo, Y., and Smith, B.: Importance of vegetation dynamics for future terrestrial carbon cycling, *Environmental Research Letters*, 10, 054 019, doi:10.1088/1748-9326/10/5/054019, <http://stacks.iop.org/1748-9326/10/i=5/a=054019>, 2015.
- 1330 Allen, C. D., Macalady, A. K., Chenchouni, H., Bachelet, D., McDowell, N., Vennetier, M., Kitzberger, T., Rigling, A., Breshears, D. D., Hogg, E. T., Gonzalez, P., Fensham, R., Zhang, Z., Castro, J., Demidova, N., Lim, J.-H., Allard, G., Running, S. W., Semerci, A., and Cobb, N.: A global overview of drought and heat-induced tree mortality reveals emerging climate change risks for forests, *Forest Ecology and Management*, 259, 660–684, doi:doi: DOI: 10.1016/j.foreco.2009.09.001, <http://www.sciencedirect.com/science/article/B6T6X-4XH566S-1/2/d00470694e3e74d70cad5ca0883162ce>, 2010.
- 1335 Arnold, J. G., Williams, J. R., Nicks, A. D., and Sammons, N. B.: *SWRRB: A Basin Scale Simulation Model for Soil and Water Resources Management*, Texas A&M University Press, College Station, Texas, 1990.
- Asseng, S., Brisson, N., Basso, B., Martre, P., Aggarwal, P. K., Angulo, C., Bertuzzi, P., Biernath, C., Challinor, A. J., Doltra, J., Gayler, S., Goldberg, R., Grant, R., Heng, L., Hooker, J., Hunt, L. A., Ingwersen, J., Izaurrealde, R. C., Kersebaum, K. C., Müller, C., Kumar, S. N., Nendel, C., Leary, G. O., Olesen, J. E., Osborne, T. M., Palosuo, T., Priesack, E., Ripoche, D., Semenov, M. A., Shcherbak, I., Steduto, P., Stöckle, C., Stratonovitch, P., Streck, T., Supit, I., Tao, F., Travasso, M., Waha, K., Wallach, D., Williams, J. R., and Wolf, J.: Uncertainty in simulating wheat yields under climate change - Supplementary Information, *Nature Climate Change*, doi:10.1038/NCLIMATE1916, 2013.
- 1340 Asseng, S., Ewert, F., Martre, P., Rotter, R. P., Lobell, D. B., Cammarano, D., Kimball, B. A., Ottman, M. J., Wall, G. W., White, J. W., Reynolds, M. P., Alderman, P. D., Prasad, P. V. V., Aggarwal, P. K., Anothai, J., Basso, B., Biernath, C., Challinor, A. J., De Sanctis, G., Doltra, J., Fereres, E., Garcia-Vila, M., Gayler, S., Hoogenboom, G., Hunt, L. A., Izaurrealde, R. C., Jabloun, M., Jones, C. D., Kersebaum, K. C., Koehler, A.-K., Muller, C., Naresh Kumar, S., Nendel, C., O'Leary, G., Olesen, J. E., Palosuo, T., Priesack, E., Eyshi Rezaei, E., Ruane, A. C., Semenov, M. A., Shcherbak, I., Stockle, C., Stratonovitch, P., Streck, T., Supit, I., Tao, F., Thorburn, P. J., Waha, K., Wang, E., Wallach, D., Wolf, J., Zhao, Z., and Zhu, Y.: Rising temperatures reduce global wheat production, *Nature Clim. Change*, 5, 143–147, doi:10.1038/nclimate2470, 2015.
- 1350 Bassu, S., Brisson, N., Durand, J.-L., Boote, K., Lizaso, J., Jones, J. W., Rosenzweig, C., Ruane, A. C., Adam, M., Baron, C., Basso, B., Biernath, C., Boogaard, H., Conijn, S., Corbeels, M., Deryng, D., De Sanctis, G., Gayler, S., Grassini, P., Hatfield, J., Hoek, S., Izaurrealde, C., Jongschaap, R., Kemanian, A. R., Kersebaum, K. C., Kim, S.-H., Kumar, N. S., Makowski, D., Müller, C., Nendel, C., Priesack, E., Pravia, M. V., Sau, F., Shcherbak, I., Tao, F., Teixeira, E., Timlin, D., and Waha, K.: How do various maize crop models vary in their responses to climate change factors?, *Global change biology*, doi:10.1111/gcb.12520, 2014.
- Bayazitoglu, Y. and Özişik, M. N.: *Elements of heat transfer*, McGraw-Hill, New York, <https://www.crcpress.com/Elements-of-Heat-Transfer/Rathakrishnan/p/book/9781439878910>, 1988.
- 1360 Becker, A., Finger, P., Meyer-Christoffer, A., Rudolf, B., Schamm, K., Schneider, U., and Ziese, M.: A description of the global land-surface precipitation data products of the Global Precipitation Climatology Centre with sample applications including centennial (trend) analysis from 1901–present, *Earth System Science Data*, 5, 71–99, doi:10.5194/essd-5-71-2013, <http://www.earth-syst-sci-data.net/5/71/2013/>, 2013.

- 1365 Beer, C., Lucht, W., Gerten, D., Thonicke, K., and Schmulius, C.: Effects of soil freezing and thawing on vegetation carbon density in Siberia: A modeling analysis with the Lund-Potsdam-Jena Dynamic Global Vegetation Model (LPJ-DGVM), *Global Biogeochem. Cycles*, 21, GB1012, doi:10.1029/2006GB002760, 2007.
- Beringer, T., Lucht, W., and Schaphoff, S.: Bioenergy production potential of global biomass plantations under environmental and agricultural constraints, *GCB Bioenergy*, 3, 299–312, doi:10.1111/j.1757-1707.2010.01088.x, 2011.
- 1370 Biemans, H., Haddeland, I., Kabat, P., Ludwig, F., Hutjes, R. W. a., Heinke, J., von Bloh, W., and Gerten, D.: Impact of reservoirs on river discharge and irrigation water supply during the 20th century, *Water Resources Research*, 47, W03 509, doi:10.1029/2009WR008929, 2011.
- 1375 Biemans, H., Speelman, L., Ludwig, F., Moors, E., Wiltshire, A., Kumar, P., Gerten, D., and Kabat, P.: Future water resources for food production in five South Asian river basins and potential for adaptation — A modeling study, *Changing water resources availability in Northern India with respect to Himalayan glacier retreat and changing monsoon patterns: consequences and adaptation*, 468–469, Supplement, S117–S131, doi:10.1016/j.scitotenv.2013.05.092, 2013.
- 1380 Boisier, J., de Noblet-Ducoudré, N., Pitman, A., Cruz, F., Delire, C., van den Hurk, B., van der Molen, M., Müller, C., and Voldoire, A.: Attributing the biogeophysical impacts of Land-Use induced Land-Cover Changes on surface climate to specific causes. Results from the first LUCID set of simulations, *J. Geophys. Res.*, 117, D12 116, doi:10.1029/2011JD017106, 2012.
- Boit, A., Sakschewski, B., Boysen, L., Cano-Crespo, A., Clement, J., Garcia-alaniz, N., Kok, K., Kolb, M., 1385 Langerwisch, F., Rammig, A., Sachse, R., van Eupen, M., von Bloh, W., Clara Zemp, D., and Thonicke, K.: Large-scale impact of climate change vs. land-use change on future biome shifts in Latin America, *Global Change Biology*, 22, 3689–3701, doi:10.1111/gcb.13355, 2016.
- Bondeau, A., Smith, P., Zaehle, S., Schaphoff, S., Lucht, W., Cramer, W., Gerten, D., Lotze-Campen, Hermann, Müller, C., Reichstein, M., and Smith, B.: Modelling the role of agriculture for the 20th century global 1390 terrestrial carbon balance, *Global Change Biology*, 13, 679–706, doi:10.1111/j.1365-2486.2006.01305.x, 2007.
- Boysen, L. R., Lucht, W., Gerten, D., and Heck, V.: Impacts devalue the potential of large-scale terrestrial CO₂ removal through biomass plantations, *Environmental Research Letters*, 11, 095 010, doi:10.1088/1748-9326/11/9/095010, 2016.
- 1395 Brouwer, C., Prins, K., and Heibloem, M.: Irrigation Water Management : Irrigation Scheduling. Training manual no. 4, Tech. Rep. 4, FAO Land and Water Development Division, Rome, Italy, <http://www.fao.org/docrep/t7202e/t7202e00.htm>, 1989.
- Brovkin, V., van Bodegom, P. M., Kleinen, T., Wirth, C., Cornwell, W. K., Cornelissen, J. H. C., and Kattge, J.: Plant-driven variation in decomposition rates improves projections of global litter stock distribution, *Biogeosciences*, 9, 565–576, doi:10.5194/bg-9-565-2012, 2012.
- 1400 Canadell, J. G., Le Quéré, C., Raupach, M. R., Field, C. B., Buitenhuis, E. T., Ciais, P., Conway, T. J., Gillett, N. P., Houghton, R. A., and Marland, G.: Contributions to accelerating atmospheric CO₂ growth from economic activity, carbon intensity, and efficiency of natural sinks, *Proceedings of the National Academy of Sciences*, 104, 18 866–18 870, doi:10.1073/pnas.0702737104, 2007.

