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Plant profit maximisation improves predictions of European forest responses to drought

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Summary

- Knowledge of how water stress impacts the carbon and water cycles is a key uncertainty in terrestrial biosphere models.
- We tested a new profit maximisation model, where photosynthetic uptake of CO₂ is optimally traded against plant hydraulic function, as an alternative to the empirical functions commonly used in models to regulate gas exchange during periods of water stress. We conducted a multi-

site evaluation of this model at the ecosystem scale, before and during major droughts in Europe. Additionally, we asked whether the maximum hydraulic conductance in the soil-plant continuum (k_{max}) – a key model parameter which is not commonly measured – could be predicted from long-term site climate.

- Compared to a control model with an empirical soil moisture function, the profit maximisation model improved the simulation of evapotranspiration during the growing season, reducing the Normalised Mean Square Error by c. 63%, across mesic and xeric sites. We also showed that k_{max} could be estimated from long-term climate, with improvements in the simulation of evapotranspiration at eight out of the 10 forest sites during drought.
- Although the generalisation of this approach is contingent upon determining k_{max} , it presents a mechanistic trait-based alternative to regulate canopy gas exchange in global models.

Key words: vegetation drought responses, land surface models, canopy gas exchange, plant optimality, plant profit maximisation, plant trait coordination, hydraulic trait adjustments to climate.

I. Introduction

Water availability is one of the primary controls of vegetation function, largely responsible for the pronounced seasonal and interannual variability in global CO₂ uptake by terrestrial ecosystems (Ahlstrom *et al.*, 2015; Jung *et al.*, 2017; Humphrey *et al.*, 2018; Green *et al.*, 2019). As a result, contemporary droughts affecting the Amazon, Australia, Europe and North America have had considerable impact on ecosystems (e.g. affecting water use efficiency, see Peters *et al.* (2018)), human populations and economic systems (Ciais *et al.*, 2005; Marengo *et al.*, 2013; van Dijk *et al.*, 2013; Boyer *et al.*, 2013). Critically, the impact of drought is often long-lasting (Kannenberg *et al.*, 2019). For example, the legacy of the 2003 European drought was apparent beyond the conclusion of the drought, as forests experienced diebacks and pest infestations (Bréda *et al.*, 2006; Stahl *et al.*, 2016). Increases in drought intensity and/or duration, as projected in response to climate change, would therefore have major implications for the terrestrial biosphere (Stocker *et al.*, 2014; Ault *et al.*, 2014; Cook *et al.*, 2015; Zhao & Dai, 2017).

In coupled climate models, land surface models (LSMs) simulate the exchange of carbon, water and energy fluxes between the land surface and the atmosphere (Pitman, 2003). Land schemes typically use empirical relationships, hereafter referred to as β functions, to reduce canopy gas exchange in response to a decline in soil water availability (Best *et al.*, 2011; Egea *et al.*, 2011). These β functions rely on soil-dependent parameters, e.g. soil moisture content at field capacity and the wilting point, instead of reflecting vegetation adaptations to water availability. β has also been defined based on soil water potential thresholds (Oleson *et al.*, 2013), causing abrupt declines in vegetation function due to declining water availability in the top soil layers (De Kauwe *et al.*, 2015b; Medlyn *et al.*, 2016). Overall, β functions lack both empirical support (Verhoef & Egea, 2014; Medlyn *et al.*, 2016) and theoretical foundation, leading to important inter-model disagreement in the shapes of the functional forms used to limit gas exchange (Desborough, 1997; Medlyn *et al.*, 2016; De Kauwe *et al.*, 2017).

Importantly, the use of β functions contradicts widespread evidence that plants differ in their sensitivity to reduced water availability (Brodribb & Cochard, 2009; Urli *et al.*, 2013; Li *et al.*, 2017). Explicit representations of plant hydraulics (segmented into e.g. root, stem, and leaf hydraulic elements) offer a mechanistic alternative to empirical soil-dependent β functions, and can be parameterised from measured plant traits (e.g. Sperry *et al.*, 1998; Williams *et al.*, 2001; Sperry & Love, 2015; Xu *et al.*, 2016; Tuzet *et al.*, 2017; Martin-StPaul *et al.*, 2017). By depicting water transport through the soil-plant-atmosphere continuum, plant hydraulic schemes open pathways to capture key vegetation responses to drought: as water availability declines, xylem water potential drops and critical cavitation might occur (Wolfe *et al.*, 2016; Martin-StPaul *et al.*, 2017), a process which could lead to representing mortality via hydraulic failure.

Another approach extends the long-standing Water Use Efficiency Hypothesis (WUEH; Cowan & Farquhar (1977); Medlyn *et al.*, (2011)) in relation to plant hydraulics (Wolf *et al.*, 2016; Sperry *et al.*, 2017). The WUEH postulates that stomata adjust to keep the relationship between the sensitivities of the stomatal conductance (g_s) to transpiration (E) and to net carbon assimilation (A_n) constant for a fixed amount of water loss (λ). Whilst the WUEH is supported by empirical data (Lin *et al.*, 2015), there is no clear way to estimate changes in λ on the timescale over which soil water changes (Wong *et al.*, 1985; Manzoni *et al.*, 2011). Consequently, the WUEH fails to relate the sensitivities of E and

 A_n with a hydraulic cost of water loss and still relies on a β function to limit gas exchange (De Kauwe et al., 2015a).

By contrast, Wolf *et al.* (2016) and Sperry *et al.* (2017) hypothesised that plants optimally control water loss on an instantaneous basis, to avoid excessive reduction in xylem hydraulic conductivity from a drying soil. In their approach, vulnerability curves are modelled from several set points, which correspond to specific plant hydraulic parameters, i.e. two P_x parameters representing the water potential drop at x% xylem conductivity loss (e.g. P_{50} at 50% xylem conductivity loss). It is worth highlighting that the set points used to model the vulnerability curves arise directly from measurements. The later allows for realistic simulations of the progressive impairment of water flow through the xylem, as water potential drops from the point of maximum hydraulic conductance. Unlike the WUEH, these approaches offer a mechanism to parameterise a spectrum of water use strategies, accounting for plant vulnerability to water stress.

In practice, Wolf *et al.* (2016) reformulated the WUEH, by expressing the cost of transpiration as a marginal carbon cost due to lost hydraulic conductivity, but direct measurements and/or a specific formulation of this marginal carbon cost are lacking. Sperry *et al.* (2017) overcame this limitation by normalising the water and carbon components of the WUEH, which avoids an explicit marginal cost. Their solution has since been shown to perform well at the plant scale in garden experiments (Venturas *et al.*, 2018; Wang *et al.*, 2019), but is yet to be tested at the ecosystem scale and across ecosystems.

Applying the Wolf *et al.* (2016) and Sperry *et al.* (2017) schemes in models requires three specific hydraulic parameters. Whereas values of the two required P_x parameters are readily available from hydraulic trait databases (e.g. Choat *et al.*, 2012), estimating the maximum hydraulic conductance in the soil-plant continuum (k_{max}) is more complicated. Wolf *et al.* (2016) proposed combining the conductivity from soil to the fine-root xylem, the hydraulic conductivity in the root zone, and the ratio of root to leaf area to calculate k_{max} . Venturas *et al.* (2018) and Wang *et al.* (2019) set the rhizosphere maximum hydraulic conductance and inferred the whole plant (roots to leaves) component of k_{max} from measured ratios of transpiration to leaf water potential in well-watered conditions. Either

formulation is challenging to parameterise globally and requires additional input parameters that are not readily available.

This study has two objectives:

to take the Sperry *et al.* (2017) model from the plant to the ecosystem scale, testing the capacity of this new model to simulate observed (eddy covariance) water and carbon fluxes across European forests during two major droughts (2003 and 2006);

to test whether the unknown key model parameter, k_{max} in the soil-plant continuum, can be estimated at the stand (or ecosystem) scale, assuming coordination between the hydraulic and photosynthetic traits, as well as hydraulic long-term behavioural plasticity and adjustments to climate.

To address objective I, we implemented a modified version of the Sperry *et al.* (2017) model into a simplified LSM. This allows us to test the behaviour of this new model in a generic tractable framework, respecting the broad assumptions commonly used in LSMs without making the implementation specific to an existing LSM. Inside the tractable LSM, we compared the behaviour of the modified Sperry *et al.* (2017) model to that of the Medlyn stomatal optimisation model (Medlyn *et al.*, 2011), which is widely used in state-of-the-art LSMs (CABLE (De Kauwe *et al.*, 2015a), CLM5.0 (Kennedy *et al.*, 2019), JULESv5.4 (Oliver *et al.*, 2018)). To address objective II, we explored alternative ways to determine k_{max} , testing combinations of various climatic and behavioural adjustments, but always assuming hydraulic and photosynthetic trait coordination.

