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Abstract

Partitioning evapotranspiration (ET) is essential for improving water resource management and understanding the global hydrological cycle. However, ET partitioning in various ecosystems is challenging as some assumptions are restricted to certain areas or plant types. Here, we developed a novel ET partitioning method coupling definitions of leaf and ecosystem water use efficiencies (WUE_{leaf} and WUE_{eco} , respectively). We used 25 eddy covariance flux sites for 196 site-years to evaluate $T:ET$ characteristics of seven plant functional types (PFTs) at different spatiotemporal scales. The results indicated the spatiotemporal characteristics of WUE_{leaf} and WUE_{eco} were not consistent, resulting in $T:ET$ variation in the seven PFTs. Deciduous broadleaf forests showed the highest mean annual $T:ET$ (0.67), followed by evergreen broadleaf forests (0.63), grasslands (0.52), evergreen needleleaf forests (0.46), and woody savanna (0.41), and C_3 croplands had higher $T:ET$ (0.65) than C_4 croplands (0.48). The annual mean leaf area index (LAI) explained about 26% of the variation in $T:ET$, with the trend in $T:ET$ consistent with the known effects of LAI. The overall trends and magnitude of $T:ET$ in this study were similar to different results of ET partitioning methods globally. Importantly, this method improved $T:ET$ estimation accuracy in vegetation-sparse and water-limited areas. Our novel ET partitioning method is suitable for estimating $T:ET$ at various spatiotemporal scales and provides insight into the conversion of WUE at different scales.

1. Introduction

Evapotranspiration (ET)—including evaporation of soil water and water intercepted by the plant canopy (E), and stomatal transpiration (T)—is crucial for understanding global ecohydrological systems [Katul *et al.*, 2012; Wang and Dickinson, 2012; Kool *et al.*, 2014]. Transpiration is directly in connection with biological processes with photosynthesis for plant productivity and usually considered as productive water loss [Granier *et al.*, 1999; Jasechko *et al.*, 2013; Yi *et al.*, 2019]. Enhancing the productive part (T) and decreasing the non-productive part (E) is critical for the sustainable water resources management in drylands [Newman *et al.*, 2010; Kool *et al.*, 2014]. Therefore, partitioning ET into E and T is essential for land surface process models [Lawrence *et al.*, 2007], quantifying water use efficiency (WUE), and coupling hydrological and biogeochemical cycles [Austin *et al.*, 2004; Mastrotheodoros *et al.*, 2017]. However, partitioning ET continuously is challenging in most ecosystems.

Some methods from the site to ecosystem scale have been used to investigate the ratio of T to ET , denoted as $T:ET$, including sap flow [Rafi *et al.*, 2019], stable isotope [Wang *et al.*, 2010; Xiao *et al.*, 2018], eddy covariance [Paul-Limoges *et al.*, 2020], and model simulation [Gu *et al.*, 2018]. Sap flow can continuously record plant transpiration, but it is difficult to scale up to field ecosystems [Kool *et al.*, 2014]. Water stable isotopes for ET partitioning are based on differences in the water vapor isotope of E and T , but this method costs a lot of human and financial resources and makes continuous observation difficult [Griffis, 2013; Xiao *et al.*, 2018]. Mechanistic and empirical models can overcome these issues, but the debatable hypothesis [Schlaepfer *et al.*, 2014] and substantial number of parameters [Kool *et al.*, 2014] have caused some uncertainties in modeling simulations. Eddy covariance technique is used to measure exchanges of carbon and water flux between atmosphere and underlying surface, and flux networks have been more than 900 sites worldwide [Baldocchi and Ryu, 2011; Zhou *et al.*, 2016]. Traditionally, eddy-covariance techniques ignore understory T and plant E from canopy rainfall interception. A new ET partitioning method was introduced by Scanlon and Sahu [2008], which assumed that the relationship between T and E is related to stomatal fluxes and non-stomatal fluxes, and involved the parameter of leaf WUE [Scanlon and Kustas, 2010]. However, this

method has not been widely adopted as it requires high frequency (10–20 Hz) data; therefore, some sites in global flux networks cannot be used due to the flux-variance similarity assumption [Wagle *et al.*, 2020]. Zhou *et al.* [2016] introduced a novel ET partitioning method based on underlying water use efficiency (uWUE), which is easy to use in practice and can estimate $T:ET$ at various spatiotemporal scales [Berkelhammer *et al.*, 2016; Zhou *et al.*, 2016; Jiang *et al.*, 2020], but has some limitations. Firstly, the assumption that $T = ET$ at times during the growing season might be invalid in arid and vegetation-sparse areas, where E cannot be ignored. Therefore, this method would overestimate $T:ET$ [Scott and Biederman, 2017; Li *et al.*, 2019;]. Secondly, a linear relationship between $(1-C_i/C_a)$ and the square root of vapor pressure deficit ($VPD^{0.5}$) is only applicable for C_3 plants, so leaf uWUE is not available for C_4 plants due to the lack of marginal water cost of carbon gain (λ_c) [Lloyd and Farquhar, 1994]. Thirdly, the marginal WUE changes when stomata no longer behave optimally under severe water deficit due to limited xylem water transport, the uWUE method is not useful [Zhou *et al.*, 2018].

Here, we developed a novel and simple ET partitioning method that accounts for leaf and ecosystem WUE (WUE_{leaf} and WUE_{eco}) and applies to all areas and plant types using accessible half-hourly eddy-covariance flux data. To provide a reliable evidence for the proposed novel ET partitioning method, this study aimed to (1) evaluate WUE_{leaf} and WUE_{eco} among plant functional types (PFTs) at spatiotemporal scales, (2) evaluate $T:ET$ variation in PFTs at spatiotemporal scales, and (3) evaluate the feasibility of the new ET partitioning method and its applicability in arid areas. The proposed method can evaluate characterization of $T:ET$ at multiple spatiotemporal scales at global flux tower networks.

2. Methods

The theoretical foundation of water loss from E , T , and ET is associated with non-stomatal, stomatal, and stomatal/non-stomatal mixing behaviors, respectively [Scanlon and Sahu, 2008; Scanlon and Kustas, 2010]. Transpiration is related to plant and regulated by leaf stomas, while E is only related to soil and environmental conditions. According to observation scales and water consumption differences, WUE is defined differently at various spatiotemporal scales. The concept of WUE includes ecosystem WUE (WUE_{eco}), which combines stomatal and non-stomatal mixing behaviors, and ecosystem transpiration efficiency ($iWUE_{eco}$), which is only associated with stomatal behaviors. WUE_{eco} is defined as the ratio of carbon fixation (GPP) to water consumption by the ecosystem (ET), and $iWUE_{eco}$ is defined as the ratio of GPP to water consumption via transpiration (T). Thus, $T:ET$ can be determined from the ratio of WUE_{eco} to $iWUE_{eco}$. WUE_{eco} can be estimated directly from GPP and ET provided by flux-covariance sites. It is difficult to obtain $iWUE_{eco}$ but it can be up-scaled from WUE_{leaf} indirectly. Ecosystem WUE is defined as the ratio of ecosystem GPP of ecosystem to water consumption via evapotranspiration (ET), and it can also be decomposed into GPP/ T and $T:ET$:

$$WUE_{eco} = \frac{GPP}{ET} = \frac{GPP}{T} \times \frac{T}{ET} = iWUE_{eco} \times \frac{T}{ET} \quad (1)$$

In this study, evapotranspiration (ET) includes transpiration (T), soil water evaporation (E_s) and canopy interception evaporation (E_i):

$$ET = T + E_s + E_i \quad (2)$$

Studies indicated that WUE at the leaf level can be directly up-scaled to the canopy level in a consistent natural environment [Barton *et al.*, 2012; Linderson *et al.*, 2012]. Zhou *et al.* [2016] also tested that uWUE at the leaf level is broadly consistent with the ecosystem level ($GPP \cdot VPD^{0.5}/T$). Thus, ecosystem $iWUE_{eco}$ (GPP/T) in Eq. (3), can be approximated the WUE at the leaf level (WUE_{leaf}) [Cheng *et al.*,

