Spatial and temporal variations in plant Water Use Efficiency inferred from tree-ring, eddy covariance and atmospheric observations

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Abstract

Plant Water Use Efficiency (WUE), which is the ratio of the uptake of carbon dioxide through photosynthesis to the loss of water through transpiration, is a very useful metric of the functioning of the land biosphere. WUE is expected to increase with atmospheric CO$_2$, but to decline with increasing atmospheric evaporative demand – which can arise from increases in near-surface temperature or decreases in relative humidity. We have used $\Delta^{13}$C measurements from tree-rings, along with eddy-covariance measurements from Fluxnet sites, to estimate the sensitivities of WUE to changes in CO$_2$ and atmospheric humidity deficit. This enables us to reconstruct fractional changes in WUE, based on changes in atmospheric climate and CO$_2$, for the entire period of the instrumental global climate record. We estimate that overall WUE increased from 1900 to 2010 by 48±22%, which is more than double that simulated by the latest Earth System Models. This long-term trend is largely driven by increases in CO$_2$, but significant inter-annual variability and regional differences are evident due to variations in temperature and relative humidity. There are several highly populated regions, such as Western Europe and East Asia, where the rate of increase of WUE has declined sharply in the last two decades. Our data-based analysis indicates increases in WUE that typically exceed those simulated by Earth System Models – implying that these models are either underestimating increases in photosynthesis or underestimating reductions in transpiration.
1. Introduction

Plant Water Use Efficiency ($WUE$) is the ratio of the $CO_2$ assimilated through photosynthesis (Gross Primary Productivity, $GPP$), to the water used by plants as the flux of Transpiration ($E_T$):

$$WUE = \frac{GPP}{E_T}$$

(1)

Carbon dioxide may affect plants through increases in photosynthesis (Ainsworth and Rogers 2007; Franks et al. 2013) and possible reductions in transpiration associated with the partial closure of leaf stomatal pores under elevated $CO_2$ (Field et al. 1995; Gedney et al. 2006; Betts et al. 2007). Both of these effects are uncertain though. $CO_2$-fertilization of photosynthesis is often found to be limited by nutrient availability (Norby et al. 2010), and large-scale transpiration may not reduce even with $CO_2$-induced stomatal closure, if plant leaf area index increases enough to counteract reduced transpiration from each leaf (Piao et al. 2007). $WUE$ does however appear to be increasing more robustly with $CO_2$, according to both tree-ring (Franks et al. 2013) and eddy-covariance flux measurements (Keenan et al. 2013).

Plant photosynthesis and transpiration are coupled through the behaviour of leaf stomatal pores, through which $CO_2$ must diffuse to be fixed during photosynthesis, and through which the transpiration flux escapes to the atmosphere. The combined behaviour of the leaf stomata leads to an environmentally dependent “canopy conductance” that controls both the water and carbon fluxes. As a consequence, both $GPP$ and $E_T$ can be written as the product of a canopy conductance and a
concentration gradient, which is sometimes described as an electrical analogue (Cowan 1972). For GPP, the concentration gradient is the difference between the atmospheric CO₂ concentration at the leaf surface \( (C_a) \) and the internal CO₂ concentration within plant leaves \( (C_i) \):

\[
GPP = g_c (C_a - C_i) \tag{2}
\]

where \( g_c \) is the canopy conductance for CO₂.

For \( E_T \), the concentration gradient is the difference between the specific humidity of the atmosphere at the leaf surface \( (q_a) \) and the specific humidity inside the plant leaves, which is saturated at the leaf temperature \( (q_{sat}) \). The canopy conductances for GPP and \( E_T \) both arise from diffusion through leaf stomatal pores, and therefore only differ by a constant factor of 1.6 (the square root of the ratio of the molecular masses of CO₂ and H₂O).

\[
E_T = 1.6g_c (q_{sat} - q_a) \tag{3}
\]