- 1405 Carvalho, N., Forkel, M., Khomik, M., Bellarby, J., Jung, M., Migliavacca, M., Mu, M., Saatchi, S., Santoro, M., Thurner, M., Weber, U., Ahrens, B., Beer, C., Cescatti, A., Randerson, J. T., and Reichstein, M.: Global covariation of carbon turnover times with climate in terrestrial ecosystems, *Nature*, 514, 213–217, 10.1038/nature13731, 2014.
- Christian, H. J., Blakeslee, R. J., Boccippio, D. J., Boeck, W. L., Buechler, D. E., Driscoll, K. T., Goodman, S. J.,
1410 Hall, J. M., Koshak, W. J., Mach, D. M., and Stewart, M. F.: Global frequency and distribution of lightning as observed from space by the Optical Transient Detector, *Journal of Geophysical Research: Atmospheres*, 108, ACL 4–1, doi:10.1029/2002JD002347, 2003.
- Collatz, G., Ball, J., Grivet, C., and Berry, J. A.: Physiological and environmental regulation of stomatal conductance, photosynthesis and transpiration: a model that includes a laminar boundary layer, *Agricultural and*
1415 *Forest Meteorology*, 54, 107–136, doi:10.1016/0168-1923(91)90002-8, 1991.
- Collatz, G. J., Ribas-Carbo, M., and Berry, J. A.: Coupled Photosynthesis-Stomatal Conductance Model for Leaves of C4 Plants, *Functional Plant Biology*, 19, 519–538, doi:10.1071/pp9920519, 1992.
- Cosby, B. J., Hornberger, G. M., Clapp, R. B., and Ginn, T. R.: A Statistical Exploration of the Relationships of Soil Moisture Characteristics to the Physical Properties of Soils, *Water Resour. Res.*, 20, 682–690,
1420 doi:10.1029/WR020i006p00682, 1984.
- Cramer, W., Bondeau, A., Woodward, F. I., Prentice, I. C., Betts, R. A., Brovkin, V., Cox, P. M., Fisher, V., Foley, J. A., Friend, A. D., Kucharik, C., Lomas, M. R., Ramankutty, N., Sitch, S., Smith, B., White, A., and Young-Molling, C.: Global response of terrestrial ecosystem structure and function to CO₂ and climate change: results from six dynamic global vegetation models, *Global Change Biology*, 7, 357–373, doi:10.1046/j.1365-
1425 2486.2001.00383.x, <http://dx.doi.org/10.1046/j.1365-2486.2001.00383.x>, 01261, 2001.
- Dee, D. P., Uppala, S. M., Simmons, A. J., Berrisford, P., Poli, P., Kobayashi, S., Andrae, U., Balmaseda, M. A., Balsamo, G., Bauer, P., Bechtold, P., Beljaars, A. C. M., van de Berg, L., Bidlot, J., Bormann, N., Delsol, C., Dragani, R., Fuentes, M., Geer, A. J., Haimberger, L., Healy, S. B., Hersbach, H., Hólm, E. V., Isaksen, I., Kållberg, P., Köhler, M., Matricardi, M., McNally, A. P., Monge-Sanz, B. M., Morcrette, J.-
1430 J., Park, B.-K., Peubey, C., de Rosnay, P., Tavolato, C., Thépaut, J.-N., and Vitart, F.: The ERA-Interim reanalysis: configuration and performance of the data assimilation system, *Quarterly Journal of the Royal Meteorological Society*, 137, 553–597, doi:10.1002/qj.828, <http://dx.doi.org/10.1002/qj.828>, 2011.
- Deryng, D., Elliott, J., Folberth, C., Muller, C., Pugh, T. A. M., Boote, K. J., Conway, D., Ruane, A. C., Gerten, D., Jones, J. W., Khabarov, N., Olin, S., Schaphoff, S., Schmid, E., Yang, H., and Rosenzweig, C.: Regional
1435 disparities in the beneficial effects of rising CO₂ concentrations on crop water productivity, *Nature Clim. Change*, advance online publication, doi:10.1038/nclimate2995, 2016.
- Ellis, E. C., Klein Goldewijk, K., Siebert, S., Lightman, D., and Ramankutty, N.: Anthropogenic transformation of the biomes, 1700 to 2000, *Global Ecology and Biogeography*, 19, 589–606, doi:10.1111/j.1466-8238.2010.00540.x, 2010.
- 1440 Fader, M., Rost, S., Müller, C., Bondeau, A., and Gerten, D.: Virtual water content of temperate cereals and maize: Present and potential future patterns, *Journal of Hydrology*, 384, 218–231, doi:10.1016/j.jhydrol.2009.12.011, 2010.