II. Materials and Methods

The Materials and Methods are organised into six sections: (1) describes the modified Sperry *et al.* (2017) model; (2) presents the approaches tested to estimate k_{max} ; (3) outlines the core components of the tractable LSM; (4) describes the forcing and site data; (5) details the model experiments performed; and (6) introduces a methodology to robustly assess model performance.

A summary of the model experiments and configurations (Material and Methods 3 and 5) is available in Table 3. A more visual summary of our experimental setup (Materials and Methods 5 to 6a) is

presented in the Supporting Information Fig. S1. For the site information and parameters, cf. Tables 1 and 2 and the Supporting Information Tables S1 and S2; Tables S3-S5 contain additional model parameters.

1. Profit Maximisation Approach

The approach developed by Sperry *et al.* (2017) proposes that plants regulate leaf water potential (Ψ_{leaf} ; MPa) on an instantaneous basis, by trading increasingly marginal carbon intakes against increasing hydraulic conductance losses. Thus, plants balance carbon gain (CG) and hydraulic cost (HC) at the optimal leaf water potential ($\Psi_{leaf,opt}$; MPa) where profit is maximised:

$$Profit_{max} = max(CG(\Psi) - HC(\Psi)) \qquad \in [0,1]$$
 (1)

 Ψ (MPa) is the water potential in the soil-plant continuum (see paragraph below). CG and HC are unitless and normalised to one, which makes them comparable.

In our implementation of the model introduced in Eqn 1, Ψ varies along a continuous transpiration stream (i.e. a single hydraulic conductor): from the root zone soil water potential (Ψ_s ; MPa) to Ψ_{leaf} . Ψ_{leaf} cannot drop below the critical leaf water potential indicative of maximum xylem hydraulic failure (Ψ_{crit} ; MPa). Using a continuous transpiration stream reduces parameterisation by removing hydraulic segmentation, but likely results in more marked drought stress because it smooths conductance changes between the root zone and the leaves. We set Ψ_{crit} to match a near complete hydraulic conductivity loss of 95%, accounting for possible high levels of embolism resistance.

a) Hydraulic Cost

The normalised hydraulic cost function reflects the increasing potential damage from cavitation as hydraulic conductance is lost:

$$HC(\Psi) = \frac{k_{i,max} - k(\Psi)}{k_{i,max} - k_{crit}} \in [0,1]$$
 (2)

where $k_{i,max}$ (mmol m⁻² s⁻¹ MPa⁻¹) is the instantaneous maximum hydraulic conductance in the soilplant continuum, after accounting for water stress (i.e. k_{max} evaluated at Ψ_s), k_{crit} (mmol m⁻² s⁻¹ MPa⁻¹) is the critically low hydraulic conductance that provokes xylem failure at Ψ_{crit} , and k (mmol m⁻² s⁻¹ MPa⁻¹) is the hydraulic conductance for any water potential in the soil-plant continuum (see Eqn 3).

Hydraulic conductance is represented by a cumulative Weibull distribution (Neufeld et al., 1992):

$$k(\Psi) = k_{max} e^{-\left(\frac{\Psi}{b}\right)^c} \tag{3}$$

where k_{max} (mmol m⁻² s⁻¹ MPa⁻¹) is the maximum hydraulic conductance in the soil-plant continuum; b (MPa) and c (unitless) are sensitivity and shape parameters of the plant hydraulic vulnerability curve, reconstructed from two P_x parameters (e.g. P_{50} and P_{88} ; cf. Eqn S8 and S9).

Here, k_{max} represents the maximum conductance of a single hydraulic conductor connecting the root zone to the leaves. In segmented representations of plant hydraulic, this parameterisation would vary between the rhizosphere, roots, stem, etc. Our approach combines all the hydraulic elements in the soil-plant continuum, hence encompassing maximum rhizosphere conductance as well as maximum whole plant hydraulic conductance.

k drops with soil water depletion, owing to xylem embolism, and recovers with soil water recharge (i.e. we assume perfect and instantaneously reversible embolism). Hence, the supply of water for transpiration from the root zone depends on plant hydraulic vulnerability and photosynthetic demand (via CG in Eqn 1), for any steady-state pressure drop between Ψ_s and Ψ_{leaf} (Sperry & Love, 2015):

$$E(\Psi_{leaf}) = \int_{\Psi_s}^{\Psi_{leaf}} k(\Psi) d\Psi$$
 (4)

where E is expressed in mmol m⁻² s⁻¹.

b) Carbon Gain

The normalised carbon gain function represents a marginally decreasing carbon gain as leaf water potential becomes more negative:

$$CG(\Psi_{leaf}) = \frac{A(\Psi_{leaf})}{A_{max}} \in [0,1]$$
 (5)

where A (µmol m⁻² s⁻¹) is the photosynthetic uptake at each corresponding Ψ_{leaf} and A_{max} (µmol m⁻² s⁻¹) is the instantaneous maximum over the range of Ψ_{leaf} .

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The diffusive supply of CO_2 uses Fick's first law to represent the CO_2 flux from the ambient air (C_a) into the leaf intercellular air spaces (C_i):

$$A(\Psi_{leaf}) = g_c(\Psi_{leaf}) \left(C_a - C_i(\Psi_{leaf}) \right) \tag{6}$$

where g_c is the CO₂ diffusive leaf conductance (mol m⁻² s⁻¹). C_i is obtained at Ψ_{leaf} by solving a system comprising Eqn 6 and a biochemical photosynthesis model (Farquhar *et al.*, 1980) of net carbon assimilation (A_n ; µmol m⁻² s⁻¹); see Supporting Information Methods S1.

In Eqn 6, g_c varies with Ψ_{leaf} . To solve for g_c , leaf-to-air vapour pressure deficit (D_{leaf} ; kPa), and leaf temperature (T_{leaf} ; °C) – at a given Ψ_{leaf} , we build a system of three expressions of $E(\Psi_{leaf})$: (i) E given by the supply function (Eqn 4); (ii) E meeting the atmospheric demand for water, i.e. equating $1.57g_cD_{leaf}$ where 1.57 converts from conductance to CO_2 to total leaf conductance to water vapour; and (iii) E given by the Penman-Monteith equation following radiative and thermodynamic constraints imposed on T_{leaf} (see Eqn S7), and thus on D_{leaf} .

Whilst the second equation of this system (atmospheric demand for *E*) implies a perfect coupling between the leaves and the atmosphere, stomatal feedbacks are theoretically accounted for via the plant hydraulic vulnerability in the supply equation (cf. Eqn 4).

2. Calculating the Maximum Hydraulic Conductance (k_{max})

As discussed in the Introduction, the key model parameter k_{max} is not readily available. Katul *et al.* (2003) and Sperry *et al.* (2017) suggested a non-instantaneous (but not clearly defined) timescale of optimality, on which hydraulic plant traits would coordinate with photosynthetic traits. This hypothesis would reduce parameterisation and be supported by experimental evidence at the leaf-(e.g. Brodribb & Feild, 2000) and the stand-level (e.g. Lai *et al.*, 2002).

a) k_{max} behavioural plasticity

Sperry *et al.* (2017) suggested that to maintain an optimal C_i : C_a ratio of 0.7, k_{max} should be coordinated with respect to both Eqn 1 and the maximum carboxylation rate at 25°C ($V_{cmax,25}$) (cf. fig. 3b in Sperry *et al.* (2017)). With that assumption, plants always follow the profit maximisation

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approach, even at longer timescales; the possibility of long-term strategies oriented toward greater carbon accumulation, or greater water conservation, is ignored.

Here, we extend the Sperry *et al.* (2017) coordination hypothesis to add two alternative solutions for k_{max} , because we have no *a priori* knowledge of how, and whether, hydraulic and photosynthetic trait coordination varies across plant species. We test the idea that a degree of plasticity in k_{max} would feed back on $\Psi_{leaf,opt}$, and by extension on the water and carbon fluxes, so as to allow for behavioural plasticity in k_{max} at longer timescales.