2017; Medlyn *et al.*, 2011].

$$iWUE_{eco} = \frac{GPP}{T} = \frac{\int A dt}{\int T dt} \approx WUE_{leaf} = \frac{A}{T} = \frac{C_a P_a}{1.6(VPD + g_1 \sqrt{VPD})} \quad (3)$$

where A is leaf net photosynthetic rate ($\mu\text{mol}(\text{CO}_2) \text{ m}^{-2} \text{ s}^{-1}$), T is leaf transpiration rate ($\mu\text{mol}(\text{H}_2\text{O}) \text{ m}^{-2} \text{ s}^{-1}$), P_a is atmospheric pressure (kPa), C_a is ambient atmospheric CO_2 concentration ($\text{mol}(\text{CO}_2) \text{ mol}^{-1}$), VPD is vapor pressure deficit (kPa), and g_l is an empirical parameter of the stomatal conductance model ($\text{kPa}^{0.5}$), representing the exchange rate between carbon uptake and water use [Knauer *et al.*, 2018; Medlyn *et al.*, 2011]. Lin *et al.* [2015] compiled the mean g_l value of different PFTs using a global-scale extensive field observation dataset. The g_l values at the fluxnet tower sites in our study were extracted from a global map of g_l parameters (see Supplementary Information of Cheng *et al.* [2017]) by interpolating g_l values of different PFTs with a global plant classification map (SYNMAP). The g_l values at the fluxnet tower sites in our study are shown in Table 1.

Substituting Eq. (3) into Eq. (1):

$$WUE_{eco} = WUE_{leaf} \times \frac{T}{ET} \quad (4)$$

Thus, the $T:ET$ can be estimated from the ratio of WUE_{eco} to WUE_{leaf} as follows:

$$\frac{T}{ET} = \frac{WUE_{eco}}{WUE_{leaf}} = \frac{\frac{GPP}{ET}}{\frac{C_a P_a}{1.6(VPD + g_1 \sqrt{VPD})}} \quad (5)$$

3. Datasets

3.1 Flux tower data

Half-hourly flux-tower dataset from the FLUXNET2015 (<https://fluxnet.org>) were extracted. Sites were selected with the following data recorded: net solar radiation, air temperature, precipitation, latent heat flux, atmospheric pressure, ambient atmospheric CO_2 concentration, vapor pressure deficit (VPD), and estimates of gross primary productivity (GPP). Air temperature and latent heat flux at half-hourly intervals data were used to calculate ET ($\text{kg H}_2\text{O m}^{-2} \text{ d}^{-1}$) [Donatelli *et al.*, 2006]. Twenty-five sites (196 site-years) were selected and their specific information were shown in Table 1, and there are 6 PFTs in this study. Cropland were separated by C_3 and C_4 crops. The selected sites are distributed in arid and humid areas, with mean annual precipitation gradients ranging from 380 to 1426 mm (Table 1). Data quality control and screening were addressed following similar criterion and processes reported in other studies [Li *et al.*, 2019; Zhou *et al.*, 2016]. First, data only at daylight with positive net solar radiation, GPP, ET , and VPD can be reserved and defective data were excluded. Second, incoming shortwave radiation less than 50 W m^{-2} and sensible heat flux less than 5 W m^{-2} were excluded to avoid stable boundary layer conditions [Li *et al.*, 2019]. The data of rainy days were excluded by referring to Zhou *et al.*, [2015]. Fourth, we only focus on the data of the growing season period and the division of the growing season period was used the method in Zhou *et al.*, [2016]. Finally, daily values were selected to estimate $T:ET$ only when the day contains at least 10 half-hourly data.

111 3.2 Leaf area index (LAI)

112 In this study, LAI ($\text{m}^2 \text{ m}^{-2}$) of vegetation was estimated using photosynthetically active radiation data, and the estimation was used the
113 method in *Xu et al.*, [2010].

114 4. Results and Discussion

115 4.1 Estimation of WUE_{leaf} and WUE_{eco}

116 The average and long-term WUE_{eco} and WUE_{leaf} of seven PFTs for the 25 sites are shown in Fig. 1. Overall, the spatiotemporal
117 characteristics of WUE_{eco} and WUE_{leaf} for seven PFTs were not consistent, which is consistent with the findings reported by *Yi et al.*
118 [2019]. For example, deciduous broad forests ($4.95 \text{ g C kg}^{-1} \text{ H}_2\text{O}$) and evergreen broad forests ($4.97 \text{ g C kg}^{-1} \text{ H}_2\text{O}$) had the highest WUE_{eco} ,
119 while evergreen needleleaf forests ($11.7 \text{ g C kg}^{-1} \text{ H}_2\text{O}$) and C_4 croplands ($11.3 \text{ g C kg}^{-1} \text{ H}_2\text{O}$) had the highest WUE_{leaf} . The inconsistent
120 trend between WUE_{eco} and WUE_{leaf} reflects the proportion of productive and non-productive water consumption in the water cycle across
121 different PFTs. The results also showed that C_4 croplands had higher WUE_{eco} ($4.0 \text{ g C kg}^{-1} \text{ H}_2\text{O}$) and WUE_{leaf} ($11.3 \text{ g C kg}^{-1} \text{ H}_2\text{O}$) than C_3
122 croplands ($2.8 \text{ g C kg}^{-1} \text{ H}_2\text{O}$ and $5.2 \text{ g C kg}^{-1} \text{ H}_2\text{O}$, respectively). The C_4 leaves have higher WUE than C_3 leaves under same conditions
123 due to the CO_2 concentration mechanism [*Ghannoum*, 2009]. However, C_4 croplands had much higher WUE_{leaf} than its WUE_{eco} and the
124 WUE_{leaf} of C_3 croplands, indicating that C_4 croplands had lower $T:ET$ than C_3 croplands. In addition, the WUE of croplands varies in
125 agricultural systems and is mostly determined by local irrigation and field management practices. For natural vegetation, deciduous broad
126 forests ($4.95 \text{ g C kg}^{-1} \text{ H}_2\text{O}$) and evergreen broad forests ($4.97 \text{ g C kg}^{-1} \text{ H}_2\text{O}$) had higher WUE_{eco} than evergreen needleleaf forests (4.2 g C
127 $\text{kg}^{-1} \text{ H}_2\text{O}$) and grasslands ($2.4 \text{ g C kg}^{-1} \text{ H}_2\text{O}$), which were consistent with the studied model simulation [*Gu et al.*, 2018] and global satellite
128 data [*Huang et al.*, 2017]. Broad forests had higher carbon uptake capacity than evergreen needleleaf forests due to their larger leaf area
129 (Fig. 2). Compared to grasslands, forests had higher WUE_{eco} , which is likely due to their (1) lower E , as their large canopy and tree height
130 reduces the gradient in VPD between the atmosphere and soil surface [*Brutsaert*, 2005] and/or (2) lower T due to the distribution of leaf
131 stomata. In most herbs, stomata exist on both the adaxial and abaxial leaf surfaces, while in many trees, stomata only exist on the adaxial
132 leaf surface [*Taiz and Zeiger*, 2006]. In this study, woody savannas had the lowest WUE_{eco} ($1.8 \text{ g C kg}^{-1} \text{ H}_2\text{O}$) and WUE_{leaf} (4.8 g C kg^{-1}
133 H_2O). The US_SRM and US_Ton sites for woody savannas belonged to arid and semiarid areas; their low WUE_{eco} and WUE_{leaf} values
134 have been attributed to low carbon uptake and large soil evaporation [*Scott and Biederman*, 2017; *Wang et al.*, 2016].