- Fader, M., Gerten, D., Thammer, M., Heinke, J., Lotze-Campen, H., Lucht, W., and Cramer, W.: Internal and external green-blue agricultural water footprints of nations, and related water and land savings through trade, *Hydrology and Earth System Sciences Discussions*, 8, 483–527, doi:10.5194/hessd-8-483-2011, 2011.
- Fader, M., von Bloh, W., Shi, S., Bondeau, A., and Cramer, W.: Modelling Mediterranean agro-ecosystems by including agricultural trees in the LPJmL model, *Geoscientific Model Development*, 8, 3545–3561, doi:10.5194/gmd-8-3545-2015, 2015.
- Fader, M., Shi, S., von Bloh, W., Bondeau, A., and Cramer, W.: Mediterranean irrigation under climate change: more efficient irrigation needed to compensate for increases in irrigation water requirements, *Hydrology and Earth System Sciences*, 20, 953–973, doi:10.5194/hess-20-953-2016, 2016.
- FAO/IIASA/ISRIC/ISSCAS/JRC: Harmonized World Soil Database (version 1.2)., <http://www.iiasa.ac.at/Research/LUC/External-World-soil-database/HTML/>, 2012.
- Farquhar, G. D. and von Caemmerer, S.: Modelling of Photosynthetic Response to Environmental Conditions, in: *Physiological Plant Ecology II: Water Relations and Carbon Assimilation*, edited by Lange, O. L., Nobel, P. S., Osmond, C. B., and Ziegler, H., pp. 549–587, Springer Berlin Heidelberg, Berlin, Heidelberg, http://dx.doi.org/10.1007/978-3-642-68150-9_17, 1982.
- Farquhar, G. D., Caemmerer, S. V., and Berry, J. A.: A Biochemical Model of Photosynthetic CO₂ Assimilation in Leaves of C₃ Species, *Planta*, 90, 78–90, doi:10.1007/BF00386231, 1980.
- Federer, C. A.: Transpirational Supply and Demand - Plant, Soil, and Atmospheric Effects Evaluated by Simulation, *Water Resources Research*, 18, 355–362, doi:10.1029/WR018i002p00355, 1982.
- Flörke, M., Kynast, E., Bärlund, I., Eisner, S., Wimmer, F., and Alcamo, J.: Domestic and industrial water uses of the past 60 years as a mirror of socio-economic development: A global simulation study, *Global Environmental Change*, 23, 144–156, doi:10.1016/j.gloenvcha.2012.10.018, 2013.
- Forkel, M., Carvalhais, N., Schaphoff, S., v. Bloh, W., Migliavacca, M., Thurner, M., and Thonicke, K.: Identifying environmental controls on vegetation greenness phenology through model–data integration, *Biogeosciences*, 11, 7025–7050, doi:10.5194/bg-11-7025-2014, <http://www.biogeosciences.net/11/7025/2014/>, 2014.
- Forkel, M., Carvalhais, N., Rödenbeck, C., Keeling, R., Heimann, M., Thonicke, K., Zaehle, S., and Reichstein, M.: Enhanced seasonal CO₂ exchange caused by amplified plant productivity in northern ecosystems, *Science*, 351, 696, doi:10.1126/science.aac4971, <http://science.sciencemag.org/content/351/6274/696.abstract>, 2016.
- Friedlingstein, P., Meinshausen, M., Arora, V. K., Jones, C. D., Anav, A., Liddicoat, S. K., and Knutti, R.: Uncertainties in CMIP5 Climate Projections due to Carbon Cycle Feedbacks, *Journal of Climate*, 27, 511–526, doi:10.1175/JCLI-D-12-00579.1, 2013.
- Friend, A. D., Lucht, W., Rademacher, T. T., Keribin, R., Betts, R., Cadule, P., Ciais, P., Clark, D. B., Dankers, R., Falloon, P. D., Ito, A., Kahana, R., Kleidon, A., Lomas, M. R., Nishina, K., Ostberg, S., Pavlick, R., Peylin, P., Schaphoff, S., Vuichard, N., Warszawski, L., Wiltshire, A., and Woodward, F. I.: Carbon residence time dominates uncertainty in terrestrial vegetation responses to future climate and atmospheric CO₂, *Proceedings of the National Academy of Sciences*, 111, 3280–3285, doi:10.1073/pnas.1222477110, 2014.
- Gelfan, A. N., Pomeroy, J. W., and Kuchment, L. S.: Modeling Forest Cover Influences on Snow Accumulation, Sublimation, and Melt, *Journal of Hydrometeorology*, 5, 785–803, doi:10.1175/1525-

7541(2004)005<0785:MFCIOS>2.0.CO;2, [https://doi.org/10.1175/1525-7541\(2004\)005<0785:MFCIOS>2.0.CO;2](https://doi.org/10.1175/1525-7541(2004)005<0785:MFCIOS>2.0.CO;2), 2004.

- 1485 Gerten, D., Schaphoff, S., Haberlandt, U., Lucht, W., and Sitch, S.: Terrestrial vegetation and water balance—hydrological evaluation of a dynamic global vegetation model, *Journal of Hydrology*, 286, 249–270, doi: DOI: 10.1016/j.jhydrol.2003.09.029, <http://www.sciencedirect.com/science/article/B6V6C-4B9D86T-3/2/aa123e159770c269994f0d74c5edc335>, 2004.
- Gerten, D., Rost, S., von Bloh, W., and Lucht, W.: Causes of change in 20th century global river discharge, *Geophysical Research Letters*, 35, 1–5, doi:10.1029/2008GL035258, 2008.
- 1490 Gerten, D., Heinke, J., Hoff, H., Biemans, H., Fader, M., and Waha, K.: Global water availability and requirements for future food production, *Journal of Hydrometeorology*, p. 110531121709055, doi:10.1175/2011JHM1328.1, 2011.
- Gerten, D., Hoff, H., Rockström, J., Jägermeyr, J., Kummu, M., and Pastor, A. V.: Towards a revised planetary boundary for consumptive freshwater use: role of environmental flow requirements, *Current Opinion in Environmental Sustainability*, 5, 551–558, doi:10.1016/j.cosust.2013.11.001, 2013.
- 1495 Gosme, M., Suffert, F., and Jeuffroy, M.-H.: Intensive versus low-input cropping systems: What is the optimal partitioning of agricultural area in order to reduce pesticide use while maintaining productivity?, *Agricultural Systems*, 103, 110–116, doi:10.1016/j.agsy.2009.11.002, 2010.
- 1500 Gumpenberger, M., Vohland, K., Heyder, U., Poulter, B., Macey, K., Anja Rammig, Popp, A., and Cramer, W.: Predicting pan-tropical climate change induced forest stock gains and losses—implications for REDD, *Environmental Research Letters*, 5, 014 013, doi:10.1088/1748-9326/5/1/014013, 2010.
- Haberl, H., Erb, K.-H., Krausmann, F., Bondeau, A., Lauk, C., Müller, C., Plutzar, C., and Steinberger, J. K.: Global bioenergy potentials from agricultural land in 2050: Sensitivity to climate change, diets and yields, *Biomass and bioenergy*, 35, 4753–4769, doi:10.1016/j.biombioe.2011.04.035, 2011.
- 1505 Harris, I., Jones, P., Osborn, T., and Lister, D.: Updated high-resolution grids of monthly climatic observations – the CRU TS3.10 Dataset, *International Journal of Climatology*, 34, 623–642, doi:10.1002/joc.3711, 2014.
- Haxeltine, A. and Prentice, I. C.: A General Model for the Light-Use Efficiency of Primary Production, *Functional Ecology*, 10, 551–561, doi:10.2307/2390165, 1996.
- 1510 Heck, V., Gerten, D., Lucht, W., and Boysen, L. R.: Is extensive terrestrial carbon dioxide removal a ‘green’ form of geoengineering? A global modelling study, *Global and Planetary Change*, 137, 123–130, doi:10.1016/j.gloplacha.2015.12.008, 2016.
- Herrero, M., Havlík, P., Valin, H., Notenbaert, A., Rufino, M. C., Thornton, P. K., Blümmel, M., Weiss, F., Grace, D., and Obersteiner, M.: Biomass use, production, feed efficiencies, and greenhouse gas emissions from global livestock systems., *Proceedings of the National Academy of Sciences of the United States of America*, doi:10.1073/pnas.1308149110, 2013.
- 1515 Heyder, U., Schaphoff, S., Gerten, D., and Lucht, W.: Risk of severe climate change impact on the terrestrial biosphere, *Environmental Research Letters*, 6, 034 036, doi:10.1088/1748-9326/6/3/034036, <http://stacks.iop.org/1748-9326/6/i=3/a=034036>, 2011.
- 1520 Huang, S., Titus, S. J., and Wiens, D. P.: Comparison of nonlinear height–diameter functions for major Alberta tree species, *Canadian Journal of Forest Research*, 22, 1297–1304, doi:10.1139/x92-172, 1992.