Changes in stomatal opening in response to changes in sunlight, atmospheric temperature and humidity, soil moisture, and CO₂, are complex and uncertain (Berry et al. 2010), as are the scaling of these leaf-level responses up to the canopy and beyond (Piao et al. 2007; Jarvis and McNaughton 1986; Jarvis 1995). However, since stomatal behaviour affects transpiration and photosynthesis similarly, WUE is relatively insensitive to these uncertainties:

\[
WUE = \frac{(C_a - C_i)}{1.6(q_{sat} - q_a)} = \frac{(C_a - C_i)}{1.6D} = \frac{C_a(1-f)}{1.6D} \tag{4}
\]
where $D$ is the atmospheric humidity deficit ($q_{\text{sat}} - q_a$) and $f$ is the ratio of the internal to the external CO$_2$ concentration ($C_i/C_a$). This equation therefore expresses $WUE$ in terms of atmospheric variables, $C_a$ and $D$ (which itself depends on relative humidity and temperature), along with the factor $f$. The remaining uncertainty associated with plant physiology is therefore contained in $f$.

In the absence of water limitations, there is good evidence that $f$ is approximately independent of $C_a$, so that $C_i$ remains proportional to $C_a$, unless $D$ changes (Jacobs 1994; Katul et al. 2010; Leuning 1995; Morison et al. 1983). Even during drought, $f$ will vary with $D$, due in part to correlations between $D$ and soil moisture (Brodribb 1996).

Stomatal optimisation theories, which assume that stomata act so as to maximise photosynthesis for a given amount of available water (Cowan and Farquhar 1977), also suggest that $f$ should depend predominantly on $C_a$ and $D$ (Katul et al. 2010; Medlyn et al. 2011). Absolute values of $WUE$ will depend on the nature of the vegetation and soil, such as the plant and soil hydraulics, but these optimisation theories imply that there will be a near universal sensitivity of fractional changes in $WUE$ to fractional changes in $C_a$ and $D$ (see SI Appendix):

$$\frac{WUE}{WUE(0)} = \left(\frac{C_a}{C_a(0)}\right)^a \left(\frac{D}{D(0)}\right)^b$$

where the subscript (0) denotes the initial state of each variable, and $a$ and $b$ are dimensionless coefficients. For given $a$ and $b$ values this equation describes how the fractional change in $WUE$ at each location varies with fractional changes in $C_a$ and $D$. Although they differ in their underlying
assumptions and detailed conclusions, it is interesting to note that the latest stomatal optimization theories (Katul et al. 2010; Medlyn et al. 2011) both imply $a=1$ and $b=-0.5$ (see SI Appendix).

We focus in this study on fractional changes in $WUE$, which are more likely to be independent of these complex factors. Therefore, we use two very different datasets of $WUE$, derived from tree ring measurements and eddy-covariance fluxes and aim to model the fractional changes in plant $WUE$ by using atmospheric data alone. The longer-term climate signals are derived from the tree rings, spanning at least the last 100 years. Monthly $WUE$ values are derived from eddy-covariance observations between 1995-2006. We do not assume the applicability of stomatal optimization theories, but instead adopt equation 5 as a parsimonious empirical model for the fractional changes in $WUE$ observed at each measurement site, given suitable fitting parameters $a$ and $b$. Tests using more elaborate statistical models, with additional environmental variables or vegetation-specific parameters, were not found to produce significant improvements in the fit to the observed changes in $WUE$ despite the introduction of extra fitting parameters. Finally, we have compared our reconstruction of the fractional change in plant $WUE$ to Earth System Models (ESMs) simulations, focussing on regional variations in the $WUE$ changes and how these compare to the long-term tree ring observations.
2. Materials and Methods

We estimate the sensitivity of WUE to $C_a$ and $D$ by fitting to WUE changes inferred from both eddy-covariance fluxes (relatively short records with high-temporal resolution) and carbon isotope records from tree-rings (longer-term records with annual resolution). We use observations from 28 eddy-covariance and 31 tree-ring sites (see Figure 1 and SI Appendix, Tables S1 and S2).