135 Multi-year and interannual variations in WUE_{eco} and WUE_{leaf} for the DE_Gri (GRA), IT_Cpz (EBF), FR_Fon (DBF), US_Ne1 (CRO),
136 US_NR1 (ENF), US_SRM (WSA) sites were estimated daily (Fig. 2). The multi-year variation was small for each site, but inter-sites were
137 large for WUE_{eco} , WUE_{leaf} , and GPP. Similar results were reported in *Nelson et al.* [2020], indicating that $T:ET$ differs more from one site
138 to another than between years for the same site. For all PFTs, the daily variation trend in WUE_{eco} is consistent with that of GPP, but
139 WUE_{leaf} showed high-frequency fluctuations over time. The high-frequency variation of WUE_{leaf} is the main reason for the high variation
140 in the estimated daily $T:ET$. In this study, the stomatal conductance model was used to estimated WUE_{leaf} half-hourly, and g_l is constant
141 when vegetation type and site are determined. Although the variations in ambient atmospheric CO_2 concentration and atmospheric
142 pressure were small on a daily scale, studies have shown that ambient atmospheric CO_2 concentration has a significant effect on WUE_{leaf}
143 [*Knauer et al.*, 2017; *Onoda et al.*, 2009]. The high-frequency variation is mostly attributed to variation in VPD , which is easily affected

144 by daily climate factors, such as light, humidity, and temperature. Similarly, *Yi et al.* [2019] indicated that WUE_{leaf} was most sensitive to
145 variations in VPD , accounting for 86% of influence factors of VPD , soil moisture, and ambient atmospheric CO_2 concentration.

146 4.2 Estimation of $T:ET$

147 The annual mean $T:ET$ varied greatly among 196 site-years with seven PFTs (Fig. 3). Overall, the mean $T:ET$ among the 196 site-years
148 was 0.54 (range 0.4–0.67), which is very close to results (0.57, range 0.5–0.64) on a global terrestrial scale [*Wei et al.*, 2017]. In this study,
149 deciduous broad forests had the highest average $T:ET$ (0.67, range 0.50–0.78), with the trend and magnitude similar to the results of a
150 global meta-analysis (0.67) [*Schlesinger and Jasechko*, 2014], transpiration estimation algorithm (TEA) method (0.7) [*Nelson et al.*, 2018]
151 and modeling study (0.6) [*Gu et al.*, 2018]. However, *Zhou et al.* [2016] reported that deciduous broad forests (0.52 ± 0.08) had the lowest
152 $T:ET$ among all PFTs, being lower than those reported by *Schlesinger and Jasechko* [2014] in a global synthesis of 81 studies on ET
153 partitioning. The reason for underestimating $T:ET$ of deciduous broad forests was explained in *Zhou et al.* [2016]. The estimated annual
154 $T:ET$ for ENF (mean 0.46) is lower than that reported (0.55) by *Schlesinger and Jasechko* [2014] but the values for each site in this study
155 (0.34–0.61) were all within the range for ENF (0.3–0.7). The reason for underestimation may be due to parameter g_l variation among the
156 four evergreen needleleaf forest sites. Our study had two sites (CA_NS3 and US_NR1) with small g_l values (2.9 and 2.6, respectively). A
157 low g_l would be accompanied by high WUE_{leaf} , resulting in low $T:ET$. The annual $T:ET$ of grasslands varied little (0.52, range 0.38–0.67),
158 compared with 0.56 ± 0.05 [*Zhou et al.*, 2016] and 0.57 ± 0.19 [*Schlesinger and Jasechko*, 2014], because we included two grasslands
159 located in arid areas and their perennial average $T:ET$ were 0.45 and 0.56 for US_Wkg and US_SRG sites, respectively. Thus, the
160 estimated $T:ET$ by using our method is very close to that reported in the published researches for deciduous broad forests, evergreen
161 needleleaf forest, and grasslands. Moreover, this study showed that C_3 croplands had higher $T:ET$ (0.65) than C_4 croplands (0.48), which is
162 consistent with those for wheat (C_3) and maize (C_4) using ET partitioning methods with a two-source model or isotope approach [*Wei et*
163 *al.*, 2018]. However, the ET partitioning method used by *Zhou et al.* [2016] showed that C_4 croplands had higher $T:ET$ (0.69) than C_3
164 croplands (0.62). The reason why $T:ET$ differs in agriculture systems may be due to human factors, such as irrigation, mulching or
165 fertilizer. In addition, two woody savannas (US_Ton and US_SRM) located in semiarid and arid sites had an average $T:ET$ of 0.41 (range
166 0.39–0.43), which is consistent with that (0.48 ± 0.12) reported by *Schlesinger and Jasechko* [2014].

167 Seasonal and interannual variations in $T:ET$ for DE_Gri (GRA), IT_Cpz (EBF), FR_Fon (DBF), US_Ne1 (CRO), US_NR1 (ENF), and
168 US_SRM (WSA) were estimated daily (Fig. 4). Overall trends in seasonal $T:ET$ characteristics for six PFTs were similar to those in the
169 TEA and uWUE methods [*Nelson et al.*, 2020]. We selected three sites and years (DE_Gri, US_Ne1, and US_NR1) that were also used in
170 a novel ET partitioning method based on soil and canopy conductances [*Li et al.*, 2019] and compared their $T:ET$ trends and characteristics
171 within and between years. The three sites showed consistent trends in $T:ET$ characteristics, albeit slightly higher in *Li et al.* [2019] because
172 they did not include evaporation from canopy interception.

173 The seasonal patterns of $T:ET$ differed for the six PFTs. For grasslands at the DE_Gri site, $T:ET$ increased to 0.8 over time but rapidly
174 declined with the harvest of herbage, which occurred several times a year. For croplands, maize grew faster than the other natural
175 vegetation, and T rapidly reached its maximum. The daily variation in $T:ET$ at the US_Ne1 site followed a single-peak pattern within a
176 growing season. For evergreen broad forests at the IT_Cpz site, the $T:ET$ was mostly above 0.4, with an unobvious peak and large
177 variation in daily $T:ET$ during the growing season. For evergreen needleleaf forests at the US_NR1 site, $T:ET$ was below 0.2 in the early

growing season, and showed high-frequency fluctuations, reaching a peak value of about 0.54 during the growing season. The reasons for the high-frequency fluctuations in $T:ET$ across plant types are provided in section 4.1. Moreover, large daily variations in $T:ET$ can be influenced by other biotic/abiotic factors, for example, soil water content, vegetation coverage, and plant phenology [Berkelhammer *et al.*, 2016; Gao *et al.*, 2019; Oishi *et al.*, 2008].

4.3 Evaluation of the ET partitioning method

The relationship between mean growing season LAI and annual $T:ET$ shows a significant R^2 of linear regression ($p < 0.01$) (Fig. 5). This result is similar to that of Li *et al.* [2019] and Gu *et al.* [2018], but with improved correlation coefficients. However, the low R^2 of 0.26 suggests the mean growing season LAI explains small (26%) variations in mean annual $T:ET$ across biomes, indicating that other biological and non-biological factors affect ET variation in addition to LAI in global ecosystem. Ambient atmospheric conditions (e.g., VPD and precipitation), soil moisture, and plant phenology are important factors affecting high $T:ET$ variation at daily or seasonal scales [Good *et al.*, 2014; Scott and Biederman, 2017; Zhao *et al.*, 2018]. Another reason is that the mean LAI values of the growing season but not the whole growing season, including the early growing season, were selected in this study. Scott and Biederman [2017] indicated that a larger percentage of $T:ET$ variation is explained by a commonly used power function with LAI rather than a linear relationship early in the growing season. Overall, the correlation between LAI and $T:ET$ further demonstrates the reliability of this new ET partitioning method.

