

AUTHOR RESPONSE

The manuscript for our paper *The importance of radiation for semi-empirical water-use efficiency models* has been revised according to the reviews. We carefully considered the suggestions made by three anonymous referees. We integrate both the minor comments regarding spelling, coherence and citations, as well as general comments made by the three referees. The proposed changes have improved the manuscript substantially, both in its intelligibility and its implications.

The major change has been the addition of a stratification of all included sites by type of vegetation, as suggested by Referee #2. The revised manuscript now addresses differences between low and high vegetation types for the two parameters $uWUE$ and r , as well as the metric ET_frac . The results of these analyses have given additional credence to the proposed explanation that equilibrium evaporation is responsible for observed effects of radiation on transpiration.

Substantial changes have been made to the supplementary materials, which now includes multiple suggestions made by the referees.

The introduction has been revised for improved clarity.

Upon consideration, we think that model limitations are adequately addressed in our discussion section. To highlight this important part, we introduced a new subsection, "4.2 Limitations".

Below, you can find our responses to the three referees, as we have made published them for each referee comment (see the respective DOI for the full comments that include the figures).

Thank you for your assistance in improving this manuscript.

Best,
Sven Boese on behalf of the co-authors

Referee 1

Please refer to our original response to Referee #1 for the full author comment including the referenced figures.

AC1: doi:10.5194/bg-2016-524-AC1

We thank Referee #1 for the positive and constructive appraisal of our article! Below, we respond to the general and specific points of the review.

Metrics For Model Evaluation

“But MEF does not provide a quantitative measure of the increase in the amount of variance explained by the successive modifications to the model. Table 2 could contain such an estimation, based i.e. on the (average) changes in the RMSE.” We agree that the Nash-Sutcliffe Efficiency (MEF) captures only a part of the overall model performance. Specifically, the absolute magnitude of the errors is not reflected in this measure. We therefore added an complementing table for the cross-validated RMSE (Fig. 1 of this response) to the supplementary materials.

Negative Intercepts

“The only problem in the model definition is that the intercept is left to be negative, which has no biological meaning [...]” We also think that negative intercepts are biologically implausible. A squared intercept caused problems in the optimization, but it was possible to use a constrained optimization that limits the intercept parameter range to positive values. As can be seen in the attached plot (Fig. 2 of this response), the differences in MEF between the original model +ETres and the bounded variant +ETres_bnd is miniscule. Because it precludes biologically implausible values, we revised all relevant plots for the manuscript to account for this.

Nonlinearity

“It remains unclear why the effect of the radiation was modelled as a linear” This is a valid point! We previously tested nonlinear models, but did not detect any notable effect on model performance. However, the decision to only address linear models can appear ad hoc. We now refer to the suggested “observed–predicted” (see below) plots to support our choice of a linear response to radiation. We also included a nonlinear model variant (“+Rg_nl”) of the form $ET =$

$(GPPVPD^{0.5}/uWUE) + rRG^a$, where $a = 0.7$ in Fig. 2. This could represent a possible, gradual saturation effect for higher radiation levels. However, including this nonlinear response yielded inferior results.

Illustrating the Improvement of the Models

“Overall the manuscript has the tendency to not display the data and the relationships between them. Showing, for a couple of examples, the gain in having an intercept and incorporating radiation in the modelling would be great.” We concur that the previous version of the manuscript did not sufficiently show the actual predictions but rather metrics reflecting the skill of the employed models. We selected two sites for which we detail how the different model formulations affect the predictions. We originally plotted this as time series. However, the difference between the models were hard to extract, as our selection of data points for the estimation yields a very sparse and irregularly sampled time series. Therefore, we propose to plot observed against predicted values with a one-to-one line indicating perfect model fit. Two such plots, with two subplots for +ETres and +Rad respectively are attached to this reply (Fig. 3a,b of this response).

Fraction of Significant Intercepts

“How many times (or in percent) has the intercept ETres been found significant?” This is an important number that is now part of the revised manuscript. The estimated intercepts were significant at 86 sites, which is 78% of all 110 sites considered. We added this number in the appropriate place in the manuscript.

In addition, all specific points referring to spelling, coherence, terminology and citations were considered and integrated in the revised manuscript.

Thank you again for your assistance in improving this paper!

Referee 2

Please refer to our original response to Referee #2 for the full author comment including the referenced figures.

AC2: doi:10.5194/bg-2016-524-AC2

AC4: doi:10.5194/bg-2016-524-AC4 (missing references)

We thank Referee #2 for the positive and constructive appraisal of our article! Below, we respond to the general and specific points of the review.

Tropical Climates and Ecosystems

“But, what is author opinion, is it possible to apply suggested WUE model to large tropical areas and especially to tropical rain and monsoon forests? Or not?” For all its extensive coverage of northern latitudes, the FLUXNET is limited regarding a representation of tropical sites. This exacerbated by our choice of sites with a “Fair and Free Use” data policy. The only tropical site in our pool was the Australian site AU-How. However, we surmise that the observed effects are not restricted to extra-tropical regions, as the potential factors driving them are likely of physical nature and not due to specific processes limited to certain plant types or ecosystems. Yet one potential limitation for tropical areas with frequent precipitation events is the necessary filtering for successively rain-free periods. This would, of course, be less a problem for the mentioned monsoon climates with pronounced dry-seasons. The revised manuscript now alludes to how our inferences can only be cautiously extrapolated beyond the coverage of our subset of FLUXNET sites.

Effect of Vegetation Structure

“Is there any difference in found transpiration - GPP relationships between forest and grassland sites?” Referee #2 raises an important question regarding potential differences between different kinds of vegetation structures, such as grasslands and forests. Following up on this suggestion, we partitioned all sites in two classes: Low vegetation for grasslands, crops and savannas and high vegetation for all other vegetation types. When stratifying the data set like this, we found that $uWUE$ was not significantly different for either vegetation type (Fig. 1). However, we noted that grasslands and crops had a significantly higher mean value of r (Fig. 2). This is a relevant finding, as it supports our proposed explanation that the radiation effect could be a sign of equilibrium transpiration (Jarvis and McNaughton, 1986). In a preceding study, McNaughton and Jarvis (1983) report that grasslands had a higher decoupling parameter Ω , which quantifies the contribution of equilibrium evaporation. As Jarvis and McNaughton (1986) discuss, a stronger atmospheric decoupling (high Ω) implies a higher relative share of equilibrium transpiration. Therefore, we repeated the analysis of the fraction of radiation-associated transpiration (Fig. 3) and found that this metric, ET_frac , was significantly higher for the low vegetation PFTs grassland and crops (0.53, 95% CI: 0.48–0.58) compared to high vegetation (0.39, 95% CI: 0.34–0.44). We revised and adapted the manuscript accordingly!

Dewfall

“What is about dew formation and its evaporation? Is it ignored?” We in fact did not consider dewfall in the original article. The evaporation of dewfall would likely be very dependent on radiation, thereby violating our assumption that the observed latent heat flux represents transpiration and hence confounding our estimates of radiation sensitivity. To verify that our results were not biased by potentially including days with dewfall, we stratified the data set according to the relative humidity during night-time. As high relative humidity is a necessary, but not a sufficient criterion for dewfall, we consider this a conservative rule for which days are likely unaffected by any dewfall. We then estimated the term ET_{res} separately for all sites that had had observations for the respective RH intervals (Fig. 4), finding that the mean ET_{res} for all sites was insensitive to this variable. We also found that excluding days with high relative humidity caused only very minor changes in the mean cross-validated model-efficiencies.

Effect of LAI

"Did you analyze the relationships between contributions of soil evaporation to ET and canopy LAI? LAI is an important factor for models of evapotranspiration. With the selection of rain-free periods, we assumed that both interception and bare-soil evaporation would be negligible in our analysis. To verify that our observed patterns are nevertheless merely the result of open canopies, we performed an analysis in which we filtered for successively higher LAI observations. In the corresponding figure (Fig. 5), we show the smoothed response of mean MEF over all sites for both the Zhou and the +Rg model. As the difference in performance actually *widens* for higher LAI values, we conclude that the observed patterns are unlikely the product of open vs. closed canopies. As we discuss in a preceding point, we have now better support for the explanation involving equilibrium transpiration.

In addition, all specific points referring to spelling, coherence and citations were considered and integrated in the revised manuscript.

Thank you again for your assistance in improving this paper!

References

Jarvis, P. G. and McNaughton, K.: Stomatal control of transpiration: scaling up from leaf to region, *Advances in ecological research*, 15, 49, 1986.

McNaughton, K. G., and Jarvis, P. G.. Predicting effects of vegetation changes on transpiration and evaporation. In: "Water Deficits and Plant Growth" (T. T. Kozlowski, ed.), Vol. 7, pp. 1–47. Academic Press, New York, 1983.

Referee 3

Please refer to our original response to Referee #3 for the full author comment including the referenced figures.

AC3: doi:10.5194/bg-2016-524-AC3

We thank Referee #3 for the positive and constructive appraisal of our article! Below, we respond to the general and specific points of the review.

Ambiguity of the Models.

"Ambiguity in concepts such as WUE models, physiological WUE models? Are the authors talking about stomatal conductance models? Please clarify them and provide details." Thank you for pointing this out. The revised manuscript gives a more detailed introduction into the different types of WUE models and treats the terminology more carefully.

Collinearity

"The concerns include whether and how the authors test the collinearity between the variables such as R_g and $GPP \cdot VPD_{0.5}$ in the model fitting" This is a very good remark! The high degree of correlation is an important issue for these kind of empirical analyses. This is particularly pertinent for isolating the fraction of evapotranspiration that we attribute to radiation. In the original paper, we analyzed the impact of collinearity on our results by accounting for the correlation of parameter uncertainties (Supplementary Materials S1). In the new manuscript the problem of collinearity and our treatment is given more prominence. We further moved the mentioned section from the supplement to the method section of the main document.