2.1 Eddy-covariance observations.

The carbon and water flux observations were taken from the *Free Fair-use* Fluxnet database (www.fluxdata.org) (Baldocchi 2008; Papale et al. 2006; Reichstein et al. 2005). We selected a total of 28 sites based on data availability (Table S1). Monthly WUE was estimated from Equation 1 with $GPP$ used directly from the database (Reichstein et al. 2005). In general the total latent heat flux ($LE$) has contributions from interception loss, soil evaporation and transpiration. We follow previous studies (Dekker et al. 2001; Groenendijk et al. 2011; Keenan et al. 2013; Law et al. 2002) in assuming that the latent heat flux is dominated by transpiration during periods with no rain in the preceding two days, when the interception loss and soil evaporation are assumed small. Monthly average values of $GPP$, $E_T$, $C_a$, $D$ and $T$ were calculated from half-hourly observations (not gap-filled) during dry periods (*i.e.* no rain in the preceding two days) when $GPP$ was larger than zero. To exclude periods with unrealistic WUE values due to the division by very small $E_T$ values, we used only months during the growing season. Annual average growing season values were calculated from the months with an average temperature above $10^\circ$C. Only sites with at least 6 annual values were used, resulting in a dataset of 222 annual growing season values of WUE, $C_a$ and $D$. Data are used between 1995 and 2006. Fractional changes were calculated relative to the mean over the observational period for each of the sites, to enable comparison between sites.
2.2. Tree-ring observations.

To derive a longer-term relationship between the fractional change in WUE and variations in $C_a$ and $D$, we used $\Delta^{13}C$ tree-ring observations from 31 locations (Figure 1), ranging from 1900 to current, as described in two previous studies (Franks et al. 2013, Hemming et al. 1998) (see SI Appendix, Table S2). The discrimination of $^{12}C$ against $^{13}C$ ($\Delta^{13}C$) is estimated from the tree-ring samples (Hemming et al. 1998; van der Sleen et al. 2015). The $\Delta^{13}C$ measurements can be used to estimate the ratio of the internal to the external CO$_2$ concentration ($f=C_a/C$) using the relationship: $f=(\Delta^{13}C-4.4)/(27-4.4)$, where $\Delta^{13}C$ is in parts per thousand ($\text{‰}$), and $C_a$ is taken from the Mauna Loa atmospheric CO$_2$ record (Farquhar et al. 1989; Franks et al. 2013; Keeling et al. 1976). WUE is estimated with Equation 4 using annual average growing season values of $D$ from the CRU dataset, taking the nearest pixel to each site (Harris et al. 2013). This large-scale dataset for $D$ ensures consistency among the sites, but may underestimate the finer spatial variation in $D$. As for the eddy-covariance sites, we estimated the fractional changes relative to the mean over the observational period at each of the sites. For this analysis we have 1007 observations of WUE derived from tree-ring observations of $\Delta^{13}C$.

2.3 Fractional WUE

To estimate $a$ and $b$ with a linear regression model we rewrite Equation 5 in a logarithmic form:

$$\ln \left(1 + \frac{\Delta WUE_{\text{WUE}(0)}}{WUE_{\text{WUE}(0)}}\right) = a \ln \left(1 + \frac{\Delta C_a}{C_{\text{a}(0)}}\right) + b \ln \left(1 + \frac{\Delta D}{D(0)}\right)$$  (6)

Here the second-term in each bracket represents the fractional change in WUE, $C_a$ and $D$, respectively. These fractional change variables are used in all our subsequent statistical analyses and modelling.
We set out to fit the fractional change in WUE at each observation site (Figure 1) from the fractional change in $C_a$ and the fractional change in $D$. For comparison and fitting we therefore need to calculate $WUE(0)$, $C_a(0)$ and $D(0)$ for the observational data, which we take as the mean over the entire observational record available at each site.

2.4 Global fractional change of WUE

The dependence of fractional changes in WUE on $C_a$ and $D$ allows us to use these relationships to estimate changes in WUE at large scales using global climate data. The fractional change in $D$ can be further partitioned into a change in temperature ($T$) and relative humidity ($RH$), which makes it possible to separate the effect of changes in these variables on WUE (see SI Appendix). To do this, we used the CRU climate dataset (Harris et al. 2013) at a $0.5\degree \times 0.5\degree$ latitude/longitude grid and the annual CO$_2$ concentration at Mauna-Loa (Keeling et al. 1976) to derive the global and local variation in WUE. We only used months during the growing season when photosynthesis occurs, assumed to be above a monthly average temperature threshold of $10\degree C$. For the period 1900-1930 the average temperature was calculated for each month from which a spatial mask was generated (SI Appendix, Fig. S4). This mask was then used to calculate annual time-evolving values of WUE from the growing season values of temperature and humidity for each year between 1901 and 2010.