All our solutions for k_{max} are calculated assuming well-watered conditions. For the first solution, $k_{max,opt}$, we calculate the value of k_{max} that yields C_i : $C_a = 0.7$ at $\Psi_{leaf,opt}$, following Eqn 1, as Sperry et al. (2017) did. $\Psi_{leaf,opt}$ corresponds to levels of embolism which vary depending on the vulnerability curve, $V_{cmax,25}$, and soil water potential. For the second solution, $k_{max,high}$, we calculate the value of k_{max} that yields C_i : $C_a = 0.7$ immediately before Ψ_{leaf} equals P_{12} . P_{12} is the water potential at a 12% loss in xylem conductivity, which coincides with the onset of xylem embolism (Choat et al., 2018). $k_{max,high}$ is independent of Eqn 1 but makes use of Eqns 3-4 and 6 and again, $k_{max,high}$ is coordinated with $V_{cmax,25}$ so as to achieve C_i : $C_a = 0.7$. For the third solution, $k_{max,low}$, we calculate the value of k_{max} that yields C_i : $C_a = 0.7$ immediately before hydraulic cost offsets net profit. Mathematically, hydraulic cost exceeds profit when $CG(\Psi) = 2HC(\Psi)$. This third solution is the one for which the percentage loss in conductivity at Ψ_{leaf} ought to be the greatest (except in cases where the parameterised vulnerability curves indicate very negative hydraulic safety margins). Eqns 2-6 are used to determine $k_{max,low}$.

 $k_{max,high}$ and $k_{max,low}$ are intended as indicators of possible alternative longer-term optimisation strategies, since coordination between them and C_i : $C_a = 0.7$ may not always be physiologically meaningful. C_i : $C_a = 0.7$ might never be observed near the onset of xylem embolism for drought avoiding species, but it is unclear what a more appropriate value would be. Note, assuming these alternative set-points is analogous to assuming different soil water availabilities upon the calculation of $k_{max,opt}$.

In all three cases, once k_{max} has been calculated, it is used as an input parameter for the profit maximisation algorithm (cf. Materials and Methods 1), which simulates actual model outputs on an instantaneous basis. Figure 1 illustrates how using values of $k_{max,high}$ and $k_{max,low}$ might alter the instantaneous profit maximisation compared to $k_{max,opt}$. In Figure 1, $k_{max,high}$ maximises profit at smaller HC and CG compared to the two other solutions, displaying a more conservative water use strategy. On the contrary, $k_{max,low}$ achieves maximum profit at higher HC and CG compared to the two other solutions, displaying a more profligate water use.

Figure 1 demonstrates an example of the relative behaviours of $k_{max,opt}$, $k_{max,high}$ and $k_{max,low}$. These relative behaviours are determined by the plant's vulnerability curve and, as such, vary from species to species.

b) k_{max} adjustments to climate

Another uncertainty pertains to the coordination of k_{max} with photosynthetic traits and climate. Figure 2 displays the theoretical response of $k_{max,opt}$ (from our first k_{max} solution) to changes in atmospheric conditions. In Figure 2a, the relationship between increasing $V_{cmax,25}$ and increasing $k_{max,opt}$ is shown. As vapour pressure deficit (*D*) increases, $k_{max,opt}$ further increases for a given $V_{cmax,25}$, albeit logarithmically. In Figure 2b, $V_{cmax,25}$ is fixed and $k_{max,opt}$ is constrained by T_{air} ; it declines above 25°C.

Figure 2 shows that assuming $T_{air} = 25$ °C and D = 1 kPa, as Sperry *et al.* (2017) did, might not be appropriate for the calculation of k_{max} . As such, we test two scenarios to estimate whether the previously introduced k_{max} behavioural plasticity displays further adjustment to: (i) average climatic conditions; or (ii) to hotter and drier conditions (see Materials and Methods 4. The first scenario represents the background conditions under which plants grow. The second scenario will help us gauge the degree to which they might make hydraulic adjustments to cope with extreme growing conditions.

3. Modelling Framework

We hereafter refer to the default model, embedding the Medlyn *et al.* (2011) stomatal model, as the 'Control model', and to the profit maximisation approach (see Materials and Methods 1) as the 'Profit_{max} model'. Both models were implemented within the same tractable LSM and forced by half-hourly meteorological inputs (i.e. photosynthetically active radiation, air temperature, precipitation, vapour pressure deficit, atmospheric pressure, and wind speed), which makes results directly comparable. Our LSM represents many of the core processes used across LSMs, albeit simplified. Notes S1 and Figures S2-S5 of the Supporting Information show a broad agreement between the Control model and the CABLE LSM (version 2.0, revision 5320; (Kowalczyk *et al.*, 2006; Wang *et al.*, 2011)). We now summarise the key features of the tractable LSM but see the Supporting Information Methods S1-S6 for more information.

We used a two-big-leaf approximation (Wang & Leuning, 1998), differentiating the radiation absorbed by sunlit and shaded leaves to simulate canopy fluxes. Soil hydrology was represented using a water balance 'tipping bucket' model.

In the Control model, water stress impacts canopy fluxes by down-regulating the slope of the sensitivity of A_n to g_s , depending on the empirical soil-moisture stress factor β :

$$\beta = \frac{\theta - \theta_{wp}}{\theta_{fc} - \theta_{wp}} \tag{8}$$

where θ (m³ m⁻³) is volumetric soil moisture and θ_{wp} (m³ m⁻³) and θ_{fc} (m³ m⁻³) are the volumetric soil moisture contents at wilting point and field capacity, respectively.

The Profit_{max} model transforms θ to Ψ_s via the Clapp and Hornberger water retention equation (Clapp & Hornberger, 1978; see Eqn S11), for use in the calculations related to the transpiration stream.

4. Forcing and site data

We tested the modelling framework at 10 European eddy-covariance sites on a latitudinal gradient (cf. Table 1) over the years preceding and during two severe droughts: the 2003 continental and 2006

regional heat wave and drought events in Europe. Attention was given to diversity in climate (cf. Table 2) and forest type (cf. Table 1), to encompass a broad range of tree functional traits (cf. Table S2). Half-hourly meteorological forcing data, as well as latent heat (LE) and Gross Primary Productivity (GPP) data used to evaluate the framework, originate from the MDS gap-filled FLUXNET2015 and LaThuile datasets (http://fluxnet.fluxdata.org) and were pre-processed using the FluxnetLSM R package (v1.0; Ukkola *et al.*, 2017). The GPP data in the FLUXNET2015 and LaThuile datasets are not actually measured, but rather estimated from measured Net Ecosystem Exchange data using a night-time partitioning approach (Reichstein *et al.*, 2005).

The model was run using site-specific prescribed phenologies (see Methods S7a) derived from the 8-day MODIS MOD15A2H Version 6 Leaf Area Index (LAI) product (Myneni *et al.*, 2015). The total effective root zone depth of the 'tipping bucket' model was set between 0.3–1.20 m across sites (cf. Table S1 for the site-specific values and Methods S8a for how they were obtained). Species-specific trait data, as well as broader Plant Functional Type (PFT), parameters used to run the LSM are provided in the Supporting Information Tables S2-S5.

5. Model spin-up and experiments

Drought events were modelled by running the Control and Profit_{max} models for the calendar years of drought occurrence (2003 and 2006). The years before drought (2002 and 2005) were also modelled to act as reference conditions. The Control model was spun up to initialise the root zone soil moisture state available to both the Control and the Profit_{max} at the beginning of each year. This spin-up was either forced with the flux tower's meteorological record for the previous year, when available, or with the average site climatology.

a) k_{max} calibration

Objective I (see Introduction) was addressed by calibrating k_{max} to assess whether the Profit_{max} model can better explain the observed LE and GPP than the Control model (cf. Notes S2 and Fig. S6 for a calibration of the Control). To calibrate k_{max} , we first calculated $k_{max,opt}$ at $T_{air} = 25$ °C and D = 1 kPa. This reference $k_{max,opt}$ was then used to generate a sequence of possible values for k_{max} . 12

values were evenly selected between 25% - 95% of $k_{max,opt}$ and a further 12 values were evenly distributed between 105% - 250% of $k_{max,opt}$. The reference $k_{max,opt}$ itself was added to the sequence, which in total led to 25 values being used as parameter inputs to run each of the drought and non-drought years. The best calibrated k_{max} value was selected following the procedure detailed below, in Section 6a of the Materials and Methods.

b) k_{max} adjustments to climate

Objective II (see Introduction) was addressed by calculating the site-level k_{max} following the strategies in Table 3: the $k_{max,opt}$, $k_{max,high}$, and $k_{max,low}$ (introduced in Materials and Methods 2a) were each adjusted depending on long-term climate. We used coarser resolution meteorological data, from the 4.03 release of the Climate Research Unit TS dataset (CRU TS v4.03; Harris *et al.*, 2014) for the 1972-2002 period, to define two contrasting climate scenarios based on T_{air} and D:

- (i) an 'Average' scenario, where CRU monthly daily mean air temperature was averaged over the growing season ($T_{air,avg}$; °C), i.e. April November between 1972-2002. Average monthly vapour pressure deficit, D_{avg} (kPa), was calculated via Tetens' equation for saturation vapor pressure (Monteith & Unsworth, 1990), using air temperature and monthly average actual vapour pressure over the growing season;
- (ii) an 'Extreme' scenario, where the 90th percentile of CRU monthly average daily maximum temperature was used to calculate $T_{air,xx}$. The associated vapour pressure deficit (D_{xx} ; kPa) was calculated as above.