Overparameterization

"In addition to MEF, index such as AIC or AICc are needed to account for possible over-parameterization?" We fully agree with Referee #3 that overparameterization is an important issue in analyses focussed on model selection. We believe that we have adequately addressed this problem by

exclusively using cross-validated Nash-Sutcliffe Efficiencies (MEF) in our model comparison. Adding further parameters to a model will generally allow the model to accommodate even observations that were the result of random errors or unattributed processes. The cross-validation penalizes such a over-parameterization by iteratively testing the model's ability to predict observations that it wasn't calibrated to. Using an information criterion, such as AIC, AICc or BIC, that directly accounts for the number of parameters used in the model is another possibility to represent model complexity. AICc can be expected to converge with cross-validation asymptotically (Stone, 1977). We are therefore confident that our results are not confounded by the number of model parameters. To illustrate this, we added a table to this comment (Fig. 1 of this response) that replicates Table 1 of the original paper (Fraction of sites with a higher or lower MEF). As is the appropriate usage for AICc, we counted the fraction of sites where the pairwise difference of AICc was smaller than -2 for the model of the row to be considered superior to that of the corresponding column. The table suggests that our conclusions are not sensitive to the choice of either AICc or cross-validated MEFs.

Interaction Terms

"How the authors deal with the interactive terms among those variables." This is an interesting question. In our analysis we aimed to obtain effective, parsimonious models with sufficient biological and physical plausibility. This is why we did not test all possible combinations of predictor variables. Attached to this comment is a plot (Fig. 2 of this response) that includes both the three models of the old manuscript (Zhou, +ETres, +Rg) and three new variants for questions raised by the other referees: +ETres_bnd has parameters constrained to positive values, +Rg_nl has a nonlinear response to radiation and Introt has an interaction term of VPD with Rg. The model evaluation was performed in a comprehensive cross-validation scheme. The original +Rg variant was again confirmed to have the highest performance in this evaluation. In addition, this model is corroborated by the new results indicating higher importance of radiation for low vegetation, which makes equilibrium evaporation a plausible candidate explanation for the observed patterns (see below in our response, section PARAMETER DISTRIBUTIONS).

Introductory Definition of Models

- "p2, lines 5–15: this paragraph needs to clarify the difference between existing WUE models."
- "P2, lines 22–29: there are several confusing/incorrect statements in this

paragraph.”

- “P2, line16: what is physiological WUE models? Did the authors mean stomatal conductance models?” Thank you for pointing this out! We have revised the introduction of the paper accordingly, to better explain our approach and contrast it with existing models. We also discuss how current models include the g_0 conductance term. “the ratio GPP/ET is never constant and is considered to be proportional to v_{pd} or squared rooted v_{pd} depending on assumptions (Zhou et al., 2014)” Here, we referred to radiation, when stating that “The models implicitly assume that, at ecosystem-scale, GPP and ET respond equally to changes in radiation and that, therefore, the ratio of both is constant with regard to this factor.” This has been clarified in the revised introduction.

Effect of Water-Limitation on Collinearity

“Not sure how water limitation can affect collinearity of parameters?” Referee #3 is right that we did not provide a sufficient explanation for our reasoning here. As we mention below, the degree of correlation between the predictor variables is a property of the additive models we identified. The dependency of GPP on radiation is an obvious case for that. We expected this correlation (and the following collinearity) to decrease under water-limitation, as GPP is then no longer as easily determined by radiation. For example, during periods of extended droughts, we would expect day-to-day variability of GPP to be no longer a function of radiation and related covariates but rather variables reflecting soil-water availability. If this dependency of the covariates decreases, it would follow that the collinearity of the parameters decreases, too. However, the results were very inconclusive when adopting the aridity index (AI) that is used by the United Nations Environmental Program (defined as: $AI = \text{Precipitation} / PET$). We decided that the results were furthermore not pertinent to the main topic, which is why we decided to exclude this part from the new manuscript.

Parameter Distributions

“In results, in addition to the MEF, I would like to see the distribution of two other parameters ($uwue$ and r) of all the sites.” The updated manuscript now includes plots showing the distribution of the two parameters $uWUE$ and r . Upon a comment by Referee #2, we stratified the data-set along the vegetation structure (low for grasslands and crops, high for all other plant functional types). Quoting from our reply to Referee #1: “When stratifying the data set like this, we found that $uWUE$ was not significantly different for either vegetation type (Kolmogorov–Smirnov test) [Fig. 3 of this response]. However, we noted that grasslands and crops had a significantly higher mean value of r [Fig. 4 of this response]. This is a

relevant finding, as it supports our proposed explanation that the radiation effect could be a sign of equilibrium transpiration (Jarvis and McNaughton, 1986). In a preceding study, McNaughton and Jarvis (1983) report that grasslands had a higher decoupling parameter Ω , quantifying the contribution of equilibrium evaporation. As Jarvis and McNaughton (1986) discuss, a stronger atmospheric decoupling (high Ω) implies a higher relative share of equilibrium transpiration. Therefore, we repeated the analysis of the fraction of radiation-associated transpiration and found that this metric, ET_{frac} , was significantly higher for the low vegetation PFTs grassland and crops (0.53, 95% CI: 0.48–0.58) compared to high vegetation (0.39, 95% CI: 0.34–0.44) [Fig. 5 of this response]. We revised and adapted the manuscript accordingly!” The relevant plots have been attached to and renumbered for this comment (Fig. 3–5 of this response).

Monthly Patterns by Site

“This is interesting finding. Could it be possible for the authors to provide this similar figure for each of the sites in the supplementary materials for the readers to eyeball the site difference or similarity?” The updated supplement now contains this figure as a matrix of monthly patterns for each site individually! The plot is also attached to this comment (Fig. 6).

Covariance Assumptions

“More details are needed on the variance and covariance for each of the variables including GPP and ET, because this variance and covariance directly affect your L-M algorithm and likely results.” The optimization approach of our analyses follows eq. 5–6 in Omlin and Reichert (1999) with a σ_{meas} of 1, hence being insensitive to the uncertainties in the forcing and target variables. In agreement with Lasslop et al. (2008), this approach does not consider correlations between the errors of the original latent heat and net ecosystem exchange fluxes.

Limitation of the Models

“All the proposed models have their own assumptions and their possible violations. Please discuss them as well on how these violations could affect the results.” This is a critical aspect for a model-selection exercise such as ours and is treated more diligently in the revised version of the manuscript.

Data Availability

The data sets can be downloaded at <http://fluxnet.fluxdata.org//data/download-data/>

In addition, all specific points referring to spelling, coherence and citations were considered and integrated in the revised manuscript.

Thank you again for your assistance in improving this paper!

References

Jarvis, P. G. and McNaughton, K.: Stomatal control of transpiration: scaling up from leaf to region, *Advances in ecological research*, 15, 49, 1986.

Lasslop, G., Reichstein, M., Kattge, J., Papale, D.: Influences of observation errors in eddy flux data on inverse model parameter estimation. *Biogeosciences*, 5, 1311–1324, 2008.

McNaughton, K. G., and Jarvis, P. G.. Predicting effects of vegetation changes on transpiration and evaporation. In: “Water Deficits and Plant Growth” (T. T. Kozlowski, ed.), Vol. 7, pp. 1–47. Academic Press, New York, 1983.

Omlin, M. and Reichert, P.: A comparison of techniques for the estimation of model prediction uncertainty, *Ecological Modelling*, 115, 45–59, 1999.

Stone, M.: An asymptotic equivalence of choice of model by cross-validation and Akaike’s criterion. *Journal of the Royal Statistical Society: Series B (Methodological)*, 39, 44–47, 1977.

The importance of radiation for semi-empirical water-use efficiency models

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Abstract

Water-use efficiency (WUE) is a fundamental property for the coupling of carbon and water cycles in plants and ecosystems. Existing model formulations predicting this variable differ in the type of response of WUE to the atmospheric vapor pressure deficit of water (VPD). We tested a representative WUE model on ecosystem scale at 110 eddy-covariance sites of the
10 FLUXNET initiative by predicting evapotranspiration (ET) based on gross primary productivity (GPP) and VPD. We found that introducing an intercept term in the formulation increases model performance considerably, indicating that an additional factor needs to be considered. We demonstrate that this intercept term varies seasonally and we subsequently associate it with radiation. Replacing the constant intercept term with a linear function of global radiation was found to further improve model predictions of ET. Our new semi-empirical ecosystem WUE formulation indicates that, averaged over all sites, this
15 radiation term accounts for up to half (39–47 %) of transpiration. These empirical findings challenge the current understanding of water-use efficiency on ecosystem-scale.

1 Introduction

Understanding the coupling of carbon and water cycles is as a central question of global change research, as changes in one
20 of the cycles could directly propagate to the other (Churkina et al., 1999; Gerten et al., 2004; Ito & Inatomi, 2012). Carbon assimilation through photosynthesis (A) and transpiration (T) constitute major fluxes in these two cycles and the Earth system (Jasechko et al., 2013; Ciais et al., 2014). On the leaf-scale, *water-use efficiency* (WUE) is defined as $WUE = A/T$, quantifying the ability of a plant to assimilate atmospheric carbon dioxide per water loss by transpiration. An understanding of this ratio can hence translate into the ability to predict one of the two fluxes from the other.

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Both fluxes are limited by the stomatal conductance (g_s) of plant leaves (Cowan and Farquhar, 1977), allowing for the definition of *intrinsic water-use efficiency*, $WUE_i = A/g_s$. This quantity is less contingent on the vapor pressure deficit of atmospheric water vapor (VPD) which affects both fluxes differently. At the ecosystem scale, neither carbon assimilation nor transpiration can be observed directly. However, the eddy-covariance (EC) method (Baldocchi et al., 2001) can be used to

obtain data of the gross-primary productivity (GPP) and the evapotranspiration (ET). On ecosystem scale, water-use efficiency is then defined as $WUE = GPP/ET$, while the intrinsic water-use-efficiency is accordingly formulated as $WUE_i = GPP/G_s$, where G_s denotes the surface conductance of the ecosystem.

5 Analytical models that predict WUE on the leaf-scale (Katul et al., 2010; Medlyn et al., 2011) were derived from theoretical optimality considerations. Corresponding variants were evaluated with ecosystem-scale flux measurements gathered by the FLUXNET¹ in empirical studies (Beer et al., 2009; Zhou et al., 2014; Zhou et al., 2015). The central difference between the existing models is the response of g_s or G_s to VPD, resulting in different dependencies of WUE on VPD. The concept of *inherent water-use efficiency (IWUE)* by Beer et al. (2009) corrects for the increasing diffusion of water vapor with higher
10 values of VPD. However, the proposed IWUE is still dependent on the difference between leaf-external and leaf-internal CO₂ concentrations and therefore stomatal conductance. Physiological models (e.g. Katul et al., 2010), suggest stomatal contraction with increasing VPD, as plants aim to minimize water loss. This was found to be an important factor on ecosystem scale as shown by Zhou et al. (2014) for half-hourly and by Zhou et al. (2015) for daily observations. In both studies, the response of WUE could be approximated to be proportional to $VPD^{-0.5}$. Despite their discrepancy, both the
15 models of Beer et al. and Zhou et al. imply that, at ecosystem-scale, WUE is strictly an inverse function of VPD or $VPD^{0.5}$, respectively.