2.5 Other independent data sources

For three locations, Western North America, Western Europe and East Asia, we have compared our simulated fractional change in plant WUE with remote sensing (RS) products of GPP and $E_T$ (Jung et al. 2011). This dataset covers the period 1982-2006; we use the period 1986-1990 as a reference period for both our estimate and the fractional change in WUE from the RS product. As the RS data
does not cover a response to changes in $C_a$, we estimated the fractional change in $WUE$ with and without the $C_a$ response.

At the global scale, we have compared our simulated fractional change in plant $WUE$ with simulations of Earth System Models (ESMs). Most of the latest ESMs calculate changes in both $GPP$ and $E_T$. This allows a comparable change in $WUE$ to be calculated for 28 CMIP5 models (Taylor et al. 2012) based-on their historical simulations. Finally, regional differences in responses are compared to the 31 tree ring observation sites.
3 Results and Discussion

Figure 2 summarises the derivation of the $a$ and $b$ parameters, which are the sensitivity of $WUE$ to $C_a$ and $D$, for the tree-ring and eddy-covariance observations. In general, eddy-covariance data alone is unable to fully constrain the CO$_2$ sensitivity of the $WUE$ (Keenan et al. 2013), because the data records are too short to sample significant changes in CO$_2$, resulting in a value of $a$ of 0.79±0.79 for all eddy-covariance site (SI Appendix, Fig. S1 and Table S3). However, the longer tree-ring records overall yield a good constraint on $a$ of 1.61±0.54. The annual data-points for the two datasets can be combined into a single dataset. Fitting against this more complete dataset gives generic sensitivity coefficients of $a$=1.51±0.57 and $b$=-0.72±0.16. These values are mainly constrained by the tree-ring observations for which the fits to equation 6 are more tightly defined (Fig. 2a and SI Appendix, Fig S1 top row).

A value of $a$ larger then 1, suggests that $WUE$ has been increasing even faster than the atmospheric CO$_2$ concentration (Fig. 2c). This is qualitatively consistent with conclusions from a previous study, which was based purely on eddy covariance data (Keenan et al. 2013), but is more robustly demonstrated here due to the much longer tree-ring records.

It is interesting to note that our overall values of $a$=1.51 ± 0.57 and $b$=-0.72 ± 0.16 are larger by about 50% than the values derived from stomatal optimization theories: $a$=1.0, $b$=-0.5 (Cowan and Farquhar 1977; Katul et al. 2010; Palmroth et al. 2013), indicating a stronger response to changes in both CO$_2$ and climate. Such theoretical sensitivities are common to variants of stomatal optimization theory, including those that assume either electron transport-limited or Rubisco-limited photosynthesis, and even when additional nitrogen limitations are accounted for (Prentice et al. 2013). The differences between the optimization theory and our empirically derived $WUE$ sensitivities may arise from
differences between leaf-surface and atmospheric values of CO$_2$ and humidity, but they may also be indicative of missing constraints and feedbacks in the optimization theories (Lin et al. 2015; Prentice et al. 2014; de Boer et al. 2011, 2016).

Testing more elaborate statistical models.

Equation 6 is motivated by empirical evidence and theory suggesting that WUE should vary predominantly with $C_a$ and $D$. However, it is conceivable that the fractional change in WUE could also depend on other environmental conditions or the detailed vegetation type. In order to test for this, we carried out two additional sets of fits against the observational data. In the first test we extended our statistical model (equation 6) to include other environmental variables that had been measured at the Fluxnet sites, most notably solar radiation, air temperature, and soil water content. Including these additional predictor variables does not significantly improve the fit to the observed changes in WUE (as measured by $r^2$), and typically results in less robust predictions (as measured by the adjusted $r^2$), because of the introduction of extra fitting parameters (SI Appendix, Table S4). In the second test, we carried out separate statistical fits for each of the sites listed in the Fluxnet dataset. Clustering of these values by vegetation type would indicate that $a$ and $b$ parameters are dependent on vegetation type, but we find no evidence of such clustering (Figure 3).

Comparison to independent WUE estimates.