- Humpenöder, F., Popp, A., Dietrich, J. P., Klein, D., Lotze-Campen, H., Bonsch, M., Bodirsky, B. L., Weindl, I., Stevanovic, M., and Müller, C.: Investigating afforestation and bioenergy CCS as climate change mitigation strategies, *Environmental Research Letters*, 9, 064 029, doi:10.1088/1748-9326/9/6/064029, 2014.
- 1525 IPCC: Summary for Policymakers, in: *Climate Change 2014: Mitigation of Climate Change. Contribution of Working Group III to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change*, edited by Edenhofer, O., Pichs-Madruga, R., Sokona, Y., Farahani, E., Kadner, S., Seyboth, K., Adler, A., Baum, I., Brunner, S., Eickemeier, P., Kriemann, B., Savolainen, J., Schlömer, S., von Stechow, C., Zwickel, T., and Minx, J. C., pp. 1–32, Cambridge University Press, Cambridge, UK and New York, NY, USA, 2014.
- 1530 Jackson, R., Canadell, J., Ehleringer, J., Mooney, H., Sala, O., and Schulze, E.: A global analysis of root distributions for terrestrial biomes, *Oecologia*, 108, 389–411, doi:10.1007/BF00333714, 1996.
- Jägermeyr, J., Gerten, D., Lucht, W., Hostert, P., Migliavacca, M., and Nemani, R.: A high-resolution approach to estimating ecosystem respiration at continental scales using operational satellite data, *Global change biology*, 20, 1191–1210, doi:10.1111/gcb.12443, 2014.
- 1535 Jägermeyr, J., Gerten, D., Heinke, J., Schaphoff, S., Kummu, M., and Lucht, W.: Water savings potentials of irrigation systems: global simulation of processes and linkages, *Hydrology and Earth System Sciences*, 19, 3073–3091, doi:10.5194/hess-19-3073-2015, 2015.
- Jägermeyr, J., Gerten, D., Schaphoff, S., Heinke, J., Lucht, W., and Rockström, J.: Integrated crop water management might sustainably halve the global food gap, *Environmental Research Letters*, 11, 025 002, doi:10.1088/1748-9326/11/2/025002, 2016.
- 1540 Jägermeyr, J., Pastor, A., Biemans, h., and Gerten, D.: Reconciling irrigated food production with environmental flows for Sustainable Development Goals implementation, *Nature Communications*, 8, doi:10.1038/ncomms15900, 2017.
- Jarvis, P. G. and McNaughton, K.: Stomatal control of transpiration: scaling up from leaf to region, *Advances in ecological research*, 15, 1–49, doi:10.1016/S0065-2504(08)60119-1, 1986.
- 1545 Jobbagy, E. G. and Jackson, R. B.: The Vertical Distribution of Soil Organic Carbon and Its Relation to Climate and Vegetation, *Ecological Applications*, 10, 423–436, doi:10.2307/2641104, 2000.
- Johansen, O.: Thermal conductivity of soils, Ph.D. thesis, University of Trondheim, Trondheim, Norway, <http://oai.dtic.mil/oai/oai?verb=getRecord&metadataPrefix=html&identifier=ADA044002>, 1977.
- 1550 Jolly, W. M., Nemani, R., and Running, S. W.: A generalized, bioclimatic index to predict foliar phenology in response to climate, *Global Change Biology*, 11, 619–632, doi:10.1111/j.1365-2486.2005.00930.x, 2005.
- Jung, M., Verstraete, M., Gobron, N., Reichstein, M., Papale, D., Bondeau, A., Robustelli, M., and Pinty, B.: Diagnostic assessment of European gross primary production, *Global Change Biology*, 14, 2349–2364, doi:10.1111/j.1365-2486.2008.01647.x, 2008.
- 1555 Kalnay, E., Kanamitsu, M., Kistler, R., Collins, W., Deaven, D., Gandin, L., Iredell, M., Saha, S., White, G., Woollen, J., Zhu, Y., Leetmaa, A., Reynolds, R., Chelliah, M., Ebisuzaki, W., Higgins, W., Janowiak, J., Mo, K. C., Ropelewski, C., Wang, J., Jenne, R., and Joseph, D.: The NCEP/NCAR 40-Year Reanalysis Project, *Bulletin of the American Meteorological Society*, 77, 437–471, doi:10.1175/1520-0477(1996)077<0437:TNYRP>2.0.CO;2, [https://doi.org/10.1175/1520-0477\(1996\)077<0437:TNYRP>2.0.CO;2](https://doi.org/10.1175/1520-0477(1996)077<0437:TNYRP>2.0.CO;2), 1996.
- 1560

- Kattge, J., Díaz, S., Lavorel, S., Prentice, I. C., Leadley, P., Bönisch, G., Garnier, E., Westoby, M., Reich, P. B., Wright, I. J., Cornelissen, J. H. C., Violle, C., Harrison, S. P., Van BODEGOM, P. M., Reichstein, M., Enquist, B. J., Soudzilovskaia, N. A., Ackerly, D. D., Anand, M., Atkin, O., Bahn, M., Baker, T. R., Baldocchi, D., Bekker, R., Blanco, C. C., Blonder, B., Bond, W. J., Bradstock, R., Bunker, D. E., Casanoves, F., Cavender-Bares, J., Chambers, J. Q., Chapin III, F. S., Chave, J., Coomes, D., Cornwell, W. K., Craine, J. M., Dobrin, B. H., Duarte, L., Durka, W., Elser, J., Esser, G., Estiarte, M., Fagan, W. F., Fang, J., Fernández-Méndez, F., Fidelis, A., Finegan, B., Flores, O., Ford, H., Frank, D., Freschet, G. T., Fyllas, N. M., Gallagher, R. V., Green, W. A., Gutierrez, A. G., Hickler, T., Higgins, S. I., Hodgson, J. G., Jalili, A., Jansen, S., Joly, C. A., Kerkhoff, A. J., Kirkup, D., Kitajima, K., Kleyer, M., Klotz, S., Knops, J. M. H., Kramer, K., Kühn, I., Kurokawa, H., Laughlin, D., Lee, T. D., Leishman, M., Lens, F., Lenz, T., Lewis, S. L., Lloyd, J., Llusià, J., Louault, F., Ma, S., Mahecha, M. D., Manning, P., Massad, T., Medlyn, B. E., Messier, J., Moles, A. T., Müller, S. C., Nadrowski, K., Naeem, S., Niinemets, Ü., Nöllert, S., Nüske, A., Ogaya, R., Oleksyn, J., Onipchenko, V. G., Onoda, Y., Ordoñez, J., Overbeck, G., Ozinga, W. A., Patiño, S., Paula, S., Pausas, J. G., Peñuelas, J., Phillips, O. L., Pillar, V., Poorter, H., Poorter, L., Poschlod, P., Prinzing, A., Proulx, R., Rammig, A., Reinsch, S., Reu, B., Sack, L., Salgado-Negret, B., Sardans, J., Shiodera, S., Shipley, B., Siefert, A., Sosinski, E., Soussana, J.-F., Swaine, E., Swenson, N., Thompson, K., Thornton, P., Waldram, M., Weiher, E., White, M., White, S., Wright, S. J., Yguel, B., Zaehle, S., Zanne, A. E., and Wirth, C.: TRY – a global database of plant traits, *Global Change Biology*, 17, 2905–2935, doi:10.1111/j.1365-2486.2011.02451.x, 2011.
- 1565 Kollas, C., Kersebaum, K. C., Nendel, C., Manevski, K., Müller, C., Palosuo, T., Armas-Herrera, C. M., Beaudoin, N., Bindi, M., Charfeddine, M., Conradt, T., Constantin, J., Eitzinger, J., Ewert, F., Ferrise, R., Gaiser, T., Cortazar-Atauri, I. G. d., Giglio, L., Hlavinka, P., Hoffmann, H., Hoffmann, M. P., Launay, M., Mandercheid, R., Mary, B., Mirschel, W., Moriondo, M., Olesen, J. E., Öztürk, I., Pacholski, A., Ripoché-Wachter, D., Roggero, P. P., Roncossek, S., Rötter, R. P., Ruget, F., Sharif, B., Trnka, M., Ventrella, D., Waha, K.,
- 1570 Wegehenkel, M., Weigel, H.-J., and Wu, L.: Crop rotation modelling—A European model intercomparison, *European Journal of Agronomy*, 70, 98–111, doi:10.1016/j.eja.2015.06.007, 2015.
- 1575 Konzmann, M., Gerten, D., and Heinke, J.: Climate impacts on global irrigation requirements under 19 GCMs, simulated with a vegetation and hydrology model, *Hydrological Sciences Journal*, 58, 88–105, doi:10.1080/02626667.2013.746495, 2013.
- 1590 Krysanova, V., Müller-Wohlfeil, D.-I., and Becker, A.: Development and test of a spatially distributed hydrological/water quality model for mesoscale watersheds, *Ecological Modelling*, 106, 261–289, doi:10.1016/S0304-3800(97)00204-4, <http://www.sciencedirect.com/science/article/pii/S0304380097002044>, 1998.
- Lapola, D. M., Oyama, M. D., and Nobre, C. A.: Exploring the range of climate biome projections for tropical South America: The role of CO₂ fertilization and seasonality, *Global Biogeochem. Cycles*, 23, GB3003, doi:10.1029/2008GB003357, 2009.
- 1595 Latham, D. and Williams, E.: Lightning and forest fires, *Forest fires: Behavior and ecological effects*, pp. 375–418, 2001.
- Latham, D. J. and Schlieter, J. A.: Ignition probabilities of wildland fuels based on simulated lightning discharges, US Department of Agriculture, Forest Service, Intermountain Research Station, 1989.