Stomatal conductance models that were derived on plant scale include an additional conductance term g_0 that is invariable to changing levels of photosynthesis (Ball et al., 1987; Medlyn et al., 2011). Any transpiration resulting from this part of
20 stomatal conductance should be expected to be proportional to the gradient of the partial pressure of water vapor, quantified by the atmospheric VPD observations. In contrast to formulations commonly used in stand, land surface and vegetation models (de Kauwe et al., 2013), this process is not considered in current ecosystem-scale WUE formulations (Zhou et al. 2014, Zhou et al., 2015).

25 Incoming solar radiation is a driving factor for both photosynthesis and transpiration. However, existing WUE models posit that the ratio of both is invariant with regard to this factor. This implicitly assumes that, at ecosystem-scale, the influence of radiation on GPP and ET cancels out, as the water-use efficiency is entirely determined by gas diffusion and its limiting factors. However, photosynthesis saturates at high radiation levels (Farquhar et al., 1980), even in well-watered conditions. Models of Potential Evapotranspiration (PET), by contrast, do not prescribe a similar limiting behavior in conditions of
30 sufficient water-availability. For example, in the Penman–Monteith equation, the evaporation rate scales linearly with the absorbed radiation, holding all other factors constant (Leuning et al., 2008). Mechanistically, the process of equilibrium transpiration (Jarvis and McNaughton, 1986) implies that sizeable transpiration can occur even when the leaf is fully

¹ <http://fluxnet.fluxdata.org/>

decoupled from the atmosphere, i.e. when VPD is very low. This is a second process that current ecosystem WUE models cannot accommodate.

In this study, we address these unresolved inconsistencies regarding the importance of additional model terms for predictions of ecosystem-scale transpiration. We do this empirically on ecosystem-scale by optimizing and assessing different WUE models with FLUXNET observations from 110 globally distributed towers. In our approach, ET is selected as target variable, while the different WUE models utilize GPP as one of multiple explanatory variables. The substantial degree of correlation between GPP and ET is thus harnessed for the predictions of ET. In a first step, we identify existing biases in ecosystem-scale WUE models. In the next step, these biases are tested for their dependency on VPD and radiation. Lastly, we infer a tentative partitioning of transpiration according to its association with radiation and discuss the substantial magnitude of this metric. [We conclude by highlighting how changes in the model structures impact the between-site variability of parameter estimates.](#)

2 Data & methods

2.1 Data

The daily day-time integrals of gross primary productivity (GPP) and evapotranspiration (ET) were taken from the La Thuile FLUXNET (open and fair use data policy sites) collection². The aggregation to day-time values was based on values of potential radiation larger than 10 W m^{-2} . Additionally, we used global radiation (R_g) and the day-time vapor pressure deficit (VPD) measured at the same eddy-covariance (EC) sites.

The EC data were processed according to the standard methods (Papale et al., 2006; Reichstein et al., 2005) to assure consistent quality of the observations. Eddy-covariance GPP results were based on the flux partitioning method of Reichstein (2005). We used only data with $\text{GPP} > 0.1 \text{ gC d}^{-1} \text{ m}^{-2}$, $\text{ET} > 0.05 \text{ mm d}^{-1}$ and $\text{VPD} > 0.001 \text{ kPa}$ to reduce the relatively large impact of random measurement errors under low flux conditions. Following the procedure of Beer et al. (2009) we further used only data after three consecutive rain-free days. This reduces contributions by evaporation to the measured evapotranspiration as physical evaporation typically declines rapidly after rain events due to the depletion of water stored in the topmost soil layer (Wythers et al., 1999). That assumption similarly applies to precipitation that is intercepted on leaf-surfaces and other plant parts in the canopy. As Miralles et al. (2010) summarize, the interception storage for forest ecosystems reported in different studies amounted to a mean of 1.2 mm ($\pm 0.4 \text{ mm}$; [per unit area of canopy cover](#)). With the mean interception evaporation rates reported as 0.3 mm h^{-1} ($\pm 0.1 \text{ mm h}^{-1}$; [per unit area of canopy cover](#)), this storage can be concluded to be typically depleted within the first days after a precipitation event. Therefore, the measured

² FLUXNET Synthesis Dataset (La Thuile 2007). Available at www.fluxdata.org

evapotranspiration after three consecutive [rain-free](#) days is expected to approximate transpiration, and we additionally verified that our results are robust when considering longer [rain-free](#) periods (see below). For the main analysis, we included sites with at least 25 data-points fulfilling the requirements noted above. A list of these 110 sites used for the parameter estimation can be found in the [supplement \(Section S4, Table S2\)](#).

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The presented analyses presume that the observed evapotranspiration is dominated by transpiration after three consecutive rain-free days. To test the robustness of our findings against this assumption, we varied the number of consecutive [rain-free](#) days from 0 to 14. For each considered step, the data were filtered accordingly and parameters of the WUE models were estimated for each site. We then calculated the mean of the parameter estimates over all sites and the associated uncertainty (95% confidence interval) of the mean via bootstrapping. We excluded some humid sites from this sensitivity analysis for which too few data points were available when filtering for longer rain-free periods. This procedure ensures that [all levels of filtering for rain-free periods included the same set of sites](#).

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2.2 Concepts & models

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For our analysis, we started out with the WUE model of Zhou et al. (2015), which we converted for an inversion against ET data:

$$ET = \frac{GPP \cdot VPD^{0.5}}{uWUE} , \quad (1)$$

20

where [uWUE](#) represents the site-specific *underlying water-use efficiency*. The introduced WUE model states that no transpiration occurs independently from $GPP \cdot VPD^{0.5}$. To test this hypothesis, we introduced an intercept term (ET_{res}) in the generic WUE model, which we call $+ET_{res}$ model:

$$25 \quad ET = \frac{GPP \cdot VPD^{0.5}}{uWUE} + ET_{res} . \quad (2)$$

Hence, any significant intercept would indicate transpiration that cannot be explained by $GPP \cdot VPD^{0.5}$. The relative contribution of ET_{res} to the mean predicted flux, C_{ETres} , was calculated as:

$$30 \quad C_{ETres} = \frac{ET_{res}}{\frac{GPP \cdot VPD^{0.5}}{uWUE} + ET_{res}} , \quad (3)$$

where the denominator contains the mean predicted daily ET of the model.

To further test whether ET_{res} relates to atmospheric variables, we postulated three different alternative models. The residual transpiration could be driven by an additional VPD term that is independent from photosynthesis, as some stomatal conductance models (Medlyn et al., 2011) already include a residual conductance term g_0 , in which case the WUE model could take the form:

$$ET = \frac{GPP \cdot VPD^{0.5}}{uWUE} + g_0 \cdot VPD. \quad (4)$$

We also considered the possibility that ET_{res} is related to global radiation (Rg):

$$ET = \frac{GPP \cdot VPD^{0.5}}{uWUE} + r \cdot Rg. \quad (5)$$

To test for possible interactions between the two additional variables, we considered a third alternative where ET_{res} was modelled by included both Rg and VPD as independent factors:

$$ET = \frac{GPP \cdot VPD^{0.5}}{uWUE} + g_0 \cdot VPD + r \cdot Rg. \quad (6)$$

In the following, we refer to the reference WUE definition (Eq. 1) as "Zhou". We abbreviate models with additional covariates by omitting the reference to the GPP-VPD-term ($\frac{GPP \cdot VPD^{0.5}}{uWUE}$) of Zhou, which is always used unless denoted otherwise. The model with an additional VPD term, for example, is thus designated "+VPD".

2.3 Parameter estimation & statistics

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In the following, we refer to Eq. 1, 2, 4, 5, 6 as models, as we optimize their fit to the EC data by estimating free parameters. The estimation was conducted with the Levenberg-Marquardt technique, minimizing the sum of squares of the model residuals. The residuals were calculated as the difference between observed and predicted ET. We used the nlsLM package in R (Elzhov et al., 2015). All parameters were restricted to positive values, preventing biologically implausible estimates.

The uncertainties and correlations of the parameters were calculated with the variance-covariance matrix returned by the fitting function (Omlin & Reichert, 1999).

The model performance was assessed for each site with the Nash-Sutcliffe model efficiency (MEF):

$$\text{MEF} = 1 - \frac{\sum(Y_{\text{prd}} - Y_{\text{obs}})^2}{\sum(Y_{\text{obs}} - \bar{Y}_{\text{obs}})^2}, \quad (7)$$

5

where Y_{obs} are observations and Y_{prd} are predictions by a model.

A MEF of 1 implies a perfect fit of the model to the data. A MEF below 0 implies that the mean of the observations outperforms the fit of the model. All MEFs were calculated in a *leave-one-out* cross-validation to account for the problem of over-fitting. Thus, the cross-validated MEFs can be used to compare models with differing numbers of free parameters.

We assessed parameter distributions stratified according to a classification of vegetation structure that was based on plant functional types (PFT). The included FLUXNET sites were classified as *low vegetation structure* if the report PFT was either grassland (GRA) or crops (CRO) and *high vegetation structure* if otherwise. In total, 40 sites had *low vegetation structure*, compared to 70 sites with *high vegetation structure*. The significance of differences between distributions of parameters and metrics for these two classes was verified by a Kolmogorov–Smirnov test (Daniel, 1990) and bootstrapped 95% confidence intervals of the mean.