Our best-fit generic $a$ and $b$ parameters are able to reasonably reproduce the fractional changes in WUE due to fractional changes in both $C_a$ and $D$ across the 59 tree-ring and eddy-covariance sites (SI Appendix, Fig. S2). However, it is important to evaluate the estimated response of WUE to $C_a$ and $D$ against independent data. We compared the change in WUE estimated with the best-fit parameters to
observations at three tropical tree-ring sites from a recent study (van der Sleen et al. 2015). At these sites a range of species of both trees and under-storey were sampled. Our estimate for these three locations passes close to the mean of the observed WUE fractional changes (Fig. 4a-c). Because the RS data does not include a response to changes in $C_a$, we estimated a fractional change in WUE with and without this response (Fig. 4d-f) for three regions, which show distinct changes in WUE: Western North America, Western Europe and East Asia. The RS fluxes show little inter-annual variability, and much less variability than we estimate. For the three regions in Fig. 4d-f our estimates with and without CO$_2$ effects sit on either side of the RS estimates. In the Amazon, South Africa and South East Asia (see SI Appendix, Fig. S3) our estimates excluding CO$_2$ effects are similar to the RS estimates, whilst the inclusion of CO$_2$ effects leads to significant increases in WUE (SI Appendix, Fig. S3) that appear to be inconsistent with the RS estimates (which do not account for CO$_2$ changes), but are more consistent with the tree-ring (Franks et al. 2013) and eddy-covariance data (Keenan et al. 2013).

**Global fractional change of WUE.**

Globally, we estimate that WUE has increased by $48\pm22\%$ since 1900 (Fig. 5a), with the CO$_2$ increase contributing $+47\pm21\%$ and relative humidity contributing $+3.6\pm1.3\%$, counteracted by a much smaller reduction in WUE due to warming of $-2.3\pm0.8\%$. Estimated fractional changes in WUE between 1901-1930 and 2001-2010 differ regionally between 0.1 and 0.6 (Fig. 5b). Uncertainties in global WUE changes were derived from the range of the parameters $a$ and $b$ within 5% of the RMSE of our best fit (Fig. 2c).

**Comparison to simulations with complex Earth System Models (ESMs).**
The CMIP5 models simulate an increase in WUE of between 2% and 28% to 2005, with an ensemble mean of 14% (SI Appendix, Table S5). For comparison, our overall fit against the tree-ring and eddy-covariance data indicates an approximately 40% increase in WUE over the same period. Figure 6 compares the annual time-series of the fractional changes in WUE from the CMIP5 models (black line and green uncertainty plumes), our statistical fit (orange lines), and the mean changes observed for the tree-ring (black marks and grey uncertainty bars) and eddy-covariance sites (dark blue marks and light blue uncertainty bars). This comparison suggests that the latest ESMs significantly underestimate the historical increase in WUE.

Regional changes in WUE.

Our global average change in WUE hides substantial regional differences (Fig. 5b). This is a result of the spatially and temporally varying impact of climate change on WUE (Fig. 7a and animation in the Supporting Information), driven by the heterogeneity of the warming (Fig. 7b) and the large variation in changes in near-surface RH (Fig. 7c). In many regions the overall impact is a significant increase in WUE, such as Western North America (Fig. 7d). However, the recent rate of increase has declined substantially in several heavily populated regions. For example, WUE shows a slower increase in Western Europe since the 1980s, as a result of increases in T, which has counteracted the WUE increase due to increasing CO₂ (Fig. 7e). This is also observed in WUE trends derived from isotopic tree-ring observations in Spain (Linares et al. 2012). Our analysis indicates that East Asia has suffered an even more significant suppressions of WUE since about 1990, due predominantly to reductions in RH (Wang et al. 2012) (Fig. 7f). This pattern of changing RH is comparable with the trends in precipitation and drought since 1950 (Dai 2011).
For the 31 tree ring observation sites, we have plotted the ensemble mean regional ESM model results against the individual observed tree ring data (Fig. S5). For 10 out of 31 observation sites, the simulated fractional $WUE$ increases between 5-10%. For 3 out of the 31 sites, the fractional $WUE$ increases by more than 50%, and for 14 out of the 31 observation sites the $WUE$ change inferred from the tree ring data is significantly higher than that simulated by the ESMs.
4. Conclusions