The mean annual $T:ET$ across 196 site-years was compared with previously published estimates using four methods (isotope, modeling, flux data, and meta-analysis). Inter-comparison of various ET partitioning methods showed a spread in magnitudes of $T:ET$ from 0.38 to 0.75 (Fig. 6). The overall $T:ET$ across the 196 site-years in this study was 0.54 (range 0.40–0.67). On the whole, the average $T:ET$ values were slightly lower than studies, although the ranges of $T:ET$ for several PFTs were all within the global given reasonable interval. There may be several possible reasons for this. Firstly, many published studies ignore the rainfall evaporation intercepted by the plant canopy, which would overestimate $T:ET$ [Baldocchi, 2014; Li *et al.*, 2019]. Moreover, measurement of E and T separately will overestimate the proportion of T in ET [Gu *et al.*, 2018]. Finally, different research regions (such as those that focus only on non-limited water regions) and various scales may influence the results of $T:ET$; for example, the isotope-based approach overestimates $T:ET$, which is constrained by hydrologic decoupling [Jasechko *et al.*, 2013]. Wang *et al.* [2014] indicated that large variabilities and observation uncertainty across sites could generate a large $T:ET$ range. Similarly, Nelson *et al.* [2020] reported a 0.45–0.77 spread in magnitude of $T:ET$ using three ET partitioning methods based on fluxnet data, despite plausible and qualitatively consistent T and $T:ET$ patterns. Therefore, various ET partitioning methods and measurement techniques should be used to reduce uncertainties in T estimation [Rafi *et al.*, 2019].

We used three ET partitioning methods to compare monthly and growing season $T:ET$ in two arid grasslands (Table 2). One of the methods—proposed by Scott and Bieberman [2017]—is a reliable method for estimating $T:ET$ at water-limited sites. In general, the three ET partitioning methods produced consistent $T:ET$ at the seasonal scale. The monthly and whole growing season $T:ET$ values from our method were similar to those of Scott and Bieberman [2017], with small differences of +1.8% and –0.3% for the US_SRG and US_Wkg sites, respectively. Unsurprisingly, Zhou *et al.*'s [2015] method overestimated growingseason $T:ET$ at the US_Wkg site by 37.8%, relative to that in Scott and Bieberman [2017]. Moreover, we compared the daily $T:ET$ estimation of our method and Zhou *et al.*'s method at two arid grassland sites (Figs 7 and 8). The Pearson's correlation coefficients of $T:ET$ on a daily scale for the two methods were 0.95 and 0.85 at the US_SRG and US_Wkg sites, respectively. The overall $T:ET$ trends on a daily scale were consistent for the two methods,

but Zhou et al.'s method produced higher $T:ET$ estimation values than our method because their assumption that T equals ET throughout the growing season is not valid for arid and vegetation-sparse areas. Our ET partitioning method is not restricted to arid and vegetation-sparse areas and improved the $T:ET$ estimation accuracy in water-limited regions.

4.4 Implications and limitations

This study developed a novel method of ET partitioning based on the relationship between WUE_{leaf} and WUE_{eco} by using easily available eddy-covariance data. This method has several advantages over other ET partitioning techniques that use eddy-covariance measurements. Unlike Zhou et al. [2016] and Scott and Biederman [2017], our novel method is applicable to all regions and PFTs and not limited by local soil moisture in dry or wet areas. Unlike Li et al. [2019], our method incorporates evaporation from canopy interception. Importantly, our method can easily estimate $T:ET$ at spatiotemporal scales using meaningful ecological interpretations of WUE_{eco} and WUE_{leaf} through reliable theoretical derivation. Moreover, this method helps understand the upscaling or downscaling of WUEs at different scales.

However, there are several limitations to this study. First, uncertainty in GPP and ET estimations would result in some uncertainty in WUE_{eco} and hence $T:ET$. Second, the lack of direct observations of E and T at the flux tower sites to validate our partitioning method. Although the comparison with other ET partitioning methods lend considerable support for our method, we also find a few inconsistent results and there is no consistent conclusion as to which ET partitioning method or measurement technique is the most accurate [Kool et al., 2014]. Thirdly, we assumed that WUE_{leaf} can be used to approximate $iWUE_{eco}$, which may not be true at sites with mixed vegetation types with distinct WUE_{leaf} or heterogeneous environmental conditions, such as VPD and light density, leading to large variations in WUE_{leaf} within the site. Finally, the g_l values were determined from different PFTs and land cover maps. As g_l is predicted to change with moisture index and temperature [Lin et al., 2015], using a constant g_l value at each site may lead to some uncertainty in WUE_{leaf} and hence $T:ET$. The reason for not considering soil moisture and temperature is to retain a simple and available parameter in the $T:ET$ model.

5. Conclusion

Half-hourly flux data from 196 eddy covariance site-years was used to develop a novel ET partitioning method coupled with WUE at various scales. According to WUE based on water consumption, $T:ET$ equals the ratio of WUE_{eco} to WUE_{leaf} numerically through reasonable derivation, and WUE_{eco} and WUE_{leaf} can be easily calculated from available data provided by flux sites. The spatiotemporal characteristics of WUE_{eco} and WUE_{leaf} for seven PFTs were not consistent, which reflects the proportion of productive and non-productive water consumption across PFTs. For natural vegetation, deciduous broad forests and evergreen broad forests had higher WUE_{eco} than evergreen needleleaf forests and grasslands, but evergreen needleleaf forests had the highest WUE_{leaf} . For agricultural systems, C_4 croplands had higher WUE_{eco} and WUE_{leaf} than C_3 croplands due to the CO_2 -concentrating mechanism. Moreover, $T:ET$ characteristics varied among PFTs. Deciduous broadleaf forests had the highest mean annual $T:ET$ (0.67), followed by evergreen broadleaf forests (0.63), grasslands (0.52), evergreen needleleaf forests (0.46), and woody savannas (0.41). The C_3 croplands (0.65) had higher $T:ET$ ratios than C_4 croplands (0.48). We also examined the feasibility and reliability of our ET partitioning method—the trends and magnitudes in $T:ET$ were consistent with those of other ET partitioning methods and the known effect from LAI. Furthermore, our method improved $T:ET$ estimation accuracy in vegetation-sparse and water-limited areas. Thus, this method is sound in principle, in addition, it is easy to used and widely implemented for all regions and plant types using data from global flux tower networks. Moreover, this method provides new

insights into the conversion of WUE at different scales.

Acknowledgments

Data for these flux sites are archived at <https://fluxnet.org/sites/site-summary> and freely obtainable. We acknowledge funding from the National Key Research and Development Program (2016YFC0400204), the National Natural Science Foundation of China (41571506, 41771316, 51579212), the Shaanxi Innovative Research Team for Key Science and Technology (2017KCT-15), the '111'Project (B12007), and CAS "Youth Scholar of West China" Program (XAB2018A04).