15

2.4 Contribution Analysis

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To assess the contribution of driving variables to the predicted fluxes, we performed an analysis of attribution to the individual model terms. Consider a simple multiple linear regression model,

$$Y = a_1 \cdot X_1 + a_2 \cdot X_2, \quad (8)$$

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where Y is the dependent variable, X_1 and X_2 are independent variables and a_1 and a_2 denote the model parameters. Due to the additive character of the model, the contribution of one variable (e.g. X_1) to the total flux is given by its product with the slope ($a \cdot X_1$). In our analysis, we tested the contribution of the linear radiation term (Eq. 5) to the total modelled evapotranspiration. Thus we defined the fraction of evapotranspiration that was attributed to the radiation term as:

30

$$\text{ET}_{\text{frac}} = \frac{\sum r \cdot R_g}{\sum \left(\frac{\text{GPP} \cdot \sqrt{\text{VPD}}}{\text{uWUE}} + r \cdot R_g \right)} \quad (9)$$

We considered two variants to estimate this metric: *parallel* and *hierarchical*. In the first case, both parameters, uWUE and r were estimated in a standard parameter estimation, i.e. concurrently. In the second case, uWUE was first estimated in the +ET_{res} model. We then defined a term M as:

5

$$M = \text{ET}_{\text{res}} + \varepsilon, \quad (10)$$

where ε denotes the residuals and ET_{res} denotes the intercept parameter of the +ET_{res} model. The parameter r was then estimated in a linear regression of the form:

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$$M = r \cdot \text{Rg} + c, \quad (11)$$

where c denotes a constant intercept. By giving precedence to the uWUE parameter in this approach, we expect to get an estimate of a reasonable lower bound for ET_{frac}. All parameters were constrained to positive values.

15

To further assess uncertainties due to problems of parameter identifiability among uWUE and r in the *parallel* variant, we sampled 200 parameter vectors from the posterior parameter uncertainty distribution for each site.

Impact of parameter correlations on the contribution analysis

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In many realistic examples, the model parameters of Eq. 8, a_1 and a_2 , are not perfectly identifiable. This could be due to the correlation of X_1 and X_2 or more fundamental model structural uncertainty. In these cases, f_{X_i} is confounded by the parameter correlation a_1 and a_2 , leading to a high uncertainty of its estimation. In our analysis, GPP and Rg are expected to be highly correlated, leading to dependent parameter uncertainties. To evaluate this effect, we estimated the contribution of the described within-site uncertainty of f_{X_i} . The variance-covariance matrix (V) of the parameter estimates could be

25 calculated for each site with the results of the regression. Consequently, V can be used to derive the respective posterior parameter distributions, from which we sampled 200 parameter vectors per site, representing the uncertainty and correlation of the two parameters. Site-specific vectors $f_{X_i,S}$ can be calculated as

25

$$f_{X_i,S_j} = \left[f_{X_i,S_j,p_1}, f_{X_i,S_j,p_2}, \dots, f_{X_i,S_j,p_{200}} \right], \quad (12)$$

30

where p_1 to p_{200} denote the 200 realizations of parameter vectors and s_j denotes a specific site.

Based on this, we tested whether the global variance of f_{X_i} is a product of differing means of the site-specific $f_{X_i,S}$ vectors or their variances. For this, we performed an ANOVA along the sites as categorical variable with the parameter realizations as random replicates.

5

3 Results

In our analysis we tested different water-use efficiency (WUE) models that predicted evapotranspiration (ET) using the product of gross-primary productivity (GPP) and the water vapor pressure deficit, $GPP \cdot VPD^{0.5}$, as predictor variable. When plotting ET as a function of [this multiplicative term](#), we observed significant intercepts, e.g. for the Mediterranean FLUXNET site IT-BCi (Fig. 1a). In these cases, significant ET was observed when the driving force of the established models, the GPP-VPD-product, was small or zero. When we explicitly included this term in the model (+ETres), the cross-validated MEF increased notably (Fig. 1b). As Table 1 shows, the +ETres variant outperformed the Zhou model at 86% of the sites. The respective mean difference in MEF between the two variants was 0.07 (Table 2). [A corresponding table that presents the mean difference of the root mean squared error \(RMSE\) for all sites can be found in the supplementary materials \(Table S1\).](#)

20 **Table 1. Fraction of the 110 sites at which the model of the respective row was superior to the model of the respective column according to the MEF_{diff} criterion.**

	Zhou	+ETres	+VPD	+Rg	+VPD+Rg
Zhou	-	0.14	0.22	0.07	0.07
+ETres	0.86	-	0.43	0.07	0.08
+VPD	0.78	0.57	-	0.08	0.08
+Rg	0.93	0.93	0.92	-	0.71
+VPD+Rg	0.93	0.92	0.92	0.29	-

25 **Table 2. Mean MEF_{diff} comparison for different model variants. Entries indicate the mean of the difference MEF(row model) – MEF(column model) for all 110 sites.**

	Zhou	+ETres	+VPD	+Rg	+VPD+Rg
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Zhou	-	-0.07	-0.09	-0.16	-0.17
+ETres	0.07	-	-0.02	-0.09	-0.10
+VPD	0.09	0.02	-	-0.07	-0.07
+Rg	0.16	0.09	0.07	-	0.00
+VPD+Rg	0.17	0.10	0.07	0.00	-

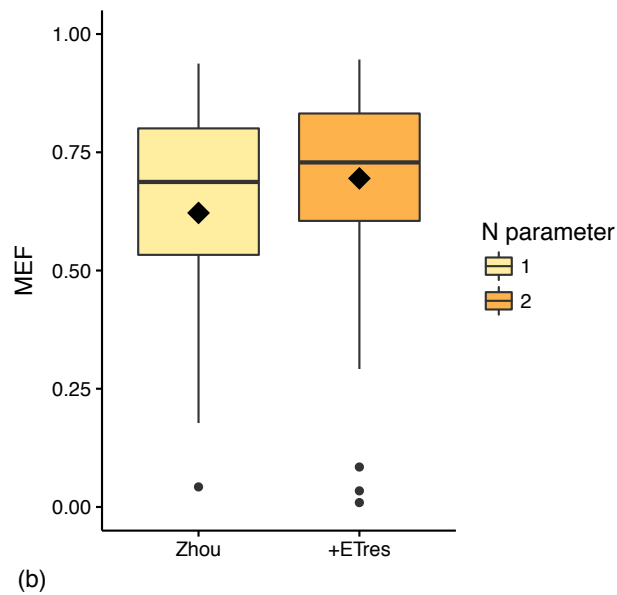
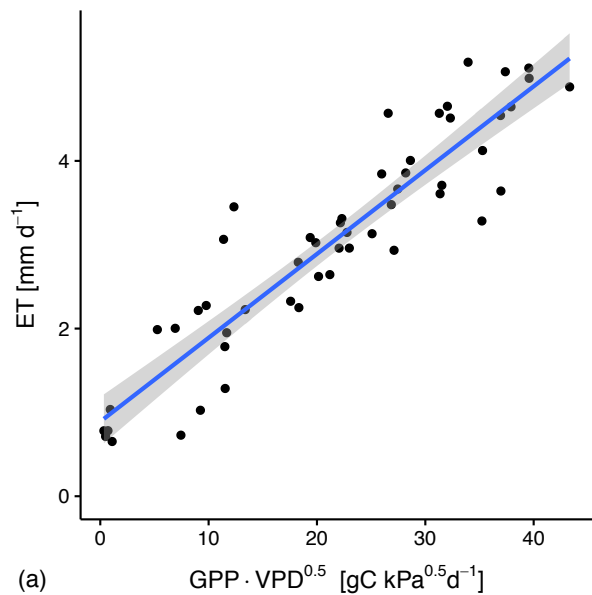


Figure 1. Linear regression between the product of GPP and $VPD^{0.5}$ and the daily ET; the shaded area indicates the 95% confidence interval of the mean of the predictions (a). Model performance of the Zhou model compared to the +ET_{res} variant (b). The thick horizontal lines of the boxplots denote the median, the diamonds denote the mean of the MEF of all sites.

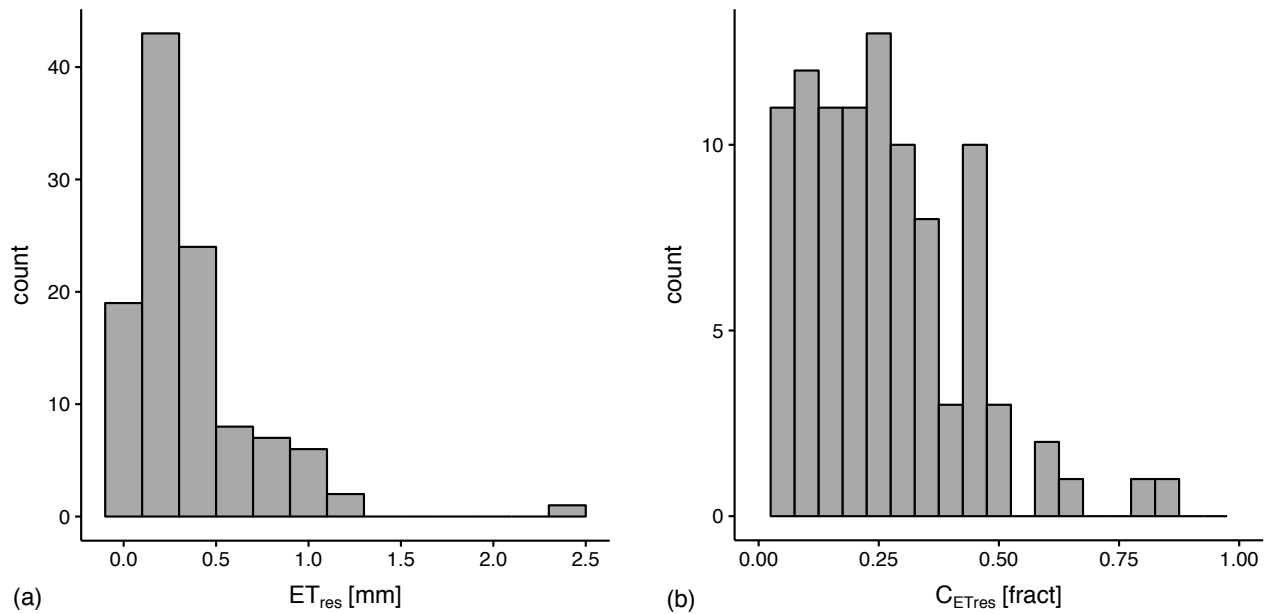


Figure 2. Distribution of the estimated intercepts. The majority of sites had values between 0–1.2 mm for the absolute intercept (a). The relative contribution of ET_{res} to the daily mean evapotranspiration as quantified by C_{ETres} reached up to 0.94 (b).