- 1600 Lawrence, D. and Slater, A.: Incorporating organic soil into a global climate model, *Climate Dynamics*, 30, 145–160, doi:10.1007/s00382-007-0278-1, 2008.
- Le Quéré, C., Moriarty, R., Andrew, R. M., Canadell, J. G., Sitch, S., Korsbakken, J. I., Friedlingstein, P., Peters, G. P., Andres, R. J., Boden, T. A., Houghton, R. A., House, J. I., Keeling, R. F., Tans, P., Arneeth, A., Bakker, D. C. E., Barbero, L., Bopp, L., Chang, J., Chevallier, F., Chini, L. P., Ciais, P., Fader, M., Feely, R. A., Gkritzalis, T., Harris, I., Hauck, J., Ilyina, T., Jain, A. K., Kato, E., Kitidis, V., Klein Goldewijk, K., Koven, C., Landschützer, P., Lauvset, S. K., Lefèvre, N., Lenton, A., Lima, I. D., Metzl, N., Millero, F., Munro, D. R., Murata, A., Nabel, J. E. M. S., Nakaoka, S., Nojiri, Y., O'Brien, K., Olsen, A., Ono, T., Pérez, F. F., Pfeil, B., Pierrot, D., Poulter, B., Rehder, G., Rödenbeck, C., Saito, S., Schuster, U., Schwinger, J., Séférian, R., Steinhoff, T., Stocker, B. D., Sutton, A. J., Takahashi, T., Tilbrook, B., van der Laan-Luijkx, I. T., van der Werf, G. R., van Heuven, S., Vandemark, D., Viovy, N., Wiltshire, A., Zaehle, S., and Zeng, N.: Global Carbon Budget 2015, *Earth System Science Data*, 7, 349–396, doi:10.5194/essd-7-349-2015, <http://www.earth-syst-sci-data.net/7/349/2015/>, 2015.
- 1610 Lehner, B. and Döll, P.: Development and validation of a global database of lakes, reservoirs and wetlands, *Journal of Hydrology*, 296, 1 – 22, doi:<http://dx.doi.org/10.1016/j.jhydrol.2004.03.028>, <http://www.sciencedirect.com/science/article/pii/S0022169404001404>, 2004.
- 1615 Lehner, B., Liermann, C. R., Revenga, C., Vörösmarty, C., Fekete, B., Crouzet, P., Döll, P., Endejan, M., Frenken, K., and Magome, J.: High-resolution mapping of the world's reservoirs and dams for sustainable river-flow management, *Frontiers in Ecology and the Environment*, 9, 494–502, doi:10.1890/100125, 2011.
- Liang, S., Stroeve, J., and Box, J. E.: Mapping daily snow/ice shortwave broadband albedo from Moderate Resolution Imaging Spectroradiometer (MODIS): The improved direct retrieval algorithm and validation with Greenland in situ measurement, *Journal of Geophysical Research: Atmospheres*, 110, doi:10.1029/2004JD005493, 2005.
- 1620 Lindeskog, M., Arneeth, A., Bondeau, A., Waha, K., Seaquist, J., Olin, S., and Smith, B.: Implications of accounting for land use in simulations of ecosystem carbon cycling in Africa, *Earth System Dynamics*, 4, 385–407, doi:10.5194/esd-4-385-2013, 2013.
- Lloyd, J. and Taylor, J. A.: On the Temperature Dependence of Soil Respiration, *Functional Ecology*, 8, 315–323, doi:10.2307/2389824, 1994.
- Lotze-Campen, H., Müller, C., Bondeau, A., Rost, S., Popp, A., and Lucht, W.: Global food demand, productivity growth, and the scarcity of land and water resources: a spatially explicit mathematical programming approach, *Agricultural Economics*, 39, 325–338, doi:10.1111/j.1574-0862.2008.00336.x, 2008.
- 1630 Luyssaert, S., Inglima, I., Jung, M., Richardson, A. D., Reichstein, M., Papale, D., Piao, S. L., Schulze, E. D., Wingate, L., Matteucci, G., Aragao, L., Aubinet, M., Beer, C., Bernhofer, C., Black, K. G., Bonal, D., Bonafond, J. M., Chambers, J., Ciais, P., Cook, B., Davis, K. J., Dolman, A. J., Gielen, B., Goulden, M., Grace, J., Granier, A., Grelle, A., Griffis, T., Grünwald, T., Guidolotti, G., Hanson, P. J., Harding, R., Hollinger, D. Y., Hutrya, L. R., Kolari, P., Kruijt, B., Kutsch, W., Lagergren, F., Laurila, T., Law, B. E., Le Maire, G., Lindroth, A., Loustau, D., Malhi, Y., Mateus, J., Migliavacca, M., Misson, L., Montagnani, L., Moncrieff, J., Moors, E., Munger, J. W., Nikinmaa, E., Ollinger, S. V., Pita, G., Rebmann, C., Roupsard, O., Saigusa, N., Sanz, M. J., Seufert, G., Sierra, C., Smith, M. L., Tang, J., Valentini, R., Vesala, T., and Janssens, I. A.: CO₂