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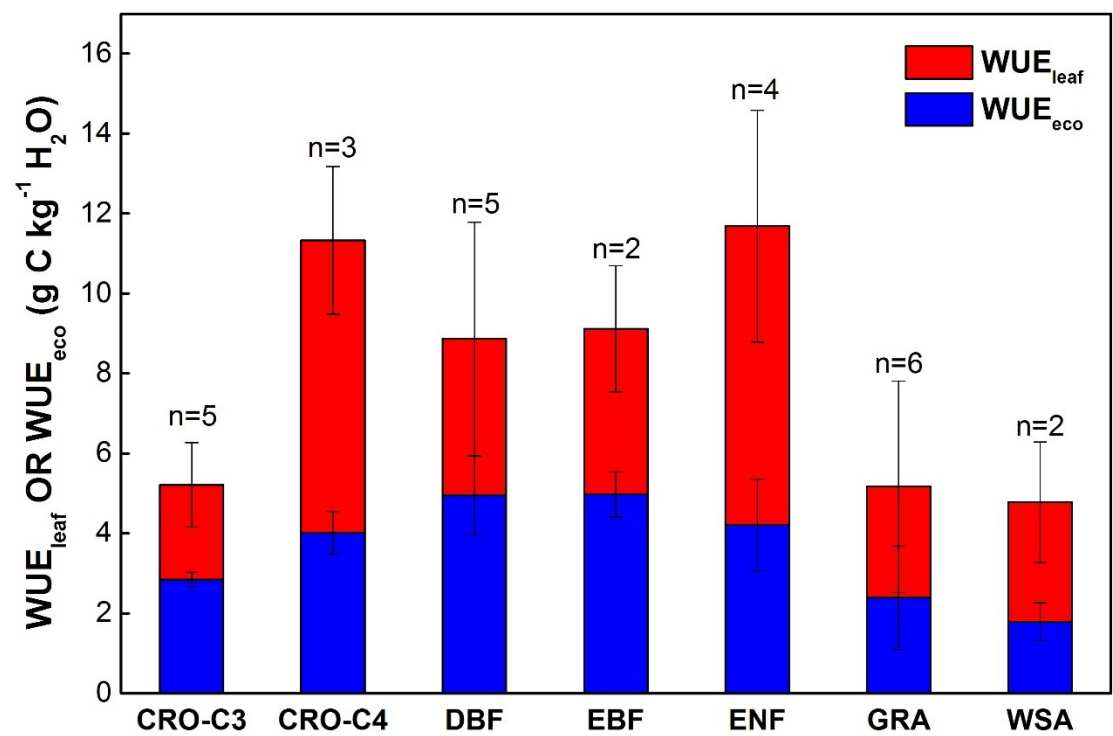
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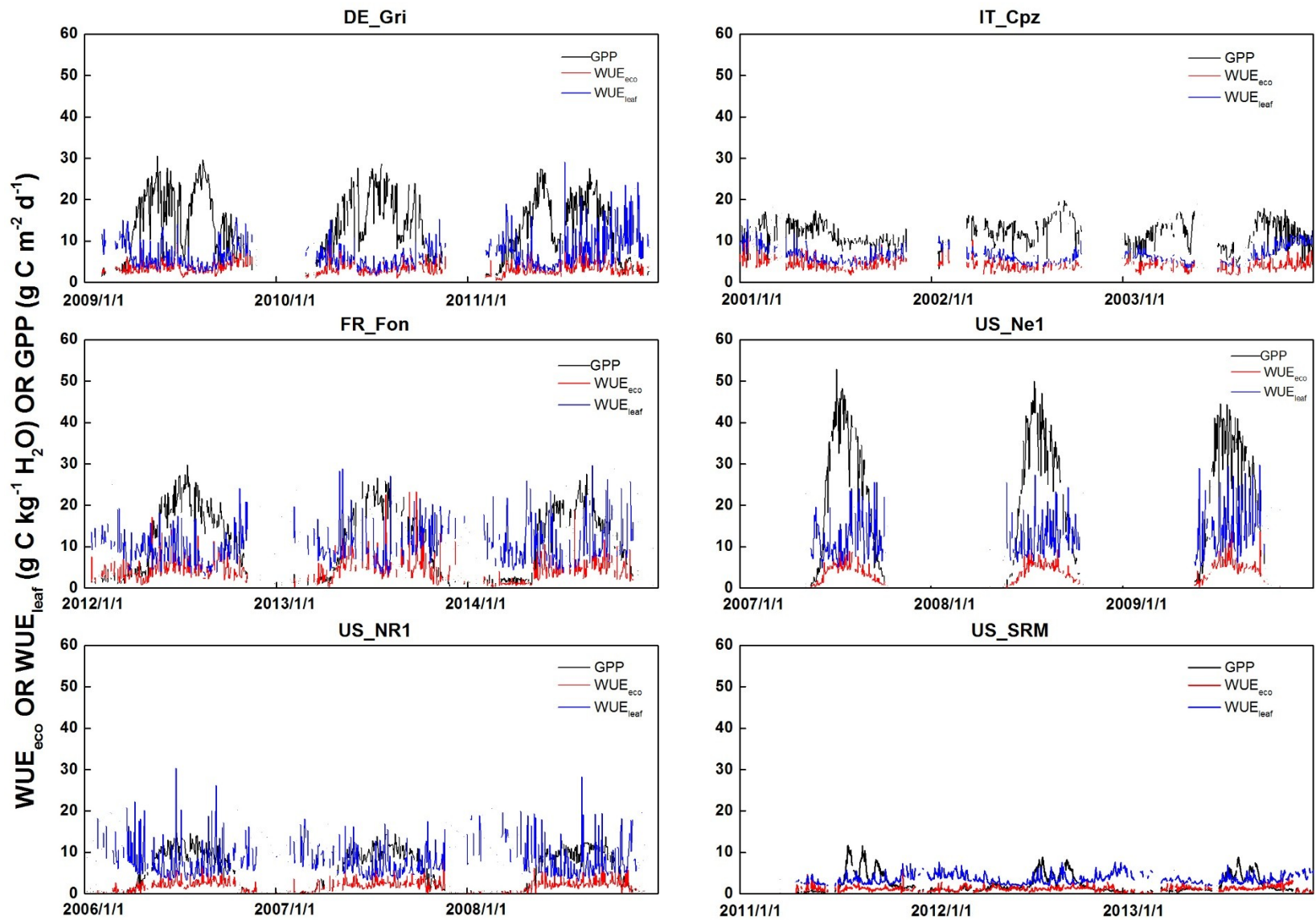
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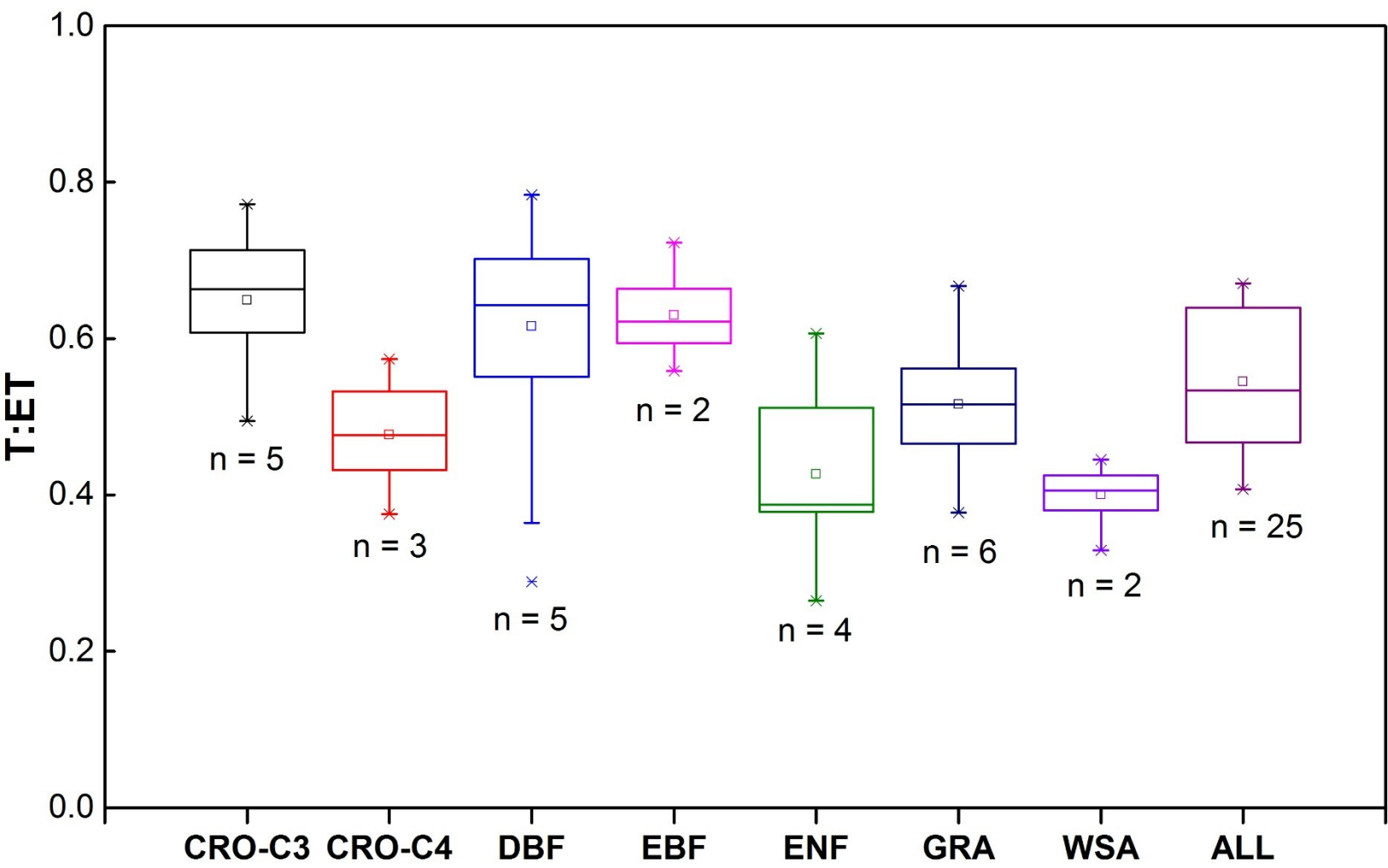
366 **Figure 1.** Distribution of mean ecosystem (WUE_{eco}) and leaf water use efficiency (WUE_{leaf}) for 25 sites across seven plant functional types
 367 (PFTs). The n value above each bar represents the sample sizes of sites. The bars represent standard deviations of WUE_{eco} and WUE_{leaf} for
 368 each PFT.