5 Of the 110 sites included in the analysis, 78% had a significant intercept. The site specifically estimated intercept values ranged from 0 to 2.36 mm (90%-percentile 0.86 mm) with a mean of 0.36 mm (Fig. 2a). The relative intercept C_{ETres} reflects the relative magnitude of the intercept to the mean daily ET of the respective sites. It ranged from 0 to 0.86 (90%-percentile 0.46) with a mean of 0.23. This implies that circa a quarter of transpiration was not attributed to $GPP \cdot VPD^{0.5}$ in this model formulation (Fig. 2b). The importance of the intercept for the prediction of daily ET and its diverging values raise the question whether the intercept compensates for the absence of a physical or biological process in the model or whether the observations are confounded by a systematic problem.

Our first hypothesis was that the ET_{res} intercept was due to the remaining contributions of soil and interception evaporation to measured ET after three consecutive rain-free days. To test this, we estimated the parameter ET_{res} for periods of successively longer consecutive rain-free days. If our hypothesis was right, we would expect a trend of declining ET_{res} with increasing consecutive rain-free days. However, no reduction of ET_{res} beyond the exclusion of the three days after precipitation proposed by Beer et al. (dotted line) could be observed (Fig. 3a). We therefore concluded that potential contributions of soil and interception evaporation to ET cannot explain the existence of the intercept term in the WUE model.

We then hypothesized that a missing process in the model would be discernible as a temporal pattern in the ET_{res} estimates. For that, we estimated the intercept for each month and site separately. The monthly means of the intercept for all sites varied in a clear seasonal pattern (Fig. 3b).³

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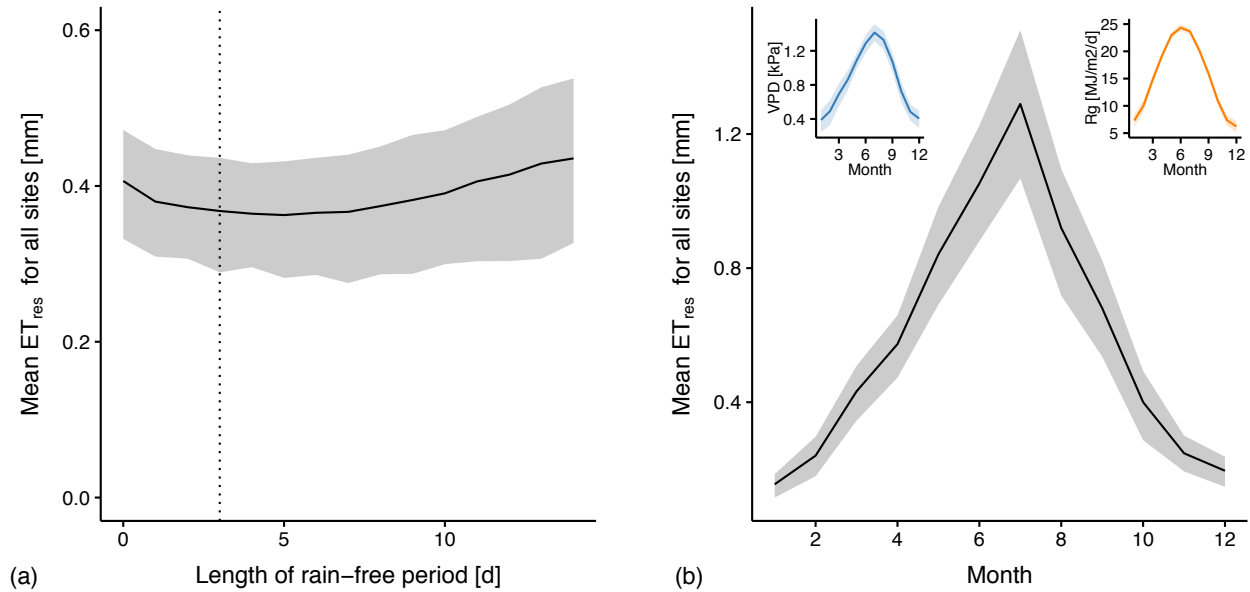


Figure 3. Sensitivity of ET_{res} to varying length of filtering after precipitation events (a). The dotted, vertical line indicates the three-day period we adopted from Beer *et al.* Monthly estimation of the intercept parameter ET_{res} . The inset plots illustrate the mean seasonal variability of VPD and Rg for all sites on a monthly scale (b). In all plots, the solid line is the mean for all sites; the band denotes the 95% confidence interval of the mean derived by bootstrapping.

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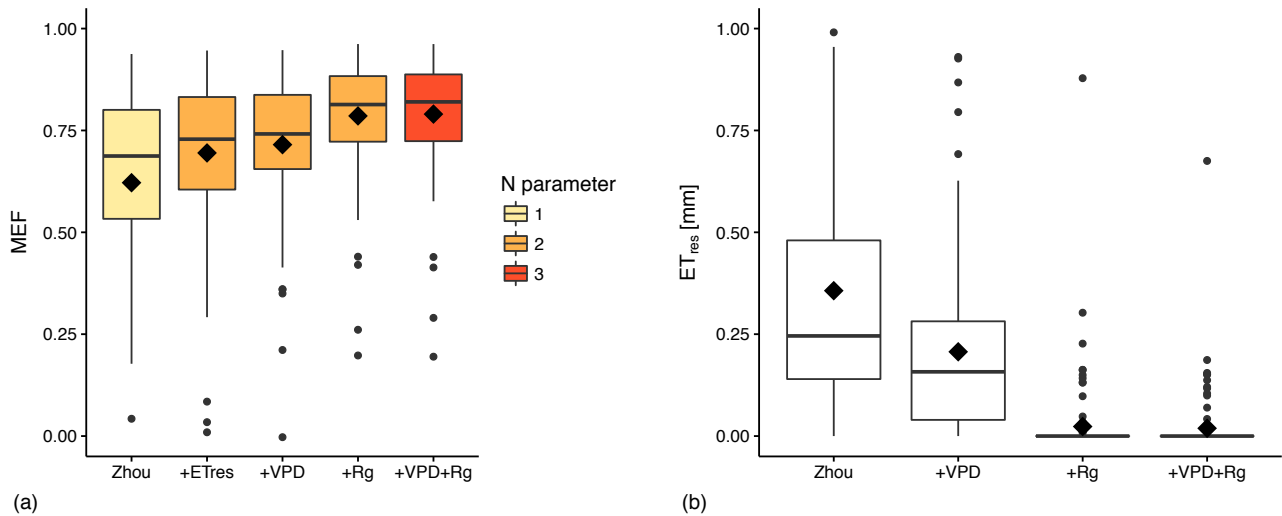
The seasonality of ET_{res} suggests a relationship to meteorological variables such as VPD or radiation that vary seasonally too. A relationship with VPD could represent the g_0 term of canopy conductance models (Ball *et al.*, 1987). It was therefore introduced in the form of an additional linear term (Eq. 4). A relationship with global radiation (Rg) (Eq. 5) could represent

15 *equilibrium transpiration* (Jarvis and McNaughton, 1986), where the energy surplus of incoming radiation forces a transpirational flux independent from the vapor pressure gradient. We tested modeling the ET_{res} intercept by including both variables separately (Eq. 4, 5) and jointly (Eq. 6).

³ The monthly variability of ET_{res} for all sites individually can be found in the supplementary materials (Section S2, Fig. S1).

We found only a small performance increase of the +VPD variant with regard to the Zhou model with intercept term (“+ET_{res}”, Fig. 4a). In fact, the MEF_{diff} suggested that the +ET_{res} model was still superior to the +VPD variant at 43% of the sites (Table 1). By comparison, the +Rg model showed a substantial mean increase of 0.09 in the MEF compared to the +ET_{res} model (Table 2). The increased model complexity (two free parameters) was justified at 93% of the sites. The model variant with both VPD and Rg terms (“+VPD+Rg”) did not improve predictions compared to the simpler +Rg variant. Plots showing the impact of the +ET_{res} and +Rg variant on the accuracy of the ET predictions for two selected sites can be found in the supplementary materials (Section S3, Fig. S2,S3)

We also tested whether the variation of ET that was unexplained by the +Rg model was still correlated with radiation. This would indicate that the chosen linear radiation-term did not fully account for all covariation between ET and Rg, implying a nonlinear dependency. The mean R^2 between the residuals of the +Rg variant and observed ET was 3.9% (maximum: 22%, 90%-percentile: 10%). This suggests that the linear variant was an adequate choice, as little of the unexplained variation of ET was still correlated with radiation.



15 **Figure 4. Cross-validated MEFs of the model variants with additional covariates (a). The +Rg variant lead to a further increase of model performance when comparing to the +ET_{res} variant. By contrast, the +VPD variant showed only a small increase of performance. Global distribution of the residual intercept for all model variants (b).**

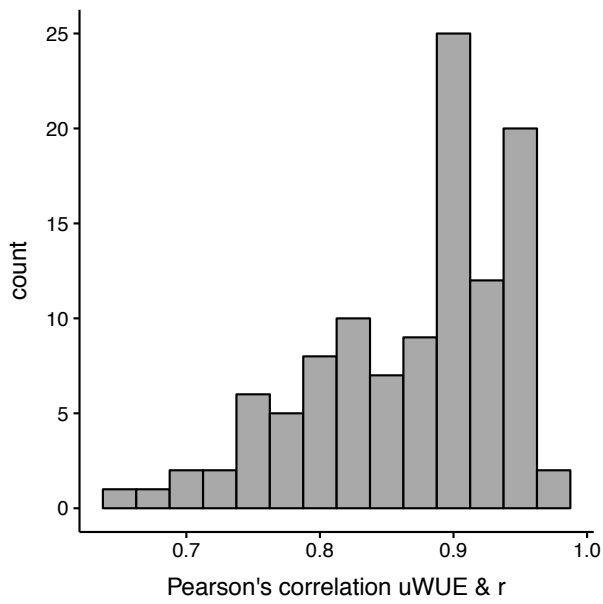


Figure 5. Correlation of the parameter estimates for r and uWUE across sites.

The difference between the +Rg and +VPD model is equally distinct, when the presence of a remaining model bias was tested. For this, the intercept was estimated with all four model variants (Zhou, +VPD, +Rg, +VPD+Rg). Only the models with an Rg-term had considerably reduced residual intercepts (Fig. 4b).