All the $k_{max,opt}$, $k_{max,high}$, and $k_{max,low}$ were calculated for the soil water potential at saturation, given that the plants' vulnerability curves already account for sensitivity to water stress. The site average growing season weighted sunlit-shaded LAI (cf. Table 2), but not the rooting depth, was used to scale the estimates from the leaf up to the ecosystem, assuming that a composite shaded-sunlit leaf proxy is representative of the stand's most productive state (cf. Methods S7b).

Table 2 summarises the sites' climate scenario information and the composite LAIs, whereas Table 3 provides an overview of the model experiments and configurations.

6. Model Performance Assessment

We applied the benchmarking methodology from the Protocol for the Analysis of Land Surface Models (PALS) Land Surface Model Benchmarking Evaluation Project (PLUMBER; Best *et al.*, 2015) to assess model performance. This methodology combines several statistical metrics to produce a comprehensive assessment of model skill, overcoming the shortcomings of individual metrics.

Five common statistical measures were used: Normalised Mean Square Error (NMSE), Mean Absolute Error (MAE), Standard Deviation (SD), and the 5th and 95th percentile values (p_5 and p_{95}) indicative of agreement between the simulated and observed distributions. Following PLUMBER (Best *et al.*, 2015), we calculated the absolute difference between 1 and the ratio of modelled to observed SD, and the absolute distances between the modelled and observed p_5 and p_{95} .

For each statistical metric, we assigned quantile ranks between 0 (best performance) and 1 (worst performance) to the model configurations within an experiment (cf. Materials and Methods 5a and 5b), i.e. we ranked the configurations relative to what the best performance could be following a quantile distribution. We then averaged each configuration's quantile ranks across the metrics. We therefore obtained an average quantile rank for each configuration, within each experiment. Notably, unlike averaging absolute ranks, averaging quantile ranks considers the configurations' relative performance across the statistical metrics.

a) Selection of best performing configurations within each of the experiments

To select: (i) the best calibrated configuration (objective I), and (ii) the best climate configuration (Average or Extreme scenario used to derive k_{max} ; objective II), we constrained the half-hourly data between April – July in 2002 and 2003. In doing so, we limited the effects of the dry downs in selecting the configurations, i.e. we did not calibrate k_{max} on the drought months in 2003 (July – October). We opted to select the best performing configurations against the observed evapotranspiration (ET) only, given that GPP is not directly observed but, rather, estimated at eddy-

covariance sites.

Although we did not train the Profit_{max} model's algorithm to match the observed data, selecting the best performing configurations based on a subset of data (ET restricted between April – July in 2002 and 2003) has consequences. We expect the ET predictions to be reasonably close to the observations between April – July in 2002 and 2003, because they are in-sample. At any other time (i.e. between July – November in 2002 and 2003; and 2005 and 2006), the Profit_{max} model's ET predictions will be out-of-sample. The model's ability to match the observations will then depend on: (i) its skill, e.g. in terms of partitioning ET into transpiration and soil evaporation; (ii) changes in the vegetation properties, including potential legacy effects (in 2005 and 2006) of the 2003 drought on k_{max} . By contrast with the ET predictions, the GPP predictions are always made out-of-sample.

b) Evaluating the experiments' best performing configurations and the Control model

In Section 1 of the Results, we gauge the effects of selecting each experiment's best performing configuration (i.e. best calibration and best climate configuration) against ET only. To conduct this analysis, we first evaluated the Profit_{max} model's ability to simulate both ET (partially out-of-sample) and GPP (effectively out-of-sample) over the full April – November periods. Then, again, we combined the statistical metrics into quantile ranks for each of the best configurations and the Control model, relative to the observations.

7. Code

All model and analysis code are freely available from https://github.com/ManonSabot/Profit_Maximisation_European_Forests (doi: 10.5281/zenodo.3566722)

III. Results

1. Model evaluation

Figure 3 shows each site's average quantile ranks, attained by the best model configurations and by the Control model, for GPP (panels a, c) and ET (panels b, d); panels a, b show drought years and

panels c, d show non-drought years. Lower ranks equate to better overall performance and box plots on the right of the figure summarise results across the entire period. Overall for ET, the calibration (light green, average quantile rank $\mu=0.44$) and the best $k_{max,opt}$ among the Average and Extreme scenarios (best climate configuration summarised in dark green, $\mu=0.55$) outperformed the Control model (purple, $\mu=0.74$). For GPP, the performances of the best calibration ($\mu=0.56$), of the best climate configuration ($\mu=0.59$), and of the Control model ($\mu=0.61$) were similar. Selecting for one model output – as we did by selecting the best configurations by evaluating their skill in reproducing ET observations alone – is likely to degrade performance of other outputs (Abramowitz *et al.*, 2019; Herger *et al.*, 2019), but in the case of the Profit_{max} model, it did not lead to significant losses in overall GPP performance.

The best Profit_{max} configurations from the two experiments improved on the simulated ET for each individual statistical metric of performance across all sites and years, relative to the Control model (not shown in Fig. 3). ET deviations (i.e NMSE) were reduced by c. 63% in the best calibration and by c. 54% in the best climate configuration, which reflects an increase in temporal coincidence with the observations. Accuracy increased (i.e. the MAE decreased) by c. 29% for the best calibration and by c. 23% for the best climate configuration. The error in variability (i.e. SD metric) decreased by c. 41% for the best calibration and by c. 24% for the best climate configuration. Finally, the Profit_{max} displayed increased ability in better capturing the tails of the distribution, by 14% and 13% for the p_5 metric (cf. Methods 6), and by 54% and 26% for the p_{95} metric, for the best calibration and the best climate configuration, respectively.

We expected a closer fit between the observed and calibrated Profit_{max} model's ET flux in the non-drought years (2002 and 2005), because k_{max} was not calibrated on the drought periods. Yet, at three sites (Parco Ticino, Puéchabon, and Roccarespampani1) the calibrated Profit_{max} model better simulated ET in the drought years than in 2002 and 2005. Relative to the Control model, improvements in the simulation of ET were larger in the drought years than in 2002 and 2005, for both the Profit_{max} model's best calibration and its best climate configuration. In the drought years, the Profit_{max} model outperformed the Control model for ET at eight out of 10 sites. In the non-drought years, it did so at seven sites for the best calibration and at six sites for the best climate configuration

(cf. Fig. 3). Figures 4 and 5 show timeseries comparisons of the Control model with the Profit_{max} model's best calibration between April – November during the drought years (the best climate configurations are shown in Figs. 6 and 7). Large improvements in the simulated simulation ET are evident at Hyytiälä, Parco Ticino, Roccarespampani1 and El Saler. At Hesse, the Profit_{max} significantly reduced ET biases (cf. Figs. 4n, 6n and S7n before June), but was still outranked by the Control (cf. Fig. 3).

2. Model behaviour

In the Control model, under well-watered conditions, photosynthetic water use efficiency (WUE) is set based on the empirical g_1 parameter and stomatal sensitivity to D (see Methods S4). In the Profit_{max} model, the instantaneous WUE varies optimally depending on the potential to increase carbon gain versus the incurred risk of hydraulic function loss, even under well-watered conditions. Figure 8a shows the effective g_1 parameter implied by the behaviour of the Profit_{max} model across the 10 sites. At Roccaresmpampani 1 and 2, the Profit_{max} model suggests a more conservative water use strategy than the Control model (lower g_1) to better match the observations. Conversely, the Profit_{max} model implies a more profligate water use strategy (higher g_1) at Hyytiäla. In either case, the resulting changes in E do not linearly translate to marked changes in GPP because of the non-linear relationship between g_s and A_n (e.g. Figs. 4a, b and 5e, f). Indeed, when stomata are fully open, A_n is primarily limited by the rate of ribulose-1,5-bisphosphate (RuBP) regeneration (see Methods S1) and thus relatively insensitive to variations in stomatal conductance.