This study shows that fractional changes in plant WUE, at large-scales and over the period of the climatological record, can be inferred from atmospheric data alone. By combining two very different datasets of WUE derived from tree-ring Δ¹³C measurements and eddy-covariance fluxes we have derived a consistent response of the fractional change in WUE to the fractional changes in $C_a$ and $D$. This generic response can be used to estimate WUE changes over the entire period of the atmospheric record. Our analysis shows that global WUE increased by approximately a half over the 20th century predominantly due to rising CO₂, which is significantly more than is simulated by the latest Earth System Models. However, this increase in WUE has been modulated downwards in recent decades by the impact of climate change. This is especially true for the highly populated regions of Western Europe and East Asia, where reductions in atmospheric relative humidity and increases in temperature have acted to offset increases in WUE due to increasing CO₂. We conclude that the effects of increasing CO₂ on plant WUE are significantly underestimated in the latest climate projections.

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Figure Legends:

Figure 1. Locations of the eddy-covariance flux sites and tree-ring sites used (see Table S2 and S3 for a list of the sites). Tropical tree-ring sites are used as independent data sources for comparison.

Figure 2. Water Use Efficiency (WUE) from tree-ring and eddy-covariance observations. The relationship between the observed fractional change in WUE and the fractional change in (A) CO₂ concentration and (B) humidity deficit of both datasets is fitted to Equation 3, with best-fit values for a (1.51±0.57) and b (-0.72±0.16). (C) The colors show the root mean square error (RMSE) of the simulated vs. observed fractional change in WUE as a function of a and b, with the black area representing the best parameters within 5% of the RMSE of the best fit (white star). The black star represents the values according to the optimality hypothesis.
Figure 3. Comparison of best-fit a and b parameters (see equation 5) by plant functional type (PFT). Here the sites are organized by dominant PFT, using classifications used for the FluxNet sites: evergreen broadleaf forest (EBF), evergreen needleleaf forest (ENF), grassland (GRA), mixed forest (MF), wetland (WET) and woody savannah (WSA).
Figure 4. Comparison of estimated Water Use Efficiency trends to independent observations. Simulated fractional change in WUE (orange) compared to observations for three tropical tree-ring sites in Bolivia (A), Cameroon (B) and Thailand (C) (blue, van der Sleen 2015). Simulated fractional change in WUE for (D) Western North America, (E) Western Europe and (F) East Asia, with (dark red) and without (orange) CO₂ effect, compared to the WUE trend derived from a remote sensing product of carbon uptake and water loss (Jung et al. 2011). The location of the tree-ring sites is presented in Fig. S1 and the regions D-F are as in Fig. 5.
Figure 5. 20th century fractional change of Water Use Efficiency (WUE). (A) Time series of the estimated global fractional change in WUE (orange, relative to the average over 1901-1930) partitioned into the effects of changes in CO₂, relative humidity and temperature. (B) Spatial pattern of the estimated fractional change in WUE between 1901-1930 and 2001-2010. These calculations use observed monthly surface air temperature and vapour pressure (Harris et al. 2013) during the growing season, and annual atmospheric CO₂ concentrations at Mauna Loa (Keeling et al. 1976).
Figure 6. Comparison of measured and modeled fractional changes in WUE from 1860 to 2010. Estimates from tree-rings and eddy-covariance data are shown by the black and blue points respectively, with the bars in each case showing +/-1 standard deviation about the mean response. The results from complex coupled Earth System Models are shown by the black continuous line and the green plume (with dark green showing one standard deviation and light green showing two standard deviations). The algorithm presented in this paper, which estimates fractional WUE changes from changes in CO2 concentration and humidity deficit alone (equation 6), is shown by the orange lines. To enable the comparison between these different estimates, we normalized over common overlapping periods (for the tree-ring data and model simulations – 1900-1930; for the tree-ring and eddy-covariance data – the period of overlap when at least 3 eddy-covariance sites are available).
Figure 7. Changes in WUE arising from climate variables. Spatial patterns of the fractional changes in WUE arising from changes in (A) climate, i.e. both temperature and relative humidity (RH) together, (B) temperature alone, and (C) RH alone, between 1901-1930 and 2001-2010. Time-series are as in Fig. 5 for (D) Western North America, (E) Western Europe and (F) East Asia, which show the large regional and temporal variations in these climate-driven changes in WUE.