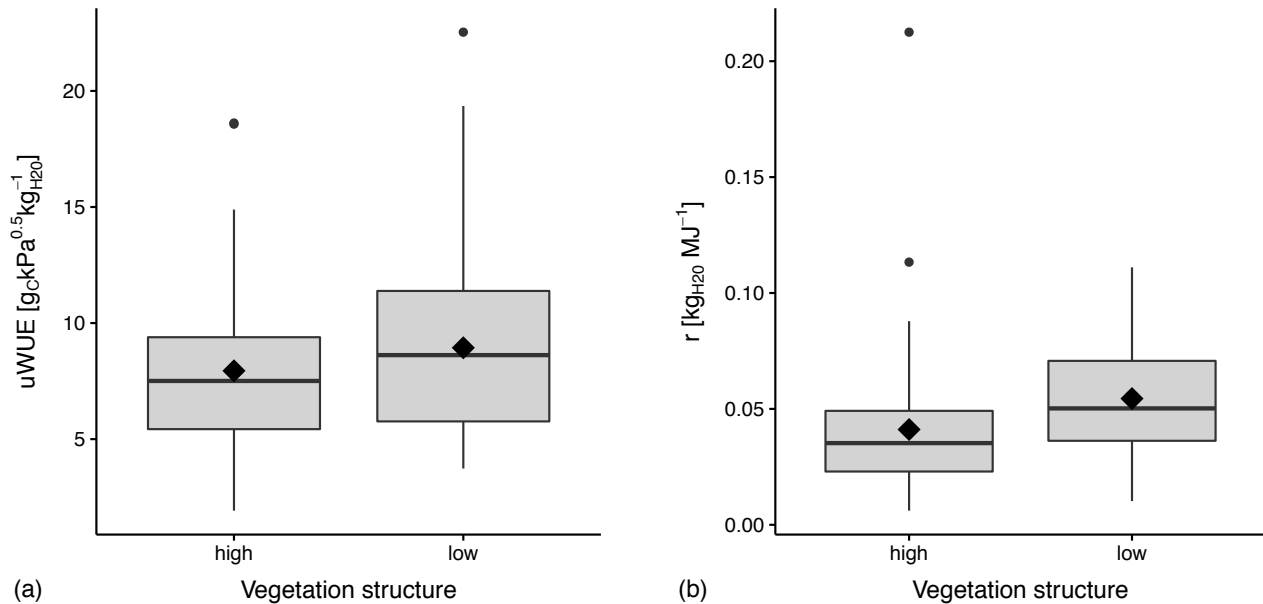
All models compared to the original definition Zhou had two or more parameters. Of those models with two parameters, the +Rg emerged as the best model after cross-validation, indicating that the additional model complexity was in fact justified. The presence of two parameters raised the question whether they could be identified independently. In fact, we found a high degree of correlation between the parameters for all sites (Fig. 5). It is likely that the correlation originates from the correlation between GPP and Rg.

Parameter distributions were separated for low and high vegetation structure (Fig. 6). The estimates of uWUE (a) were not significantly different for low and high vegetation (Kolmogorov–Smirnov test, $p = 0.35$). By contrast, the estimates for r (b) were significantly higher for low (mean: 0.054, 95%-CI: 0.047–0.062) compared to high vegetation structure (mean: 0.041, 95%-CI: 0.034–0.049; Kolmogorov–Smirnov test, $p = 0.003$).

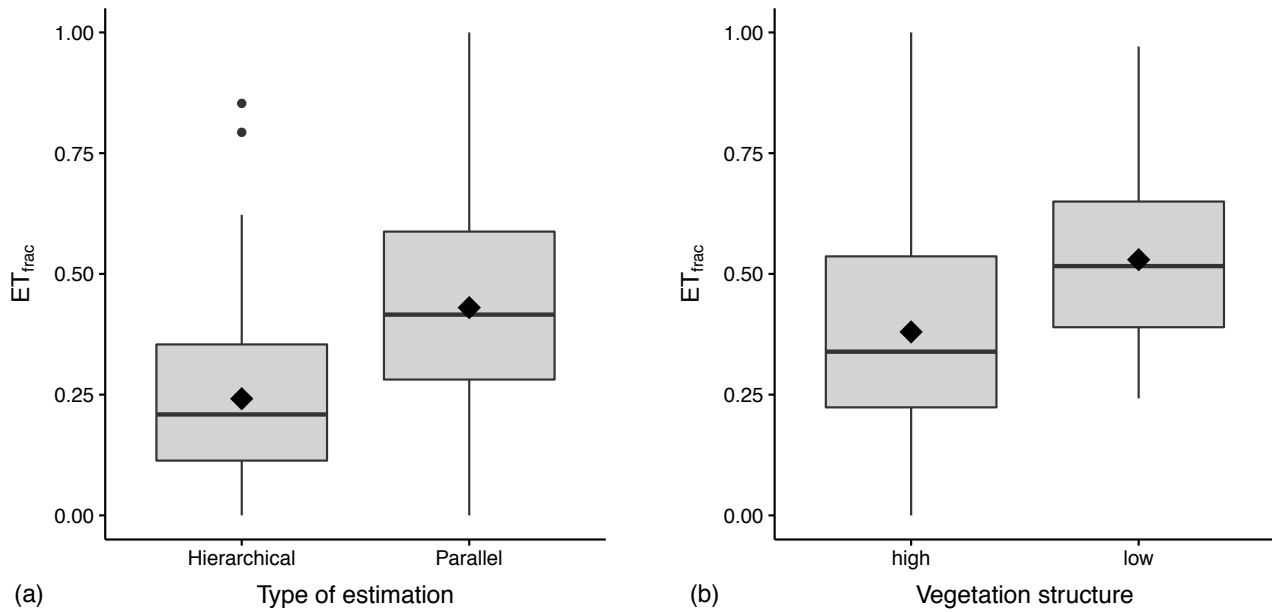
As transpiration of the +Rg model is a linear combination of stomatal and radiation-driven components, it is possible to calculate the relative contribution of each component to daily ET fluxes. We compared two approaches to calculate this

quantity: *Parallel* for estimating both parameters concurrently, *hierarchical* for estimating r only after uWUE has been calibrated (Eq. 10, 11). In both approaches, we observed that a sizeable fraction of mean daily ET could be attributed to the radiation term (Fig. 7a). For the *hierarchical* approach, the mean global ET_{frac} was 24% (95%-CI: 21–27%); for the *parallel* approach, the mean global ET_{frac} was 43% (95%-CI: 39–47%). Similar to the assessment of parameter distributions, we stratified the ET_{frac} index for low and high vegetation structures (Fig. 7b). ET_{frac} was significantly higher for low (mean: 0.53; 95%-CI: 0.48–0.58) compared to high vegetation structure (mean: 0.38; 95%-CI: 0.33–0.43; Kolmogorov–Smirnov test $p < 0.001$).

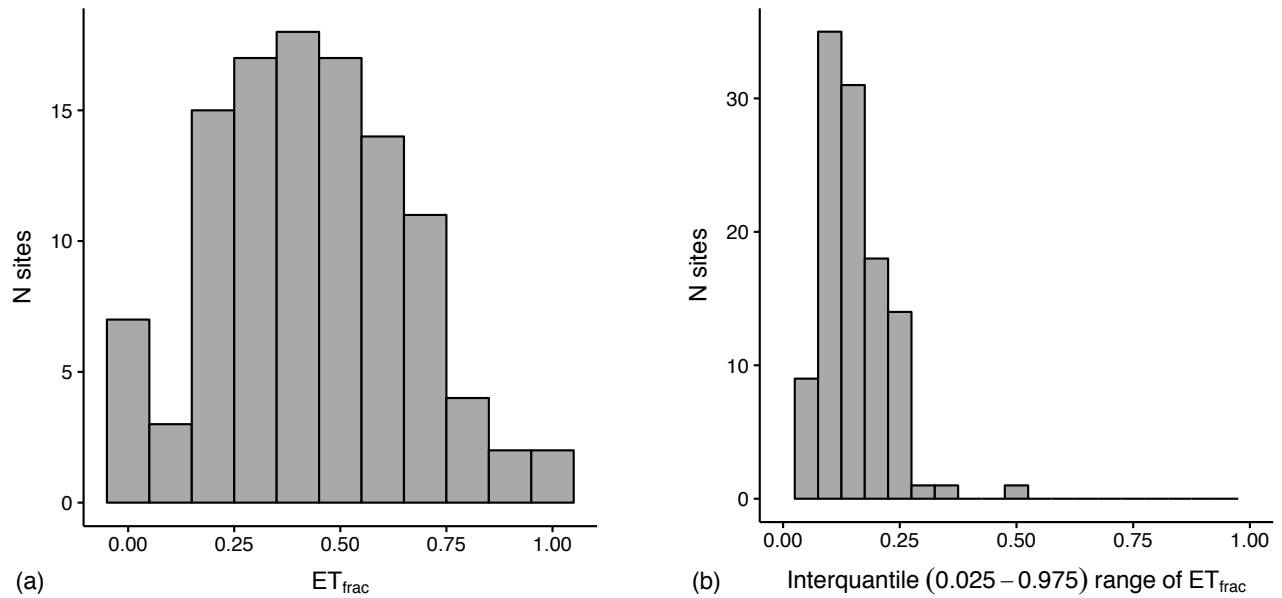
Notably, ET_{frac} estimated with the *parallel* approach varied widely between the sites (Fig. 8a). We assessed to which degree this variability can be interpreted as between-site variability of the expected value of ET_{frac} or whether it is due to poorly constrained and correlated parameters *due to collinearity* (Fig. 5). The ‘within-site’ uncertainty of ET_{frac} caused by parameter uncertainty was quantified as the range between the 97.5 and the 2.5 percentiles (95% confidence interval, CI) of ET_{frac} estimates from 200 parameter vectors sampled from their respective posterior distributions. The vast majority of sites had an CI lower than 0.3 (Fig. 8b). This suggests that the large variability of ET_{frac} was not a result of parameter uncertainties. The conducted ANOVA supported this conclusion, as it revealed that 96% of the global ET_{frac} variability could be attributed to the variability between sites.