Behaviour differences between the models are also explained by their contrasting sensitivities to D. The Profit_{max} approach does not *a priori* set the sensitivity to D, whereas the Control model assumes g_s to be proportional to $1/\sqrt{D}$ in the absence of water stress (Medlyn *et al.*, 2011). Both panels of Figure 8 show a wide variation in the sensitivity of g_s to D across the 10 sites, varying between 0.48 and 0.84, and 0.62 on average in line with the findings of Oren *et al.* (1999).

To understand how the Profit_{max}'s instantaneous regulation of WUE affects plant water use during the growing season, we analysed the best calibration's partitioning of ET into transpiration and soil evaporation, compared to that of the Control model. At all sites besides Roccarespampani 1 and 2, the

ratio of transpiration to ET was increased and the ratio of soil evaporation to ET was decreased. These changes in partitioning led to delays in the overestimations of the rate and magnitude of dry-down simulated by the Control model (seen in many LSMs; Ukkola *et al.*, 2016) by up to two months (e.g. Fig. 5f, p in the Profit_{max}), on par with the observations. Figure 9 illustrates how the Profit_{max} mechanistically shifted the plants' ability to transpire as a function of soil water availability, from where canopy gas exchanges are maximised to where plants begin to wilt. This led to greater transpiration under drier conditions (e.g. Fig. 5j) whereby, at seven out of the 10 sites (not shown), the stomata were still not fully closed past the threshold for the wilting point of -1.5 MPa commonly used in LSMs (see Fig. 9 for an example at two sites). By contrast, the soil water potential never dropped below -0.9 MPa in the Control model.

Crucially, the Profit_{max} model's ability to simulate more realistic WUEs (e.g. Figs. 4b, p and 5b, p) and to transpire for longer could lead to more realistic annual ET and GPP modelled fluxes. Puéchabon excepted, the best calibration's total ET estimates between April – November were always closer (> 75% closer at five out of 10 sites) to the observations than the Control model's. For example, at Hyytiälä, the calibrated Profit_{max} model simulated a total ET of 562 mm between April – November, compared to 537 mm and 253 mm in the observed data and Control model, respectively; over that period, its total simulated GPP was 1620 g C, compared to 2055 g C and 1423 g C in the observed data and Control model, respectively.

3. Predicted stand k_{max}

In this study, we explored whether each site's stand k_{max} could be derived from the multi-decadal (30-year) climate, assuming hydraulic adjustments on that timescale. Figure 10 shows the wide range of k_{max} values (0.08–2.53 mmol m⁻² s⁻¹ MPa⁻¹) obtained from the two climate scenarios (Average and Extreme). We found those values to be in good agreement with the literature (cf. Notes S3 and Table S6 for a qualitative comparison). Even though the calculations of k_{max} were performed without knowledge of site precipitation, the linear regressions (cf. dotted lines) show increases in k_{max} with increasing Mean Annual Precipitation (MAP; see Table 2), suggesting predictable variation in the hydraulic traits across sites. The regression slope was the steepest for the Extreme scenario, with k_{max}

values ranging c. 2.8 times those of the Average scenario. Note that the relationship between k_{max} and MAP can partially be explained by the relationship between P_{50} and MAP (Li $et\ al.$, 2018), but that latter relationship alone only explains c. 1/3 of the degree of predictability in k_{max} for a given MAP because species-level values of P_{50} weaken any direct link between P_{50} data and site MAP. The expected relationship between LAI and MAP (Yang $et\ al.$, 2018) is not a good candidate to explain more of the predictably, as the relationship between k_{max} and MAP holds irrespectively of changes in LAI.

Where the best climate predicted k_{max} broadly agreed with the calibrated k_{max} , the Profit_{max} model's climate configuration also had the ability to outperform the Control model (cf. Fig. 3). The calibrated k_{max} exceeded that of the Extreme scenario at Hyytiälä and Loobos, but otherwise lay between the values of the Average and Extreme scenarios. The $k_{max,high}$ and $k_{max,low}$ were often smaller than the $k_{max,opt}$ (e.g. Hesse in Fig. 10), as we would expect from species trying to avoid the onset of xylem embolism (Köcher *et al.*, 2012), but this was not always the case (e.g. Puéchabon where the $k_{max,opt}$ was enveloped by the $k_{max,high}$ and $k_{max,low}$, indicating higher drought-tolerance). Overall, no unique climate-driven (i.e. Average or Extreme) or behaviour-driven (i.e. $k_{max,high}$, $k_{max,opt}$, or $k_{max,low}$) pattern explained the improved model performance.

IV. Discussion

Climate models are particularly challenged when it comes to projecting how drought will change in the future (Orlowsky & Seneviratne, 2013). A number of offline LSMs have been shown to dry out too quickly (Martínez-de la Torre *et al.*, 2019; Ukkola *et al.*, 2016) and coupled-climate models markedly diverge from observations (Ukkola *et al.*, 2018) due to differences in their representation of soil hydraulic processes and vegetation water stress during drought. As root-zone soil moisture availability decreases, the partitioning of net radiation at the vegetated land surface increases sensible heat relative to latent heat. Therefore, it is likely that if climate models incorrectly represent vegetation responses to drought, they also erroneously represent feedbacks to the boundary layer (Donat *et al.*, 2018). This hinders the capacity to capture any land surface amplification of climate extremes (Miralles *et al.*, 2018; Yunusa *et al.*, 2015).

As an alternative to approaches widely used in LSMs, we investigated an optimisation approach that considers an evolving trade-off between hydraulic cost and carbon gain to limit vegetation function during periods of water stress. When calibrated (via k_{max}), the Profit_{max} model was able to outperform our Control model, largely improving the simulated ET at eight out of 10 sites (cf. Fig. 3). Importantly, the Profit_{max} model, with its parameterised behaviour emerging from measured hydraulic traits, showed enhanced skill both outside and during drought. During drought, the k_{max} model parameter estimated from long-term site climate also led to improvements in simulated ET at nine out of 10 sites. The positive nature of this evaluation at the ecosystem scale opens the door to the incorporation of hydraulic optimisation approaches in models.

1. Can k_{max} be derived from climate?

The Profit_{max} model relies on key hydraulic traits: values of water potential relating to a specific percentage loss of conductivity (e.g. P_{50}) and k_{max} . P_x traits are widely measured; it is less clear how one might reliably estimate k_{max} at the ecosystem scale. k_{max} is a conceptual representation of the maximum conductance that plants could achieve given their ability to access water under the most favourable conditions, but it has not been commonly reported/derived in existing plant hydraulic literature. Here, we sought to explore whether k_{max} varied predictively with a measure of climate (i) because there is evidence of plant trait adjustments with climate (Atkin & Tjoelker, 2003; Carins Murphy *et al.* 2012; Marchin *et al.*, 2016; Mencuccini, 2003), and (ii) because it led to a more parsimonious model with fewer parameters.

Although MAP is a broad measure of water availability which ignores ground-water effects, we found that k_{max} increased with increasing MAP (cf. Fig. 10). This finding makes physical sense because a stand of plants with low water availability would be limited in its ability to draw water from the soil and to conduct it through to the canopy, compared to a similar stand at a location with higher water availability. We also expected k_{max} to increase with vapour pressure deficit (cf. Fig. 2a), yet be inhibited at high temperatures (cf. Fig. 2b). Across our 10 sites, we found that the Extreme climate scenario consistently predicted the highest values of k_{max} , meaning that D was the primary predictor of k_{max} in the model.

Assuming co-variation between k_{max} and $V_{cmax,25}$, we estimated k_{max} for a single value of $V_{cmax,25}$. In reality, it is likely that k_{max} varies with $V_{cmax,25}$, for instance across seasons (Wilson *et al.*, 2000), depending on factors such as climate and soil properties (which likely influence rhizosphere conductance). k_{max} is also likely to be coupled to plant allocation in individual trees (e.g. root to leaf area ratio) and to LAI and/or tree density changes at the community scale. Further difficulty arises from the range of hydraulic behaviours observed across phenotypes and within species (Aranda *et al.*, 2005), with plants adjusting their function to different climatic thresholds and/or at different timescales. At intermediate levels of precipitation (between 700 – 800 mm y⁻¹), diverse hydraulic strategies existed for similar MAPs (cf. Fig. 10).