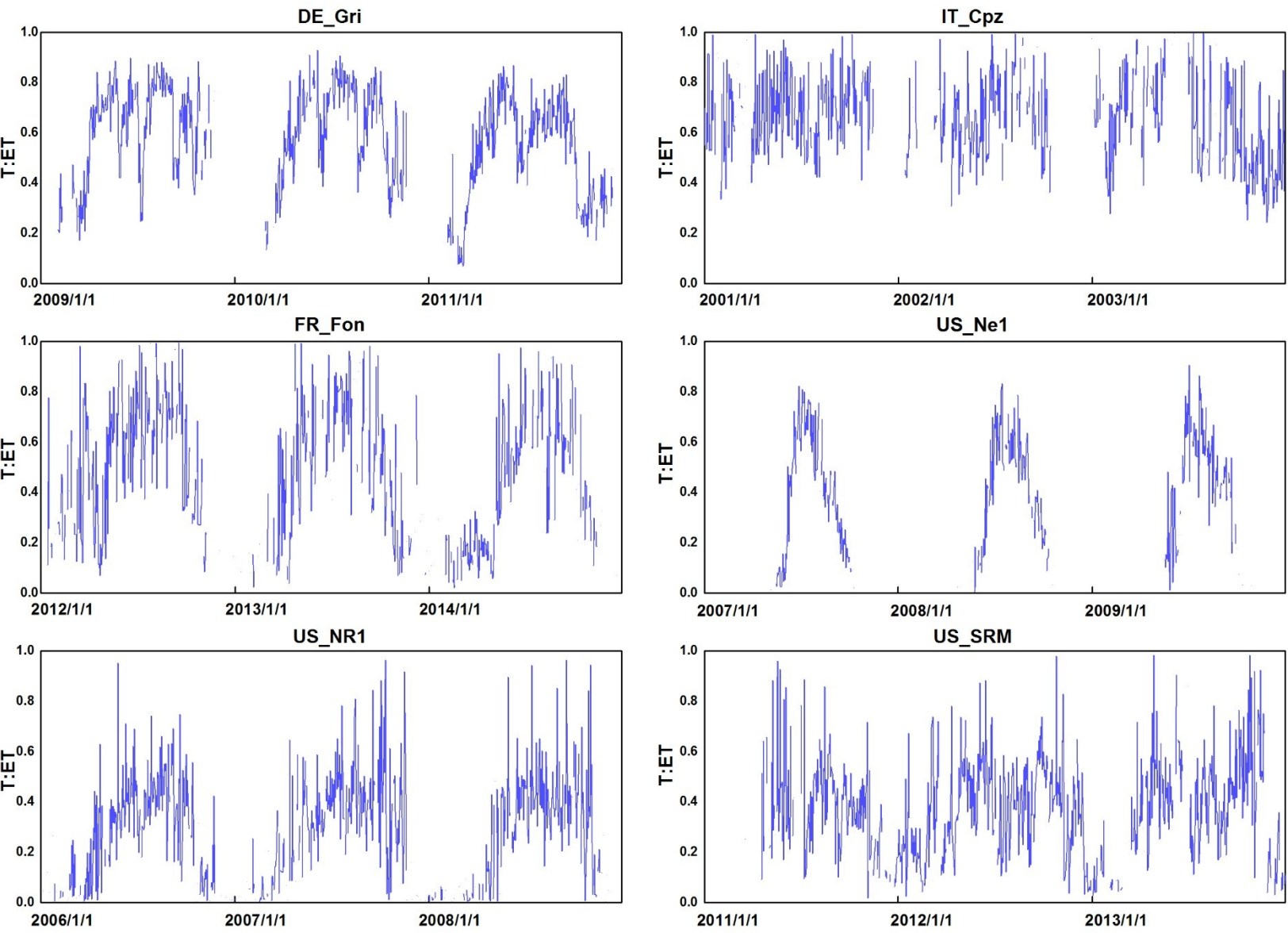
371 **Figure 2.** Seasonal and interannual variation of gross primary productivity (GPP), ecosystem (WUE_{eco}), and leaf water use efficiency
 372 (WUE_{leaf}) at the daily scale for the DE_Gri (GRA), IT_Cpz (EBF), FR_Fon (DBF), US_Ne1 (CRO), US_NR1 (ENF), and US_SRM
 373 (WSA) sites.