- balance of boreal, temperate, and tropical forests derived from a global database, *Global Change Biology*, 13, 2509–2537, doi:10.1111/j.1365-2486.2007.01439.x, 2007.
- 1640 Malik, M. J., van der Velde, R., Vekerdy, Z., and Su, Z.: Assimilation of Satellite-Observed Snow Albedo in a Land Surface Model, *Journal of Hydrometeorology*, 13, 1119–1130, doi:10.1175/JHM-D-11-0125.1, 2012.
- Monfreda, C., Ramankutty, N., and Foley, J. a.: Farming the planet: 2. Geographic distribution of crop areas, yields, physiological types, and net primary production in the year 2000, *Global Biogeochemical Cycles*, 22, 1–19, doi:10.1029/2007GB002947, 2008.
- 1645 Monsi, M.: Über den Lichtfaktor in den Pflanzengesellschaften und seine Bedeutung für die Stoffproduktion, *Japanese Journal of Botany*, 14, 22–52, doi:10.1093/aob/mci052, 1953.
- Monteith, J. and Unsworth, M.: *Principles of Environmental Physics*, Edward Arnold, New York, 2nd edn., 1990.
- 1650 Monteith, J. L.: Accommodation between transpiring vegetation and the convective boundary layer, *Journal of Hydrology*, 166, 251–263, doi:doi: DOI: 10.1016/0022-1694(94)05086-D, 1995.
- Müller, C. and Robertson, R. D.: Projecting future crop productivity for global economic modeling, *Agricultural Economics*, 45, 37–50, doi:10.1111/agec.12088, 2014.
- Müller, C., Elliott, J., Chrysanthacopoulos, J., Deryng, D., Folberth, C., Pugh, T. A., and Schmid, E.: Implications of climate mitigation for future agricultural production, *Environmental Research Letters*, 10, 125 004, doi:10.1088/1748-9326/10/12/125004, 2015.
- 1655 Müller, C., Stehfest, E., Minnen, J. G. v., Strengers, B., Bloh, W. v., Beusen, A. H. W., Schaphoff, S., Kram, T., and Lucht, W.: Drivers and patterns of land biosphere carbon balance reversal, *Environmental Research Letters*, 11, 044 002, doi:10.1088/1748-9326/11/4/044002, 2016.
- 1660 Nachtergaele, F., van Velthuisen, H., Verelst, L., Batjes, N., Dijkshoorn, K., van Engelen, V., Fischer, G., Jones, A., Montanarella, L., and Petri, M.: Harmonized world soil database, Food and Agriculture Organization of the United Nations, <http://www.fao.org/soils-portal/soil-survey/soil-maps-and-databases/harmonized-world-soil-database-v12/en/>, 2008.
- Nash, J. E.: The form of the instantaneous unit hydrograph, in: *Comptes Rendus et Rapports Assemblée Generale de Toronto*, vol. III, pp. 114–121, Toronto, <http://nora.nerc.ac.uk/508550/>, 1957.
- 1665 New, M., Hulme, M., and Jones, P.: Representing Twentieth-Century Space–Time Climate Variability. Part II: Development of 1901–96 Monthly Grids of Terrestrial Surface Climate, *Journal of Climate*, 13, 2217–2238, doi:10.1175/1520-0442(2000)013<2217:RTCSTC>2.0.CO;2, 2000.
- Olson, D. M., Dinerstein, E., Wikramanayake, E. D., Burgess, N. D., Powell, G. V. N., Underwood, E. C., D’amico, J. A., Itoua, I., Strand, H. E., Morrison, J. C., Loucks, C. J., Allnutt, T. F., Ricketts, T. H., Kura, Y., Lamoreux, J. F., Wettengel, W. W., Hedao, P., and Kassem, K. R.: Terrestrial Ecoregions of the World: A New Map of Life on Earth, *BioScience*, 51, 933, doi:10.1641/0006-3568(2001)051[0933:TEOTWA]2.0.CO;2, 2001.
- 1670 Ostberg, S., Lucht, W., Schaphoff, S., and Gerten, D.: Critical impacts of global warming on land ecosystems, *Earth System Dynamics*, 4, 347–357, doi:10.5194/esd-4-347-2013, 2013.
- Parton, W. J. and Logan, J. A.: A model for diurnal variation in soil and air temperature, *Agricultural Meteorology*, 23, 205–216, doi:10.1016/0002-1571(81)90105-9, 1981.

- Popp, A., Dietrich, J., Lotze-Campen, H., Klein, D., Bauer, N., Krause, M., Beringer, T., Gerten, D., and Edenhofer, O.: The economic potential of bioenergy for climate change mitigation with special attention given to implications for the land system, *Environmental Research Letters*, 6, 034017, doi:10.1088/1748-9326/6/3/034017, 2011.
- Popp, A., Humpenöder, F., Weindl, I., Bodirsky, B. L., Bonsch, M., Lotze-Campen, H., Müller, C., Biewald, A., Rolinski, S., Stevanovic, M., and Dietrich, J. P.: Land-use protection for climate change mitigation, *Nature Climate Change*, 4, 1095–1098, doi:10.1038/nclimate2444, 2014.
- Portmann, F. T., Siebert, S., and Döll, P.: MIRCA2000 - Global monthly irrigated and rainfed crop areas around the year 2000: A new high-resolution data set for agricultural and hydrological modeling, *Global Biogeochemical Cycles*, 24, 1–24, doi:10.1029/2008GB003435, 2010.
- Poulter, B., Aragão, L., Heyder, U., Gumpenberger, M., Heinke, J., Langerwisch, F., Rammig, A., Thonicke, K., and Cramer, W.: Net biome production of the Amazon Basin in the 21st century, *Global Change Biology*, 16, 2062–2075, doi:10.1111/j.1365-2486.2009.02064.x, 2010.
- Prentice, C. I., Sykes, M. T., and Cramer, W.: A simulation model for the transient effects of climate change on forest landscapes, *Ecological Modelling*, 65, 51–70, doi:10.1016/0304-3800(93)90126-D, 1993.
- Prentice, I., Bondeau, A., Cramer, W., Harrison, S., Hickler, T., Lucht, W., Sitch, S., Smith, B., and Sykes, M.: Dynamic Global Vegetation Modeling: Quantifying Terrestrial Ecosystem Responses to Large-Scale Environmental Change, in: *Terrestrial Ecosystems in a Changing World*, Global Change - The IGBP Series, pp. 175–192–192, Springer Berlin Heidelberg, http://dx.doi.org/10.1007/978-3-540-32730-1_15, 2007.
- Prentice, I. C., Heimann, M., and Sitch, S.: The Carbon Balance of the Terrestrial Biosphere: Ecosystem Models and Atmospheric Observations, *Ecological Applications*, 10, 1553–1573, doi:10.1890/1051-0761(2000)010[1553:TCBOTT]2.0.CO;2, 2000.
- Priestley, C. H. B. and Taylor, R. J.: On the Assessment of Surface Heat Flux and Evaporation Using Large-Scale Parameters, *Monthly Weather Review*, 100, 81–92, doi:10.1175/1520-0493(1972)100<0081:OTAOSH>2.3.CO;2, 1972.
- Pugh, T., Müller, C., Arneth, A., Haverd, V., and Smith, B.: Key knowledge and data gaps in modelling the influence of CO₂ concentration on the terrestrial carbon sink, *Plants facing Changing Climate*, 203, 3–15, doi:10.1016/j.jplph.2016.05.001, <http://www.sciencedirect.com/science/article/pii/S0176161716300529>, 2016a.
- Pugh, T., Müller, C., Elliott, J., Deryng, D., Folberth, C., Olin, S., Schmid, E., and Arneth, A.: Climate analogues suggest limited potential for intensification of production on current croplands under climate change, *Nature Communications*, 7, 12608, doi:10.1038/ncomms12608, 2016b.
- Rammig, A., Jupp, T., Thonicke, K., Tietjen, B., Heinke, J., Ostberg, S., Lucht, W., Cramer, W., and Cox, P.: Estimating the risk of Amazonian forest dieback, *New Phytologist*, 187, 694–706, doi:10.1111/j.1469-8137.2010.03318.x, 2010.
- Rammig, A., Wiedermann, M., Donges, J. F., Babst, F., von Bloh, W., Frank, D., Thonicke, K., and Mahecha, M. D.: Coincidences of climate extremes and anomalous vegetation responses: comparing tree ring patterns to simulated productivity, *Biogeosciences*, 12, 373–385, doi:10.5194/bg-12-373-2015, 2015.