Our attempt to estimate k_{max} based on climate is a promising proof of concept. We could further explore alternative climate descriptors, or scenarios, and alternative definitions of the stand composite LAI (which we assumed to be a fixed value), to derive k_{max} . Future work may otherwise include exploring the relation between k_{max} , (optimal ecosystem) climate, and vegetation height, not unlike Liu *et al.* (2019) who showed some coordination between hydraulic traits and maximum plant height across biomes. Approaches relying on measurements of the mean diameter of xylem, which correlates with hydraulic conductivity (cf. Hagen-Poiseuille law; Limousin *et al.*, 2010), could also be explored as they have previously been tested with some success (Cruiziat *et al.*, 2002). Finally, the use of data-driven approaches, like Bayesian model emulation (Fer *et al.*, 2018), might be an avenue for the calibration of hydraulic traits, allowing optimisation schemes to be more widely used in global LSMs.

2. Improving the simulation of GPP

Marked improvements in simulated ET, using the Profit_{max} model, were not consistently followed by marked improvements to the simulation of GPP. Selecting the best configurations (see Table 3 for a reminder on what consists in a configuration) based on ET alone was likely to impair the GPP simulations because changes in ET do not linearly translate into changes in GPP. Here, remarkably, the GPP simulations were not degraded. Nevertheless, the recovery of the vegetation following an episode of drought was largely instantaneous (e.g. Fig. 5n). Indeed, the Profit_{max} model only considers instantaneous fluxes and does not directly incorporate mechanistic links between loss of hydraulic function and key turnover of plant tissues (e.g. leaves; see Wolfe *et al.*, (2016) and Xu *et al.*, (2016)).

Since we prescribed LAI based on multi-year climatologies, the model could not capture large observed drops in GPP for a given year. For example, at Puéchabon, where previous years' water stress and drought legacy effects have been shown to affect leaf production in the subsequent years (Rambal *et al.*, 2014), the model failed to capture the GPP drops observed after 2003 (see the Control and Profit_{max} models in Figs. 5c and S8c) because it overestimated LAI in scaling from the fluxes from the leaf-level up.

3. Implementation into state-of-the-art LSMs

Given the good performance of the Profit_{max} model, implementing it into state-of-the-art LSMs to improve the representation of drought-vegetation feedbacks is an attractive option. However, it should be noted that our implementation of the profit maximisation approach assumes that the cavitation can be fully recovered upon soil water recharge (as in other hydraulic schemes, e.g. Xu *et al.*, 2016). Whilst this assumption is inconsistent with the plant hydraulic literature showing that hydraulic repair is not routine in trees (Cochard & Delzon, 2013; Delzon & Cochard, 2014), it is consistent with LSMs that do broadly not account for legacy effects. Incorporating legacy effects is a research gap for the LSM community.

Optimisation models can be perceived as computationally expensive for a climate model. Recently, Eller *et al.* (2018) proposed a simpler implementation of the Sperry et al. (2017) model with the Stomatal Optimisation Model based on Xylem Hydraulics (SOX), by removing the continuous transpiration stream. SOX changed the fundamental optimisation question asked by Profit_{max} from 'what is the cavitation risk plants are willing to take to maximise carbon intake?' to 'what is the maximum carbon intake plants can achieve given a set hydraulic cost?'. We do not know how the practical simplification made by SOX might affect model behaviour across ecosystems, so it is important for future work to compare both assumptions and explore their relative merits. Until such a comparison is made, we argue in favour of maintaining a full optimisation on the transpiration stream. Besides, Venturas *et al.* (2018) have shown how the Profit_{max} could be used to infer plant mortality, with the advantage of moving past prescribing mortality thresholds (Mencuccini *et al.*, 2019).

In our implementation, we used computationally optimised matrices to solve the optimisation (see code), keeping the computation expense small. We also provide a framework to vary: (i) the

frequency of instantaneous profit maximisation (i.e. Eqn 1 every 30 minutes or longer), (ii) the solving window on the transpiration stream (i.e. instead of Ψ_s to Ψ_{crit} , Ψ_s to $\Psi_{leaf,opt}(t-1) \pm x\%$ with t-1 being the previous timestep and the assumption that $\Psi_{leaf,opt}(t)$ would not depart by more than x% from the previous one), and (iii) the resolution of the transpiration stream itself (i.e. the increment between two adjacent Ψ). Preliminary testing (not shown) did not indicate any significant benefit from increasing the resolution of the transpiration stream (i.e. iii). Issues arose when the solving window on the transpiration stream was too narrow (i.e. ii). Lastly, whilst optimising up to every third hour did not significantly impact the simulations (i.e. i), daily optimisations flattened the model's response to environmental conditions, because they under-estimated variations in leaf water potential throughout the day.

Finally, we could apply the $Profit_{max}$ across sites globally, to generate functions to reduce stomatal conductance with decreasing soil water (e.g. Fig. S9). These functions could then be embedded in LSMs in place of the β functions. While such an approach would be empirical, it would maintain a traceable link to measurable hydraulic traits at no added computation cost to LSMs. This approach will be the subject of future work, because globally connecting hydraulic traits and water limitations on ET in land surface models used in climate models would reduce existing weaknesses during periods of water stress.

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Authors Contributions

MEBS, MGDK, and AJP designed the experiments. AMU pre-processed the eddy covariance data. MEBS developed the theory, implemented the model, and performed the analysis. MEBS, MGDK, AJP, BEM, AV, AMU, and GA all contributed to the final manuscript.

TablesTable 1. Summary of the 10 eddy-covariance sites

Site name	Country	Latitude	Longitude	PFTa	Dominant Species	Datab	Reference
Hyytiälä	Finland	61.85°N	24.29°E	ENF	Pinus sylvestris	FN	Rinne et al. (2000)
Sorø	Denmark	55.49°N	11.64°E	DBF	Fagus sylvatica	FN	Pilegaard et al. (2001)
Loobos	Netherlands	52.17°N	5.74°E	ENF	Pinus sylvestris	FN	Moors (2012)
Hesse Forest-	France	48.67°N	7.07°E	DBF	Fagus sylvatica	LTc	Granier et al. (2008)
Sarrebourg							
Parco Ticino	Italy	45.20°N	9.06°E	DBF	Populus x	FN ^c	Valentini & Miglietta
Forest					canadensis		(2015)
Puéchabon	France	43.74°N	3.60°E	EBF	Quercus ilex	FN	Rambal <i>et al.</i> (2004)
Roccarespampani1		42.41°N	11.93°E				
Roccarespampani2	Italy	42.39°N	11.92°E	DBF	Quercus cerris	FN	Rey et al. (2002)
El Saler1	Spain	39.35°N	0.32°W	ENF	Pinus halepensis	LTc	Kivimäenpää et al.
							(2010)
Espirra	Portugal	38.64°N	8.60°W	EBF	Eucalyptus	LT^{c}	Rodrigues et al.
					globulus		(2011)

^a Plant functional types (PFTs) are defined as: evergreen needleleaf forest (ENF), evergreen broadleaf forest (EBF), deciduous broadleaf forest (DBF)

^b Datasets are either FLUXNET2015 (FN) or LaThuile (LT)

^c Data for the years 2002 and 2003 only are used, either due to a lack of availability in the later years (LT dataset), or to missing data (at Parco Ticino Forest)

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Table 2. Climate information and average growing season weighted Leaf Area Index (LAI_{gs}) at the 10 eddy-covariance sites

Site name	Köppen Climate	MAPb	$T_{air,avg}^{b}$	D_{avg}^{b}	$T_{air,xx}^{,\mathbf{b}}$	$D_{\chi\chi}^{b}$	LAIgsc
	Classa	(mm y ⁻¹)	(°C)	(kPa)	(°C)	(kPa)	$(m^2 m^{-2})$
Hyytiälä	Dfc	570	9.88	0.34	20.20	1.19	1.34
Sorø	Cfb	568	12.82	0.31	19.70	0.77	1.58
Loobos	Cfb	778	13.40	0.34	22.00	1.14	1.34
Hesse Forest-Sarrebourg	Cfb	753	14.29	0.48	23.50	1.36	2.31
Parco Ticino Forest	Cfa	1026	18.14	0.69	28.00	1.95	0.76
Puéchabon	Csa	772	17.56	0.77	28.00	2.39	1.13
Roccarespampani1	Csa	675	19.71	0.79	29.40	2.13	1.83
Roccarespampani2							1.64
El Saler1	Csa	383	21.29	0.90	30.00	2.21	0.67
Espirra	Csa	736	20.02	0.84	29.00	2.34	0.85

^a Köppen Climate Classes are defined as: continental without dry season and with cold summer (Dfc), temperate without dry season and with warm summer (Cfb), temperate without dry season and with hot summer (Cfa), temperate with dry and hot summer (Csa)

^b MAP (Mean Annual Precipitation), average (*avg*) and extreme (*xx*) temperature and vapour pressure deficit values are calculated using the CRU TS v4.03 datasets (Harris *et al.*, 2014) over the 1972-2002 period

^c Average weighted sunlit-shaded LAI values are calculated over the growing season (i.e. April – November) following the procedure detailed in the Supporting Information Methods S7b.