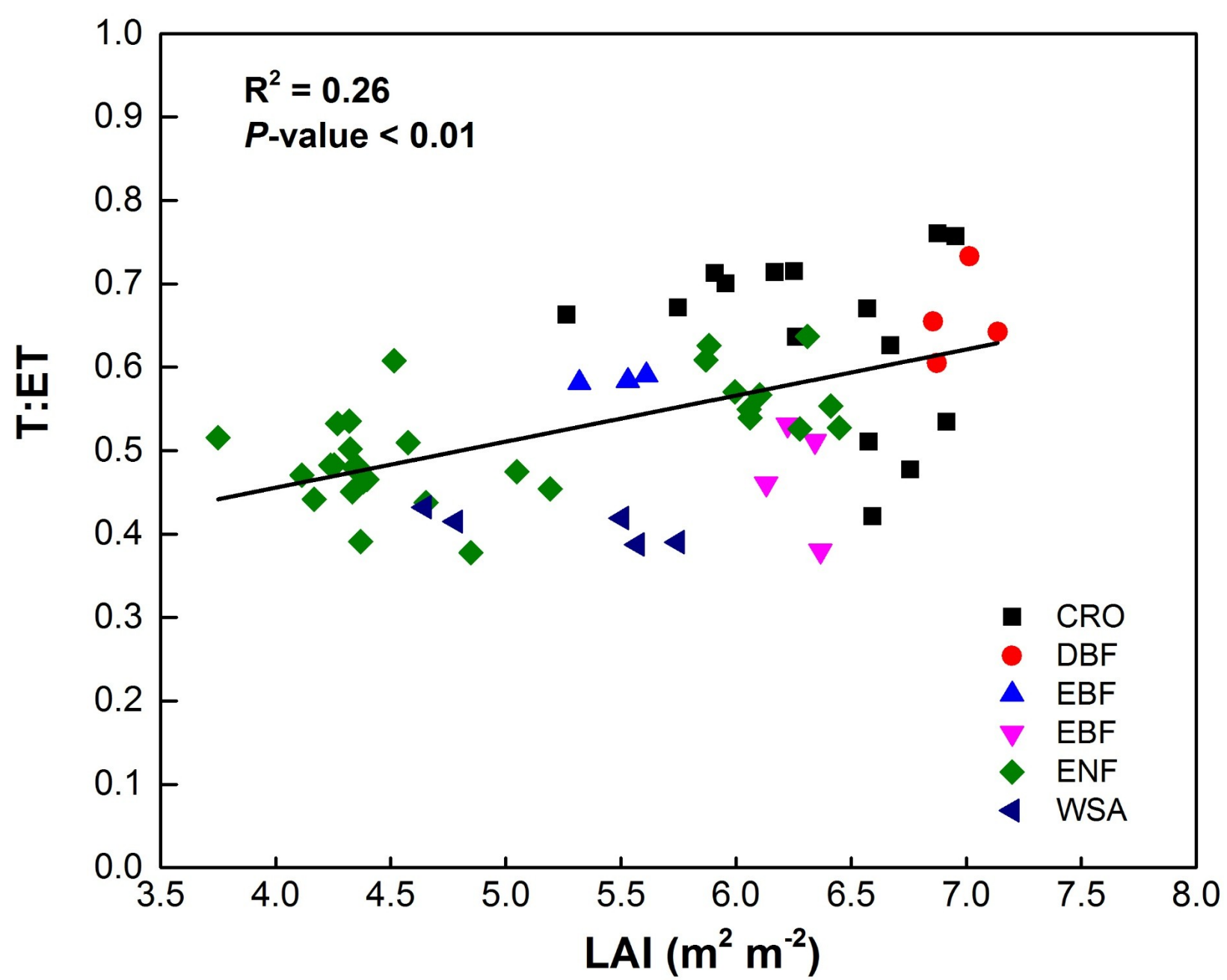
376 **Figure 3.** Characteristics of $T:ET$ of various plant functional types (PFTs) for the 196 site-years. The square and solid lines in the boxes
 377 represent the average and median values, the upper and lower error bars represent the quantiles of 75th and 25th. The n values below boxes
 378 represents the sample sizes of sites.



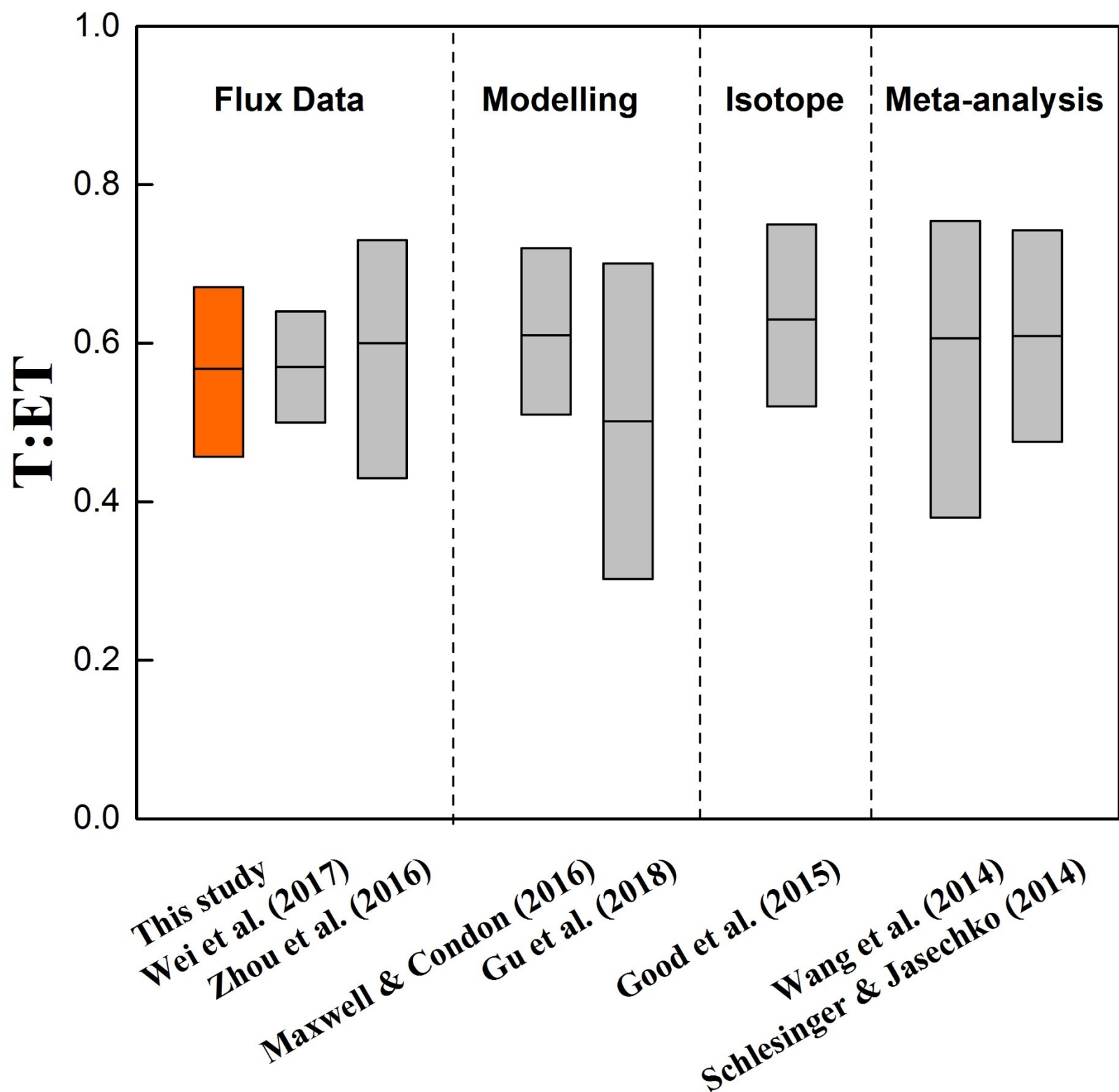
381 **Figure 4.** Interannual and seasonal and $T:ET$ characteristics daily for the DE_Gri (GRA), IT_Cpz (EBF), FR_Fon (DBF), US_Ne1 (CRO),
 382 US_NR1 (ENF), and US_SRM (WSA) sites.



385 **Figure 5.** The relationship between mean growing season leaf area index (LAI) and mean annual $T:ET$.