- Reich, P. B., Walters, M. B., and Ellsworth, D. S.: From tropics to tundra: Global convergence in plant functioning, *Proceedings of the National Academy of Sciences*, 94, 13 730–13 734, doi:10.1073/pnas.94.25.13730, 1997.
- 1720 Rockström, J., Steffen, W., Noone, K., Persson, A., Chapin, F. S., Lambin, E. F., Lenton, T. M., Scheffer, M., Folke, C., Schellnhuber, H. J., Nykvist, B., de Wit, C. A., Hughes, T., van der Leeuw, S., Rodhe, H., Sorlin, S., Snyder, P. K., Costanza, R., Svedin, U., Falkenmark, M., Karlberg, L., Corell, R. W., Fabry, V. J., Hansen, J., Walker, B., Liverman, D., Richardson, K., Crutzen, P., and Foley, J. A.: A safe operating space for humanity, *Nature*, 461, 472–475, doi:10.1038/461472a, 2009.
- 1725 Rogers, A., Medlyn, B. E., Dukes, J. S., Bonan, G., von Caemmerer, S., Dietze, M. C., Kattge, J., Leakey, A. D. B., Mercado, L. M., Niinemets, Ü., Prentice, I. C., Serbin, S. P., Sitch, S., Way, D. A., and Zaehle, S.: A roadmap for improving the representation of photosynthesis in Earth system models, *New Phytologist*, 213, 22–42, doi:10.1111/nph.14283, 2017.
- 1730 Rolinski, S., Rammig, A., Walz, A., von Bloh, W., van Oijen, M., and Thonicke, K.: A probabilistic risk assessment for the vulnerability of the European carbon cycle to weather extremes: the ecosystem perspective, *Biogeosciences*, 12, 1813–1831, doi:10.5194/bg-12-1813-2015, 2015.
- Rosenzweig, C., Elliott, J., Deryng, D., Ruane, A. C., Müller, C., Arneth, A., Boote, K. J., Folberth, C., Glotter, M., and Khabarov, N.: Assessing agricultural risks of climate change in the 21st century in a global gridded crop model intercomparison, *Proceedings of the National Academy of Sciences*, 111, 3268–3273, doi:10.1073/pnas.1222463110, 2014.
- 1735 Rost, S., Gerten, D., Bondeau, A., Lucht, W., Rohwer, J., and Schaphoff, S.: Agricultural green and blue water consumption and its influence on the global water system, *Water Resour. Res.*, 44, W09405, doi:10.1029/2007WR006331, 2008.
- Ryan, M.: Effects of climate change on plant respiration, *Ecological Applications*, 1, 157–167, doi:10.2307/1941808, 1991.
- 1740 Schaphoff, S., Heyder, U., Ostberg, S., Gerten, D., Heinke, J., and Lucht, W.: Contribution of permafrost soils to the global carbon budget, *Environmental Research Letters*, 8, 014 026, doi:10.1088/1748-9326/8/1/014026, 2013.
- Schaphoff, S., Forkel, M., Müller, C., Knauer, J., von Bloh, W., Biemans, H., Gerten, D., Heinke, J., Jaegermyer, J., Lucht, W., Rammig, A., Thonicke, K., and Waha, K.: The LPJmL4 Dynamic Global Vegetation Model with managed Land: PART II - Evaluation of a global consistent vegetation, hydrology and agricultural model, *Geoscientific Model Development*, under Revision.
- 1745 Schewe, J., Heinke, J., Gerten, D., Haddeland, I., Arnell, N. W., Clark, D. B., Dankers, R., Eisner, S., Fekete, B. M., Colón-González, F. J., Gosling, S. N., Kim, H., Liu, X., Masaki, Y., Portmann, F. T., Satoh, Y., Stacke, T., Tang, Q., Wada, Y., Wisser, D., Albrecht, T., Frieler, K., Piontek, F., Warszawski, L., and Kabat, P.: Multi-model assessment of water scarcity under climate change, *Proceedings of the National Academy of Sciences*, 111, 3245–3250, doi:10.1073/pnas.1222460110, <http://www.pnas.org/content/111/9/3245.abstract>, 2014.
- 1750 Shinozaki, K., Yoda, K., Hozumi, K., and Kira, T.: A quantitative analysis of plant form-the pipe model theory: I. Basic analyses, *Japanese Journal of Ecology*, 14, 97–105, doi:10.18960/seitai.14.3_97, 1964.

- Siebert, S., Kummu, M., Porkka, M., Döll, P., Ramankutty, N., and Scanlon, B. R.: A global data set of the extent of irrigated land from 1900 to 2005, *Hydrology and Earth System Sciences*, 19, 1521–1545, doi:10.13019/M20599, 2015.
- Sitch, S., Smith, B., Prentice, I. C., Arneth, A., Bondeau, A., Cramer, W., Kaplan, J. O., Levis, S., Lucht, W., Sykes, M. T., Thonicke, K., and Venevsky, S.: Evaluation of ecosystem dynamics, plant geography and terrestrial carbon cycling in the LPJ dynamic global vegetation model, *Global Change Biology*, 9, 161–185, doi:10.1046/j.1365-2486.2003.00569.x, 2003.
- Sitch, S., Huntingford, C., Gedney, N., Levy, P., Lomas, M. R., Piao, S. L., Betts, R. A., Ciais, P., Cox, P. M., Friedlingstein, P., Johns, C. D., Prentice, I. C., and Woodward, F. I.: Evaluation of the terrestrial carbon cycle, future plant geography and climate-carbon cycle feedbacks using five Dynamic Global Vegetation Models (DGVMs), *Global Change Biology*, 14, 2015–2039, doi:10.1111/j.1365-2486.2008.01626.x, <http://dx.doi.org/10.1111/j.1365-2486.2008.01626.x>, 2008.
- Smith, B., Wårlind, D., Arneth, A., Hickler, T., Leadley, P., Siltberg, J., and Zaehle, S.: Implications of incorporating N cycling and N limitations on primary production in an individual-based dynamic vegetation model, *Biogeosciences*, 11, 2027–2054, doi:10.5194/bg-11-2027-2014, 2014.
- Smith, T., Shugart, H., Woodward, F., and Burton, P.: Plant functional types, in: *Vegetation Dynamics & Global Change*, pp. 272–292, Springer, <http://www.springer.com/de/book/9780412036712>, 1993.
- Sprugel, D. G., Ryan, M. G., Brooks, J. R., Vogt, K. A., and Martin, T. A.: Respiration from the organ level to the stand, *Resource physiology of conifers*, pp. 255–299, https://books.google.de/books?hl=de&lr=&id=KJl1zNzgJzYC&oi=fnd&pg=PA255&dq=Respiration+from+the+organ+level+to+the+stand&ots=IihnaKEehl&sig=UrtmXN4v0OKHK7WkE65hf_F3m3M, 1995.
- Steffen, W., Richardson, K., Rockström, J., Cornell, S. E., Fetzer, I., Bennett, E. M., Biggs, R., Carpenter, S. R., de Vries, W., de Wit, C. A., Folke, C., Gerten, D., Heinke, J., Mace, G. M., Persson, L. M., Ramanathan, V., Reyers, B., and Sörlin, S.: Planetary boundaries: Guiding human development on a changing planet, *Science*, doi:10.1126/science.1259855, 2015.
- Strengers, B. J., Müller, C., Schaeffer, M., Haarsma, R. J., Severijns, C., Gerten, D., Schaphoff, S., van den Houdt, R., and Oostenrijk, R.: Assessing 20th century climate-vegetation feedbacks of land-use change and natural vegetation dynamics in a fully coupled vegetation-climate model, *International Journal of Climatology*, 30, 2055–2065, doi:10.1002/joc.2132, 2010.
- Strugnell, N. C., Lucht, W., and Schaaf, C.: A global albedo data set derived from AVHRR data for use in climate simulations, *Geophysical Research Letters*, 28, 191–194, doi:10.1029/2000GL011580, 2001.
- Thonicke, K., Venevsky, S., Sitch, S., and Cramer, W.: The role of fire disturbance for global vegetation dynamics: coupling fire into a Dynamic Global Vegetation Model, *Global Ecology and Biogeography*, 10, 661–677, doi:10.1046/j.1466-822X.2001.00175.x, 2001.
- Thonicke, K., Spessa, A., Prentice, I. C., Harrison, S. P., Dong, L., and Carmona-Moreno, C.: The influence of vegetation, fire spread and fire behaviour on biomass burning and trace gas emissions: results from a process-based model, *Biogeosciences*, 7, 1991–2011, doi:10.5194/bg-7-1991-2010, <http://www.biogeosciences.net/7/1991/2010/>, 2010.
- Thornley, J. H. M.: Respiration, Growth and Maintenance in Plants, *Nature*, 227, 304, <http://dx.doi.org/10.1038/227304b0>, 1970.