Table 3. Summary of the two experiments conducted to estimate the maximum hydraulic conductance (k_{max}) and of the model configurations evaluated within each experiment

	Experiments						
	Calibration	Adjustments to climate					
	25 values around	six k_{max} values: three solutions \times two climate scenario					
Profit _{max}	the $k_{max,opt}$ at	$k_{max,high}$	$k_{max,opt}$		$k_{max,low}$		
Configurations	$T_{air} = 25$ °C and -		i		j		
1	D = 1 kPa	Average Scenario		Extreme Scenario			

Figure Captions

- Fig. 1. An example of the instantaneous profit maximisation algorithm. The carbon gain (green), hydraulic cost (purple), and net profit (blue) are shown as functions of the transpiration stream, which ranges between the soil water potential at saturation ($Ψ_{sat}$) and the critical water potential ($Ψ_{crit}$). Maximum hydraulic conductance (k_{max}) was calculated for each of the three behavioural solutions (i.e. $k_{max,high}$, $k_{max,opt}$, and $k_{max,low}$; cf. Section 2 of the Materials and Methods), before being used as an input the model. The dashed and dotted lines illustrate the impacts of alternative strategies for k_{max} on the maximum profit; the optimal leaf water potentials ($Ψ_{leaf,opt}$) at which profit is maximised span a range of 0.4 MPa between the instantaneous model run using $k_{max,high}$ and the one using $k_{max,low}$. The species used in this example is $Juniperus\ virginiana\ (P_{50} = -6.6\ MPa\ and\ P_{88} = -10.5\ MPa$; Choat $et\ al.$, 2012), with $V_{cmax,25} = 100\ \mu mol\ m^{-2}\ s^{-1}$, $T_{air} = 25\ ^{\circ}C$ and $D = 1\ kPa$, $Ψ_s = -0.8\ kPa$, and LAI = 2 m² m⁻².
- **Fig. 2.** The sensitivity of the modelled optimal coordination between the maximum hydraulic conductance ($k_{max,opt}$) and (a) the maximum carboxylation rate at 25°C ($V_{cmax,25}$) and (b) air temperature (T_{air}), both depending on vapour pressure deficit (D). In panel a, $k_{max,opt}$ increases near proportionally with $V_{cmax,25}$ and in a logarithmic fashion with D; T_{air} is fixed to 25 °C. In panel b, $k_{max,opt}$ increases with T_{air} , before decreasing (sharply at the two highest D) starting between 20 25 °C; $V_{cmax,25}$ is set to 100 μmol m⁻² s⁻¹. The valid range for $k_{max,opt}$ is constrained by physically plausible co-occurring values of T_{air} and D under a relative humidity spanning 5 95%. The species used in this example is *Juniperus virginiana* ($P_{50} = -6.6$ MPa and $P_{88} = -10.5$ MPa; Choat *et al.*, 2012), with $\Psi_s = -0.8$ kPa and LAI = 2 m² m⁻².
- **Fig. 3**. Quantile ranks of the best configurations of the Profit_{max} model compared to the Control model, across drought (panels a and b) and non-drought years (panels c and d), and for Gross Primary Productivity (GPP; panels a and c) and Evapotranspiration (ET; panels b and d). The vertical lines blue (Average scenario) or red (Extreme scenario) correspond to the best climate configuration range of ranks across the three behavioural solutions for the maximum hydraulic conductance. The box and whisker plots to the right of the figure show summaries of the ranks across sites, but do not

account for the behavioural range shown by the vertical lines. In the box and whiskers plots, the horizontal yellow line shows the average overall quantile rank, and the box shows the interquartile range; the whiskers extend to the 10th and 90th percentile values of the ranks, with dots outside of the whiskers showing outliers.

- **Fig. 4**. A 14-day running average of the carbon and water fluxes predicted by the best selected calibration configuration from the Profit_{max} model (green line) at the five northernmost eddy-covariance sites during the 2003 (panels a, b, e, f, i, j, m, n, o, p) and 2006 (panels c, d, g, h, k, l) European drought events, compared to the Control model (purple line), and to the observations (black line). Grey lines show the prescribed phenologies (LAI, m² m⁻²) and blue bars the daily precipitation (PPT, mm d⁻¹). The Gross Primary Productivity (GPP) is in g C m⁻² d⁻¹ and the Evapotranspiration (ET) in mm d⁻¹.
- **Fig. 5.** A 14-day running average of the carbon and water fluxes predicted by the best selected calibration configuration from the Profit_{max} model (green line) at the five southernmost eddy-covariance sites during the 2003 (panels a, b, e, f, i, j, m, n, o, p) and 2006 (panels c, d, g, h, k, l) European drought events, compared to the Control model (purple line), and to the observations (black line). Grey lines show the prescribed phenologies (LAI, m² m⁻²) and blue bars the daily precipitation (PPT, mm d⁻¹). The Gross Primary Productivity (GPP) is in g C m⁻² d⁻¹ and the Evapotranspiration (ET) in mm d⁻¹.
- **Fig. 6**. A 14-day running average of the carbon and water fluxes predicted by the best selected climate configuration from the Profit_{max} model (green) at the five northernmost eddy-covariance sites during the 2003 (panels a, b, e, f, i, j, m, n, o, p) and 2006 (panels c, d, g, h, k, l) European drought events, compared to the Control model (purple line), and to the observations (black line). The green line is the $k_{max,opt}$ strategy and the green shadings encompass the instantaneous range of fluxes predicted by the three behavioural solutions for k_{max} . Grey lines show the prescribed phenologies (LAI, m² m⁻²) and blue bars the daily precipitation (PPT, mm d⁻¹). The Gross Primary Productivity (GPP) is in g C m⁻² d⁻¹ and the Evapotranspiration (ET) in mm d⁻¹.
- **Fig. 7.** A 14-day running average of the carbon and water fluxes predicted by the best selected climate configuration from the Profit_{max} model (green) at the five southernmost eddy-covariance sites during

the 2003 (panels a, b, e, f, i, j, m, n, o, p) and 2006 (panels c, d, g, h, k, l) European drought events, compared to the Control model (purple line), and to the observations (black line). The green line is the $k_{max,opt}$ strategy and the green shadings encompass the instantaneous range of fluxes predicted by the three behavioural solutions for k_{max} . Grey lines show the prescribed phenologies (LAI, m² m⁻²) and blue bars the daily precipitation (PPT, mm d⁻¹). The Gross Primary Productivity (GPP) is in g C m⁻² d⁻¹ and the Evapotranspiration (ET) in mm d⁻¹.

Fig. 8. A comparison of the sensitivity of the Control and calibrated Profit_{max} models' stomatal conductance (g_s) to vapour pressure deficit (D) across the 10 eddy-covariance sites. Panel (a) shows the relationship between the implied water use efficiency $(g_1, kPa^{0.5})$ of the calibrated Profit_{max} model and its sensitivity to D (σ , unitless). The values of g_1 obtained for the Profit_{max} model were converted from unit kPa^{σ} to unit $kPa^{0.5}$ for comparison with the values of g_1 used in the Control model. g_1 and σ were obtained by least-square fitting of the g_s simulated by the calibrated Profit_{max} model to the Medlyn *et al.* (2011) stomatal conductance model. The estimates were produced using the site hydraulic and photosynthetic parameters, for temperatures ranging 10 - 40 °C and D ranging 0.05 - 3 kPa. The values of g_1 used in the Control model are plotted against the respective sites' σ for visual comparison with those of the Profit_{max} model only, as they correspond to $\sigma = 0.5$. Panel (b) shows the effect of the various σ on the g_s given by the Control model at 25 °C. The input parameter g_1 was set to 2 $kPa^{0.5}$ for all the generated curves, but it was transformed to kPa^{σ} upon running the Control model with the site-specific values of σ shown in panel a. The reference g_s of the Control model $(\sigma = 0.5)$ is plotted for comparison. For both panels a and b, the models were run assuming well-watered conditions.

Fig. 9. A comparison of the decline in stomatal conductance (g_s) with predawn soil water potential (Ψ_s), for the Control (plain lines) and the calibrated Profit_{max} (dotted lines) models at a sub-selection of sites. The functional forms emerge from the soil parameters and the β functions in the Control, and from the plant hydraulic traits and the profit maximisation algorithm in the best selected calibration. The inset zooms on the functional forms of the g_s - Ψ_s curves from the Control model for Ψ_s > -0.3 MPa. The functional forms are made comparable by normalising g_s by its maximum at a given site.