20 **Fig. 6. Distribution of the parameter estimates partitioned by type of vegetation structure. For uWUE (a), a Kolmogorov–Smirnov test yielded no difference in the distribution between the two classes of vegetation structure ($p = 0.35$). For r (b), the same test indicated a difference in the distribution between the two classes of vegetation structure ($p = 0.003$).**



5 **Fig. 7. Distributions of ET_{frac} estimates for all sites. The diamonds indicate the mean, the bold horizontal lines indicate the median. The hierarchical approach yielded substantially lower values and can be interpreted as a conservative estimate of the quantity (a). ET_{frac} was significantly higher in sites with low vegetation structure (b).**



5 **Figure 8ab.** Distribution of the fraction of mean daily transpiration (ET_{frac}) attributed to the radiation term for all sites, calculated with the *parallel* approach (a). Distribution of the within-site variability of ET_{frac} calculated as the difference between the 97.5 and 2.5 percentiles of the ET_{frac} estimates derived by sampling from the posterior parameter densities, parallel approach (b).

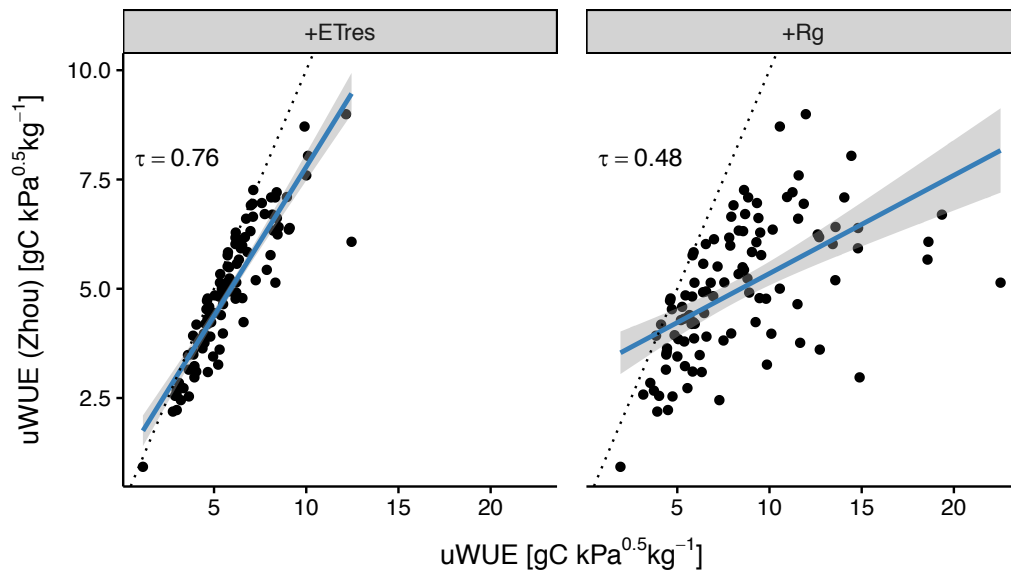


Figure 9. Effect of different model variants on global uWUE estimates. The 1:1 line is dotted. A linear dependency would indicate that uWUE estimates are commensurable across sites, even if only after a linear scaling. When comparing with the estimates of the original Zhou model, the correlation is notably weaker for +Rg than for +ET_{res}. Sites with high parameter uncertainties were removed for all models.

5

Finally, we assessed the impact of ET_{res} and Rg as additional covariates on the global variability of uWUE estimates (Fig. 9). We calculated Kendall's τ rank correlation coefficient (Kendall, 1938) between the site-level estimates of uWUEs derived from the Zhou model and two different variants: +ET_{res} and +Rg. The degree of correlation of the uWUE estimates quantifies whether changes in the model structure permutes the ordering of the estimates between sites. The extent to which parameter estimates are affected by the model structure is crucial because any explanation or prediction of parameter values between sites would be highly desirable. While a moderate correlation between the uWUE estimates of the Zhou model with the +ET_{res} variant can be seen ($\tau = 0.76$), the correlation of the uWUE estimates of the Zhou model with the respective values of the +Rg variant is low ($\tau = 0.48$). The between-site variability of uWUE explained by the estimates of the Zhou variant was 79% for the +ET_{res} variant and 31% for the +Rg variant, as quantified by the R^2 .

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4 Discussion

4.1 Findings and mechanisms

20 In this study, we identified radiation as an important variable for ecosystem-scale transpiration and water-use efficiency. Depending on the approach used, we attributed between a quarter and half of mean daily transpiration of all included FLUXNET sites to a linear radiation term. These findings raise the question which biophysical or ecophysiological processes can account for the estimated magnitudes of this attributed fraction.

25 The influence of radiation on stomatal conductance has been noted and discussed in the literature (Whitehead et al., 1981; Jarvis and McNaughton, 1986) and is also reflected in existing transpiration models (Leuning et al., 2008). By contrast, we detected a substantial transpiration component that was statistically independent from the product of GPP and VPD^{0.5}, tentatively suggesting an insensitivity to stomatal conductance. Our results suggest that this additional flux could not be associated with a g_0 conductance term, as this would imply the dependency of the additional ET on a linear VPD term (Ball et al., 1987). By contrast, the intercept was shown to be more consistent with radiation. Consequently, models that integrated radiation-driven transpiration with such an additive linear response had a superior predictive performance across flux towers.

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The observed effect of radiation indicates that equilibrium transpiration could play an important role in ecosystem-scale transpiration. Equilibrium transpiration (or equilibrium evaporation rate) is the transpiration occurring if the leaf is completely decoupled from the atmosphere (Jarvis and McNaughton, 1986). Therefore, equilibrium transpiration is independent from stomatal conductance and driven solely by the dissipation of energy, provided by incoming solar radiation.

5 According to McNaughton and Jarvis (1983), the decoupling parameter Ω is notably higher in grasslands compared to forests. This parameter reflects the proportion of evaporation that is independent from the driving gradient in the water vapour pressure. Our observation that both the radiation parameter r and the index ET_{frac} were significantly higher for ecosystems with low vegetation structure is therefore consistent with the explanation that equilibrium evaporation is responsible for the observed role of radiation.

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This can be contrasted with a competing explanation that the radiation dependency of transpiration does not reflect an additional process but rather a systematic problem of the VPD observations. VPD is measured together with the fluxes above the canopy. While the recorded water and carbon fluxes do in fact represent the net fluxes of the tower footprint, the same cannot be said for VPD. Its measurement above the canopy may differ substantially from the relevant magnitude of the variable at the leaf-scale. However, leaf temperature and thus the VPD of the leaf boundary layer is dependent on solar radiation (Tenhunen et al., 1990). Adding solar radiation to the equation could therefore be seen as compensating for the lack of the aforementioned leaf-scale VPD observations.

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4.2 Limitations

One limitation stems from the selection of rain-free periods for the parameter estimation. As previously described, this is a necessary step to justifiably assume that the observed latent heat fluxes constitute mostly a transpiration flux, rather than evaporation from bare-soil and leaf surfaces in addition to transpiration. It also makes our work comparable with the study of Beer et al. (2009) that used this method to derive their estimates. For the observed residual evapotranspiration ET_{res} we could show that it is not an artefact of insufficient exclusion of days after precipitation events. However, the environmental conditions during and after rain events generally represent some of these specific conditions: Low VPD due to the moisture available for evaporation, a higher share of diffuse radiation and a more or less sudden increase in soil moisture among others. All mentioned variables could plausibly be assumed to have an influence on the stomatal opening of plants. Therefore, the presented WUE model must not necessarily predict flux relationships during or immediately after precipitation events due to the underrepresentation of similar conditions in the sample of observations used for parameter estimation.

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The effect of atmospheric CO₂ concentrations on WUE (Morison, 1985; Conley et al., 2001) was not considered in this analysis. One could suspect that the seasonal variability of ET_{res} is affected by seasonal CO₂ variability if both were in phase. However, ET_{res} showed a global maximum during June–July, while northern hemisphere CO₂ concentrations are at its

minimum in September–October (Keeling et al., 1976), implying that CO₂ concentrations are unlikely to cause the seasonal variation of ET_{res}.

- 5 The presented results are subject to the eco-climatological representativeness of the FLUXNET in general and sites with a *fair-and-free-use* data policy specifically. This means that tropical areas in particular are underrepresented in our analysis (with only one Australian site, AU-How). However, all posited mechanisms that could be responsible for the observed effect of radiation are not expected to be restricted to extra-tropical regions. Further research is however required to quantify the importance of radiation for water-use efficiency and transpiration in tropical ecosystems.
- 10 The model structures we tested in this study were evaluated according to their global empirical adequacy. Thus, despite the possible identification of probable mechanisms responsible for the observed patterns, the model structure selected in the end likely does not reflect the exact physical mechanisms by which the ecosystem operates. Furthermore, this limitation can be specifically important in the case of water limitation. As the R_g-term is completely independent from any variable reflecting vegetation activity, our model would predict transpiration scaling with radiation during periods of severe drought. By
- 15 principle, the same problem could affect periods of low temperatures before leaf flushing, although these periods are generally also associated with low radiation levels and hence may not be as problematic.

4.3 Implications

- 20 Our empirical analysis suggests that ecosystem-scale transpiration depends to a sizeable degree on *radiation rather than only the product GPP and VPD^{0.5}*. This implies that photosynthesis and transpiration might be less strongly coupled on ecosystem scale than commonly assumed. We speculate that the additional effect of radiation could be due to equilibrium transpiration where radiation drives transpiration even when the canopy is fully decoupled from the atmosphere. *The provided evidence for the differences between low and high vegetation structures gives additional credence to this explanation.* However, we
- 25 cannot disentangle direct physiological effects of radiation on transpiration from other radiation effects that are relevant on ecosystem scale at this point. For the latter, leaf to canopy scaling, micrometeorological conditions, and boundary layer dynamics might contribute to the observed relationship between ecosystem scale water use efficiency and radiation. Thus, further research is needed to reconcile our empirical findings with detailed ecosystem-scale modeling and theory on the one hand and plant-physiological research under controlled conditions on the other hand.

30

Finally, we caution against prematurely interpreting the between-site variability of underlying water-use efficiency (uWUE). We showed that estimates of uWUE derived with the Zhou model explained only a third of the observed variability of

ecosystem uWUE derived from our empirically superior model formulation. The dependence of uWUE estimates on the chosen model formulation makes the interpretation of uWUE as an ecosystem property problematic. This concern would also hold for assessing temporal dynamics of uWUE such as long-term trends. For example, the unexpectedly large global trend of WUE across FLUXNET sites (Keenan et al., 2013) would need to be tested for its omission of radiation in the model that was used. Overall, this study highlights the importance of model structure uncertainty for interpretations of parameter variability.