Thornthwaite, C. W.: An Approach toward a Rational Classification of Climate, *Geographical review*, 38, 55–94, doi:10.2307/210739, 1948.

Tjoelker, M., Oleksyn, J., Reich, P. B., and others: Acclimation of respiration to temperature and CO₂ in seedlings of boreal tree species in relation to plant size and relative growth rate, *Global Change Biology*, 5, 679–691, doi:10.1046/j.1365-2486.1999.00257.x, 1999.

University of East Anglia Climatic Research Unit; Harris, I.C.; Jones, P. : CRU TS3.23: Climatic Research Unit (CRU) Time-Series (TS) Version 3.23 of High Resolution Gridded Data of Month-by-month Variation in Climate (Jan. 1901- Dec. 2014)., Centre for Environmental Data Analysis, <http://dx.doi.org/10.5285/4c7fdfa6-f176-4c58-acee-683d5e9d2ed5>, 2015.

Von Bloh, W., Rost, S., Gerten, D., and Lucht, W.: Efficient parallelization of a dynamic global vegetation model with river routing, *Environmental Modelling & Software*, 25, 685–690, doi:10.1016/j.envsoft.2009.11.012, 2010.

Vorosmarty, C. and Fekete, B.: ISLSCP II River Routing Data (STN-30p), in: ISLSCP Initiative II Collection. Data set., edited by Hall, F. G., Collatz, G., Meeson, B., Los, S., Brown de Colstoun, E., and Landis, D., ORNL Distributed Active Archive Center, <https://doi.org/10.3334/ORNLDAAAC/1005>, 2011.

Vörösmarty, C. J., McIntyre, P. B., Gessner, M. O., Dudgeon, D., Prusevich, A., Green, P., Glidden, S., Bunn, S. E., Sullivan, C. A., Liermann, C. R., and Davies, P. M.: Global threats to human water security and river biodiversity, *Nature*, 467, 555, <http://dx.doi.org/10.1038/nature09440>, 2010.

Waha, K., van Bussel, L. G. J., Müller, C., and Bondeau, A.: Climate-driven simulation of global crop sowing dates, *Global Ecology and Biogeography*, 21, 247–259, doi:10.1111/j.1466-8238.2011.00678.x, 2012.

Waha, K., Müller, C., Bondeau, a., Dietrich, J., Kurukulasuriya, P., Heinke, J., and Lotze-Campen, H.: Adaptation to climate change through the choice of cropping system and sowing date in sub-Saharan Africa, *Global Environmental Change*, 23, 130–143, doi:10.1016/j.gloenvcha.2012.11.001, 2013.

Waring, R., Schroeder, P., and Oren, R.: Application of the pipe model theory to predict canopy leaf area, *Canadian Journal of Forest Research*, 12, 556–560, doi:10.1139/x82-086, 1982.

Waring, R. H.: Estimating Forest Growth and Efficiency in Relation to Canopy Leaf Area, *Advances in Ecological Research*, 13, 327–354, doi:10.1016/S0065-2504(08)60111-7, 1983.

Xu, R., Dai, J., Luo, W., Yin, X., Li, Y., Tai, X., Han, L., Chen, Y., Lin, L., and Li, G.: A photothermal model of leaf area index for greenhouse crops, *Agricultural and Forest Meteorology*, 150, 541–552, doi:10.1016/j.agrformet.2010.01.019, 2010.

Zachle, S., Sitch, S., Prentice, I. C., Liski, J., Cramer, W., Erhard, M., Hickler, T., and Smith, B.: The importance of age-related decline in forest NPP for modeling regional carbon balances, *Ecological Applications*, 16, 1555–1574, doi:10.1890/1051-0761(2006)016[1555:TIOADI]2.0.CO;2, 2010.

Zeide, B.: Primary Unit of the Tree Crown, *Ecology*, 74, 1598–1602, doi:10.2307/1940088, 1993.

Zhou, X., Zhu, Q., Tang, S., Chen, X., and Wu, M.: Interception of PAR and relationship between FPAR and LAI in summer maize canopy, in: *Geoscience and Remote Sensing Symposium, 2002. IGARSS'02. 2002 IEEE International*, vol. 6, pp. 3252–3254, IEEE, doi:10.1109/IGARSS.2002.1027146, 2002.

Zscheischler, J., Mahecha, M., Von Buttlar, J., Harmeling, S., Jung, M., Rammig, A., Randerson, J. T., Schölkopf, B., Seneviratne, S. I., Tomelleri, E., Zachle, S., and Reichstein, M.: Few extreme events dominate

global interannual variability in gross primary production, *Environmental Research Letters*, 9, 035 001, doi:10.1088/1748-9326/9/3/035001, 2014a.

- 1835 Zscheischler, J., Reichstein, M., Harmeling, S., Rammig, A., Tomelleri, E., and Mahecha, M.: Extreme events in gross primary production: a characterization across continents, *Biogeosciences*, 11, 2909–2924, doi:10.5194/bg-11-2909-2014, 2014b.

Table 1. Abbreviation of PFTs, BFTs and CFTs.

Tropical broadleaved evergreen tree	TrBE
Tropical broadleaved raingreen tree	TrBR
Temperate needleleaved evergreen tree	TeNE
Temperate broadleaved evergreen tree	TeBE
Temperate broadleaved summergreen tree	TeBS
Boreal needleleaved evergreen tree	BoNE
Boreal broadleaved summergreen tree	BoBS
Boreal needleleaved summergreen tree	BoNS
Tropical herbaceous	TrH
Temperate herbaceous	TeH
Polar herbaceous	PoH
<hr/>	
Bioenergy tropical tree	BTrT
Bioenergy temperate tree	BTeT
Bioenergy C4 grass	BGrC4
<hr/>	
Temperate cereals	TeCer
Rice	Rice
Maize	Maize
Tropical cereals	TrCer
Pulses	Pul
Temperate roots	TeRo
Tropical roots	TrRo
Sunflower	SunFl
Soy-bean	Soy
Groundnut	GrNu
Rapeseed	Rape
Sugar-cane	SuCa