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Note that seemingly slow decreases in g_s with Ψ_s can be attributed to the non-linear relationship between Ψ_s and volumetric soil moisture, whereby small variations in the latter can lead to large variations in the former. To avoid rainfall effects, the data up to 48 hours after rain were excluded. To avoid low solar radiation and low temperature effects, the g_s data were restricted between 9:00 h – 15:00 h from April – November across all years. The curves were fitted with a linear generalised additive model and the shadings show the 95% confidence interval of the fit.

Fig. 10. The estimated site maximum hydraulic conductance (k_{max}), for each climate configuration of the Profit_{max} model and for the best selected calibration, shown as a function of Mean Annual Precipitation (MAP; as listed in Table 2). Note, the MAP was not used in the estimation of k_{max} , however k_{max} was multiplied by the sites' weighted composite LAI, which normalises it to ground area and makes it comparable across sites. Linear regressions are used to show the positive relationship between k_{max} and MAP, with a r^2 of 0.53 and a p-value of 0.02 for the best selected calibration, a r^2 of 0.21 and a p-value of 0.01 for the Average climate scenario, and a r^2 of 0.30 and a r-value of 0.002 for the Extreme climate scenario.

Supporting Information

Additional Supporting Information is available for this article.

- Fig. S1 A schematic showing the modelling experiments.
- **Fig. S2** A 14-day running average of the carbon and water fluxes predicted by the Control model at the five northernmost eddy-covariance sites during the 2003 and 2006 European drought events, compared to the CABLE LSM and to the observations.
- **Fig. S3** A 14-day running average of the carbon and water fluxes predicted by the Control model at the five southernmost eddy-covariance sites during the 2003 and 2006 European drought events, compared to the CABLE LSM and to the observations.
- **Fig. S4** The predawn volumetric soil water available to the vegetation, as simulated by the Control model and by CABLE, at the five northernmost eddy-covariance sites during the 2003 European drought event.
- **Fig. S5** The predawn volumetric soil water available to the vegetation, as simulated by the Control model and by CABLE, at the five southernmost eddy-covariance sites during the 2003 European drought event.
- **Fig. S6** A 14-day running average of the carbon and water fluxes predicted by two different calibrations of the Control model at a sub-selection of sites in 2002, 2003, 2005, and 2006, compared to the calibrated $Profit_{max}$ model, to the reference Control model, and to the observations.
- **Fig. S7** A 14-day running average of the carbon and water fluxes predicted by the best selected Calibration at the five northernmost eddy-covariance sites in 2002 and 2005, compared to the Control model and to the observations.
- **Fig. S8** A 14-day running average of the carbon and water fluxes predicted by the best selected Calibration at the five southernmost eddy-covariance sites in 2002 and 2005, compared to the Control model and to the observations.

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Fig. S9 The stomatal conductance as a function of the predawn total volumetric soil water available to the vegetation.

Table S1 Soil parameters at the 10 eddy-covariance sites

Table S2 Plant trait inputs at the 10 eddy-covariance sites

 Table S3
 Surface properties per plant functional type (PFT)

Table S4 Water available in the five soil-sub layers located below the soil top layer

 Table S5
 Parameters used in the biochemical photosynthesis model

Table S6 Predicted ranges of k_{max} at the 10 sites compared to species-specific measured values of

 k_{max}

Methods S1 Biochemical photosynthesis model

Methods S2 Energy balance model

Methods S3 Shape of the vulnerability curves

Methods S4 Coupling carbon and water

Methods S5 Scaling from leaf to canopy

Methods S6 Soil hydrology

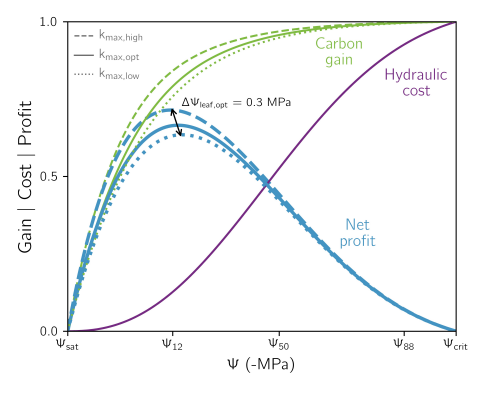
Methods S7 Prescribed LAI

Methods S8 Parameter calibrations

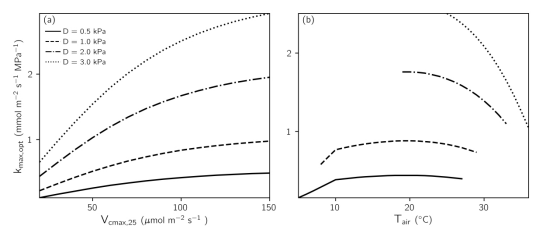
Notes S1 Comparison of the Control to CABLE

Notes S2 Why did we not calibrate the Control model?

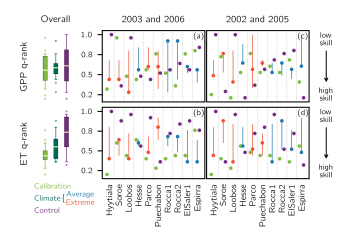
Notes S3 Comparison of the predicted values of k_{max} to the literature



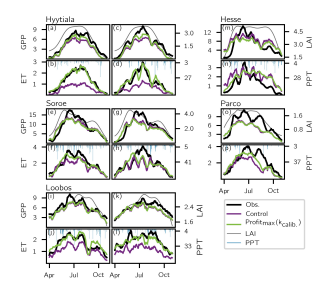
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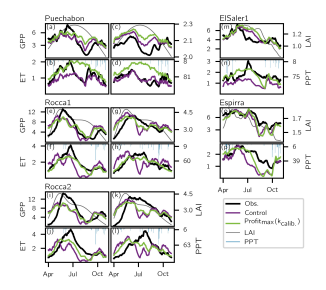
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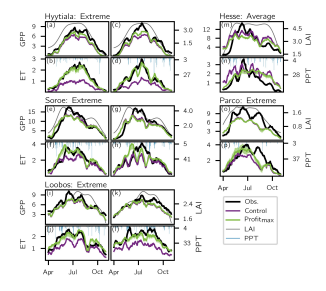
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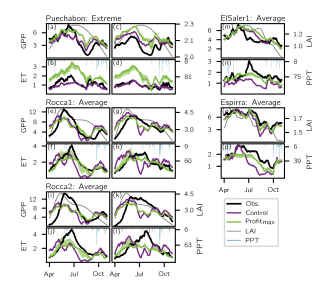
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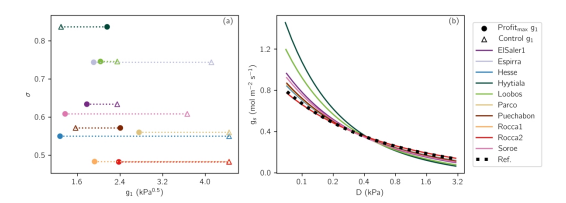
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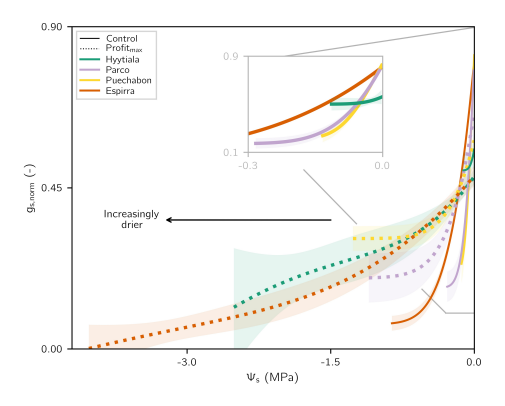
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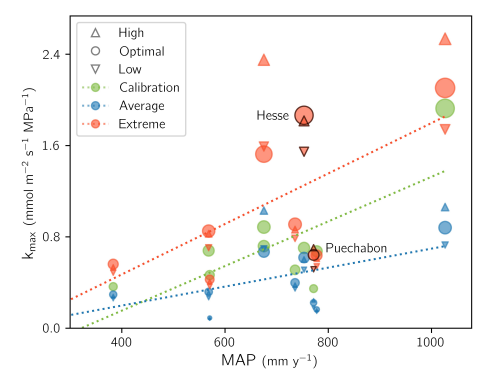
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