References

- Baldocchi, D., Falge, E., Gu, L., Olson, R., Hollinger, D., Running, S., Anthoni, P., Bernhofer, C., Davis, K., Evans, R., Fuentes, J., Goldstein, A., Zhou, G., Law, B., Lee, X., Malhi, Y., Meyers, T., Munger, W., Oechel, W., Paw U, K. T., Pilegaard, K., Schmid, H., Valentini, R., Verma, S., Vesala, T., Wilson, K., and Wofsy, S.: FLUXNET: A new tool to study the temporal and spatial variability of ecosystem-scale carbon dioxide, water vapor, and energy flux densities, *Bulletin of the American Meteorological Society*, 82, 2415–2434, 2001.
- Ball, J. T., Woodrow, I. E. and Berry, J. A.: A Model Predicting Stomatal Conductance and its Contribution to the Control of Photosynthesis under Different Environmental Conditions, in: *Progress in Photosynthesis Research* (Eds. Biggins, J.), Springer Netherlands, Dordrecht, 221–224, 1987.
- Beer, C., Ciais, P., Reichstein, M., Baldocchi, D., Law, B. E., Papale, D., Soussana, J.-F., Ammann, C., Buchmann, N., Frank, D., Gianella, D., Janssens, I. A., Knohl, A., Köstner, B., Moors, E., Rouspard, O., Verbeeck, H., Vesala, T., Williams, C. A., and Wohlfahrt, G.: Temporal and among-site variability of inherent water use efficiency at the ecosystem level, *Global Biogeochemical Cycles*, 23, GB2018, 2009.
- Churkina, G., Running, S. W., and Schloss, A. L.: Comparing global models of terrestrial net primary productivity (NPP): the importance of water availability, *Global Change Biology*, 5, 46–55, 1999.
- Ciais, P., Sabine, C., Bala, G., Bopp, L., Brovkin, V., Canadell, J., Chhabra, A., DeFries, R., Galloway, J., Heimann, M., et al.: Carbon and Other Biogeochemical Cycles, in: *Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change* (Eds. Stocker T. F., Qin D., Plattner G. K., Tignor M., Allen S. K., Boschung J., Nauels A., Xia Y., Bex V., Midgley P. M.), Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 465–570, 2014.
- Conley, M. M., Kimball, B., Brooks, T., Pinter, P., Hunsaker, D., Wall, G., Adam, N., LaMorte, R., Matthias, A., Thompson, T., et al.: CO₂ enrichment increases water-use efficiency in sorghum, *New Phytologist*, 151, 407–412, 2001.
- Cowan, I. R. and Farquhar, G. D.: Stomatal function in relation to leaf metabolism and environment, in: *Integration of Activity in the Higher Plant*, edited by Jennings, D. H., pp. 471–505, Cambridge University Press, 1977.

[Daniel, W. W.: Applied Nonparametric Statistics. PWS-Kent, Boston, 1990.](#)

de Kauwe, M.G., Medlyn, B.E., Zaehle, S., Walker, A.P., Dietze, M.C., Hickler, T., Jain, A.K., Luo, Y., Parton, W.J., Prentice, I.C. and Smith, B. Forest water use and water use efficiency at elevated CO₂: a model-data intercomparison at two contrasting temperate forest FACE sites. *Global Change Biology*, 19, 1759–1779, 2013.

5

Elzhov, T.V., Mullen, K.M., Spiess, A.N., and Bolker B.: minpack.lm: R Interface to the Levenberg-Marquardt Nonlinear Least-Squares Algorithm Found in MINPACK, Plus Support for Bounds. R package version 1.2-0. 2015

Farquhar, G.D., von Caemmerer, S. and Berry, J.A.: A biochemical model of photosynthetic CO₂ assimilation in leaves of C₃ species, *Planta*, 149, 78–90, 1980.

10

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, 2004.

15 Ito, A. and Inatomi, M.: Water-use efficiency of the terrestrial biosphere: a model analysis focusing on interactions between the global carbon and water cycles, *Journal of Hydrometeorology*, 13, 681–694, 2012.

Jarvis, P. G. and McNaughton, K.: Stomatal control of transpiration: scaling up from leaf to region, *Advances in ecological research*, 15, 49, 1986.

20

Jasechko, S., Sharp, Z. D., Gibson, J. J., Birks, S. J., Yi, Y., and Fawcett, P. J.: Terrestrial water fluxes dominated by transpiration, *Nature*, 496, 347–351, 2013.

25 Katul, G., Manzoni, S., Palmroth, S., and Oren, R.: A stomatal optimization theory to describe the effects of atmospheric CO₂ on leaf photosynthesis and transpiration, *Annals of Botany*, p. mcp292, 2009.

Keeling, C. D., Bacastow, R. B., Bainbridge, A. E., Ekdahl, C. A., Guenther, P. R., Waterman, L. S., and Chin, J. F.: Atmospheric carbon dioxide variations at Mauna Loa observatory, Hawaii, *Tellus*, 28, 538–551, 1976.

30 Keenan, T. F., Davidson, E., Moffat, A. M., Munger, W., and Richardson, A. D.: Using model–data fusion to interpret past trends, and quantify uncertainties in future predictions of terrestrial ecosystem carbon cycling, *Global Change Biology*, 18, 2555–2569, 2012.

Keenan, T. F., Hollinger, D. Y., Bohrer, G., Dragoni, D., Munger, J. W., Schmid, H. P. and Richardson, A. D.: Increase in forest water-use efficiency as atmospheric carbon dioxide concentrations rise, *Nature*, 499, 324–327, 2013.

Kendall, M.: A New Measure of Rank Correlation, *Biometrika*, 30, 81–89, 1938.

5

Koster, R. D., Dirmeyer, P. A., Guo, Z., Bonan, G., Chan, E., Cox, P., Gordon, C. T., Kanae, S., Kowalczyk, E., Lawrence, D., Liu, P., Lu, C.-H., Malyshev, S., McAvaney, B., Mitchell, K., Mocko, D., Oki, T., Oleson, K., Pitman, A., Sud, Y. C., Taylor, C. M., Verseghy, D., Vasic, R., Xue, Y., and Yamada, T.: Regions of Strong Coupling Between Soil Moisture and Precipitation, *Science*, 305, 1138–1140, 2004.

10

Leuning, R., Zhang, Y., Rajaud, A., Cleugh, H., and Tu, K.: A simple surface conductance model to estimate regional evaporation using MODIS leaf area index and the Penman-Monteith equation, *Water Resources Research*, 44, 2008.

McNaughton, K. G., and Jarvis, P. G.: Predicting effects of vegetation changes on transpiration and evaporation, in: *Water Deficits and Plant Growth*, edited by T. T. Kozlowski, Vol. 7, pp. 1-47. Academic Press, New York, 1983.

15

Medlyn, B. E., Duursma, R. A., Eamus, D., Ellsworth, D. S., Prentice, I. C., Barton, C. V. M., Crous, K. Y., de Angelis, P., Freeman, M., and Wingate, L.: Reconciling the optimal and empirical approaches to modelling stomatal conductance, *Global Change Biology*, 17, 2134–2144, 2011.

20

Miralles, D. G., Gash, J. H., Holmes, T. R. H., de Jeu, R. A. M., and Dolman, A. J.: Global canopy interception from satellite observations, *Journal of Geophysical Research*, 115, D16122, 2010.

Morison, J. I.: Sensitivity of stomata and water use efficiency to high CO₂, *Plant, Cell & Environment*, 8, 467–474, 1985.

35

Nash, J. E. and Sutcliffe, J. V.: River flow forecasting through conceptual models part I — A discussion of principles, *Journal of Hydrology*, 10, 282–290, 1970.

Oki, T. and Kanae, S.: Global hydrological cycles and world water resources, *Science*, 313, 1068–1072, 2006.

40

Omlin, M. and Reichert, P.: A comparison of techniques for the estimation of model prediction uncertainty, *Ecological Modelling*, 115, 45–59, 1999.

Papale, D., Reichstein, M., Aubinet, M., Canfora, E., Bernhofer, C., Kutsch, W., Longdoz, B., Rambal, S., Valentini, R., Vesala, T. and Yakir, D.: Towards a standardized processing of Net Ecosystem Exchange measured with eddy covariance technique: algorithms and uncertainty estimation, *Biogeosciences*, 3, 571-583, 2006.

5 Reichstein, M., Falge, E., Baldocchi, D., Papale, D., Aubinet, M., Berbigier, P., Bernhofer, C., Buchmann, N., Gilmanov, T., Granier, A., Grünwald, T., Havránková, K., Ilvesniemi, H., Janous, D., Knohl, A., Laurila, T., Lohila, A., Loustau, D., Matteucci, G., Meyers, T., Miglietta, F., Ourcival, J.-M., Pumpanen, J., Rambal, S., Rotenberg, E., Sanz, M., Tenhunen, J., Seufert, G., Vaccari, F., Vesala, T., Yakir, D. and Valentini, R.: On the separation of net ecosystem exchange into assimilation and ecosystem respiration: review and improved algorithm, *Global Change Biology*, 11, 1424–1439, 2005.

10

Tenhunen, J., Serra, A. S., Harley, P., Dougherty, R., and Reynolds, J. F.: Factors influencing carbon fixation and water use by Mediterranean sclerophyll shrubs during summer drought, *Oecologia*, 82, 381–393, 1990.

Whitehead, D., Okali, D., and Fasehun, F.: Stomatal response to environmental variables in two tropical forest species
15 during the dry season in Nigeria, *Journal of Applied Ecology*, pp. 571–587, 1981.

Wythers, K. R., Lauenroth, W. K., and Paruelo, J. M.: Bare-soil evaporation under semiarid field conditions, *Soil Science Society of America Journal*, 63, 1341–1349, 1999.

20 Zhou, S., B. Yu, Y. Huang, and Wang, G.: The effect of vapor pressure deficit on water use efficiency at the subdaily time scale, *Geophysical Research Letters*, 41, 5005–5013, 2014.

Zhou, S., Yu, B., Huang, Y., and Wang, G.: Daily underlying water use efficiency for AmeriFlux sites, *Journal of Geophysical Research: Biogeosciences*, 120, 887–902, 2015.

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Data Availability

For this study, we used observations of the FLUXNET initiative from sites with an *open and fair use data policy*⁴. The data sets can be downloaded at <http://fluxnet.fluxdata.org//data/download-data/>

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⁴ http://www.fluxdata.org/Shared%20Documents/Policy_Free_Final.pdf