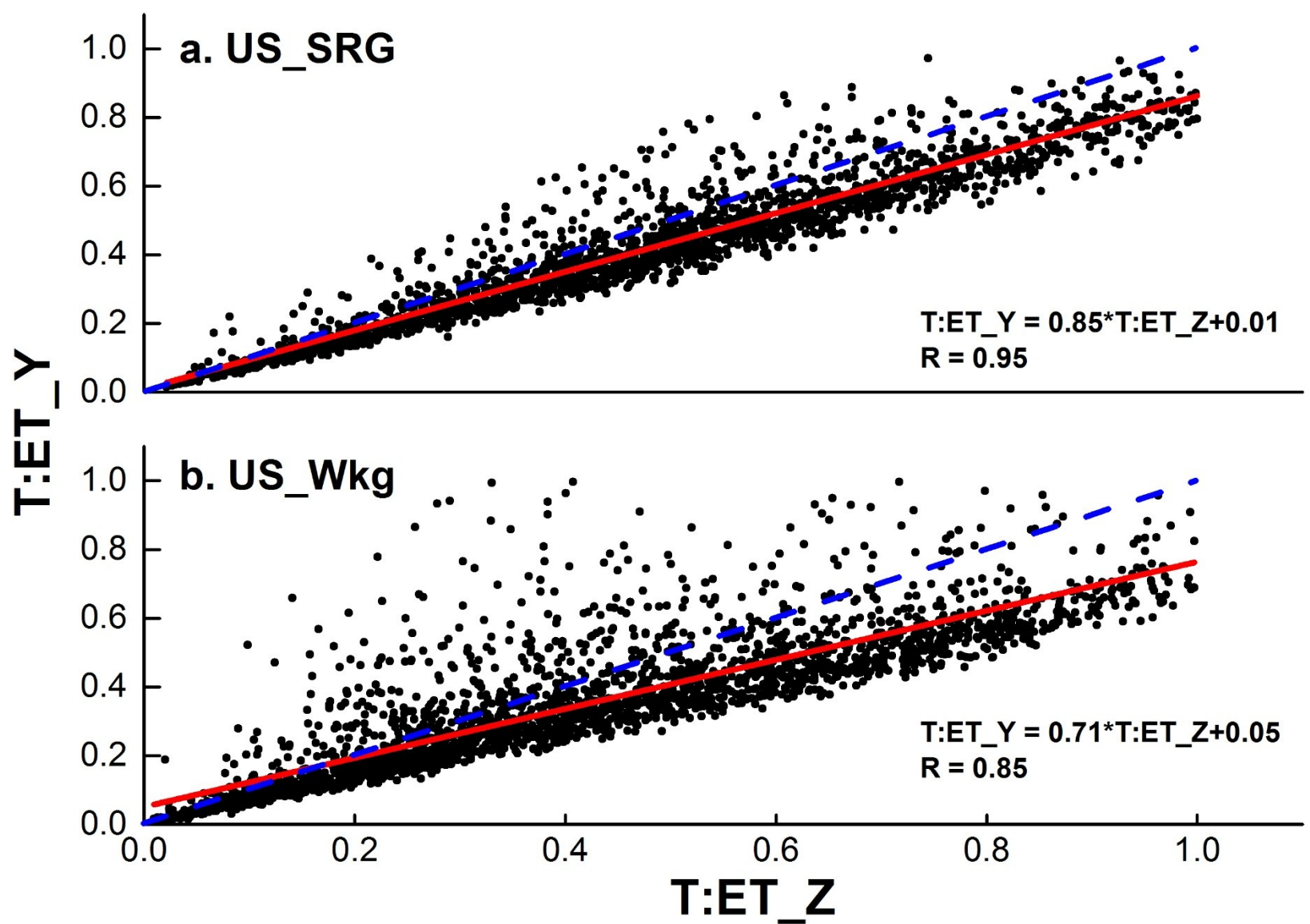


387 **Figure 6.** The comparison of $T:ET$ estimation among various methods. The rectangle represents the ranges of $T:ET$ values and the value
388 corresponding to the horizontal line in the middle represents the average value of $T:ET$ reported in the published literature.



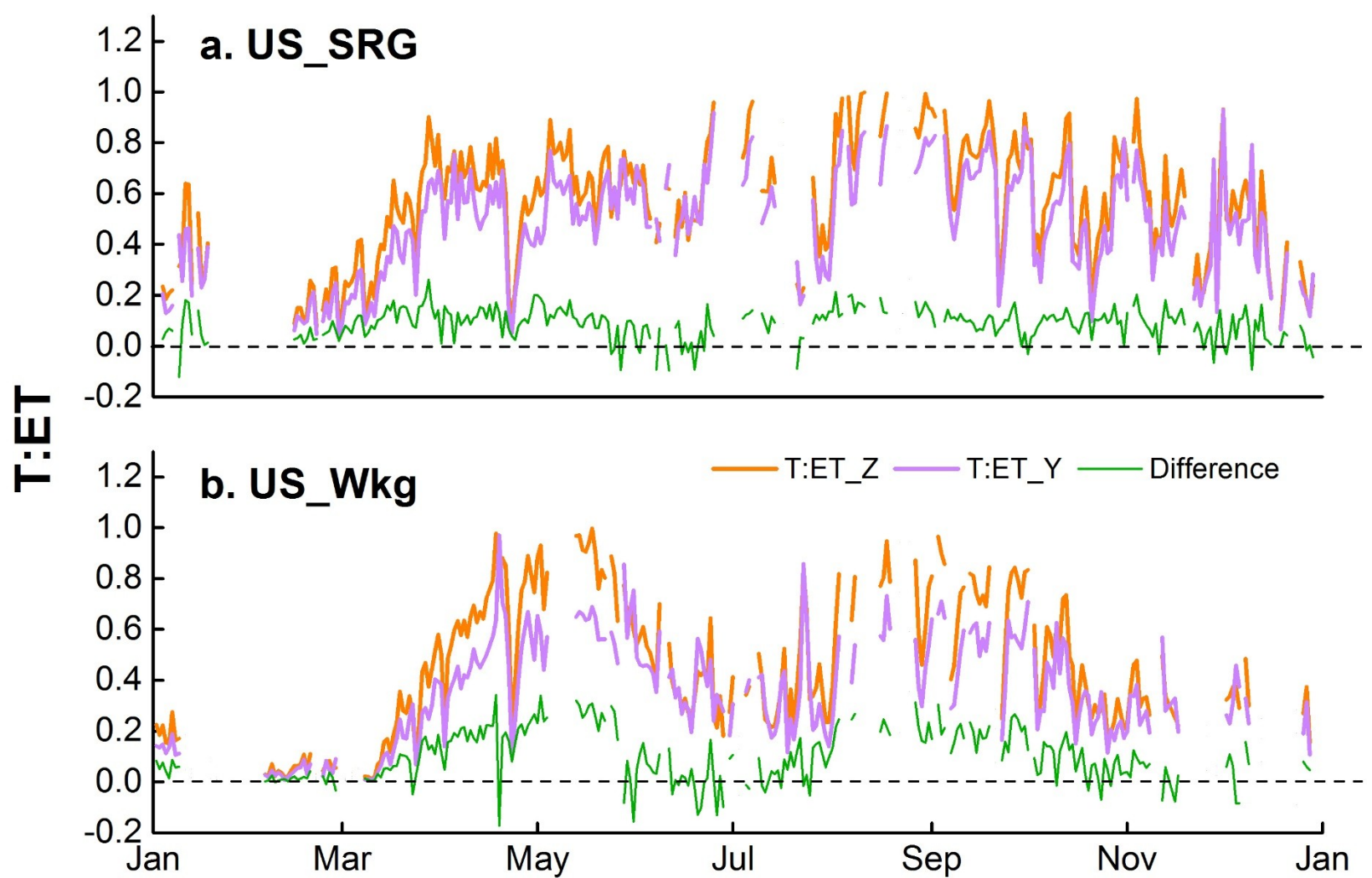
389

390 **Figure 7.** Comparison of $T:ET$ estimations on a daily scale between two arid grassland sites in this study ($T:ET_Y$) and Zhou's method
 391 ($T:ET_Z$). The points located at the (a) US_SRG and (b) US_Wkg sites are 1,987 and 2,591, respectively. The blue dotted line represents
 392 the 1:1 trend line. The red line represents the linear equation estimated using orthogonal-least-squares regression.



393

394 **Figure 8.** Comparison of daily *T:ET* characteristics in 2010 between two arid grassland sites in this study (*T:ET_Y*) and Zhou’s method
 395 (*T:ET_Z*). The difference is expressed as *T:ET* value estimated by Zhou’s method minus *T:ET* value estimated by this method.



396

| SITE CODE | LAT | LON | Elevation (m) | PFT | MAT (°C) | MAP (mm) | g_l (KPa ^{0.5}) | YEARS USED | REFERENCE |
|--------------|-------|---------|------------------|-----|-------------|-------------|---|------------|-----------------------------------|
| CA_NS3 | 55.91 | -98.38 | 260 | ENF | -2.87 | 502.22 | 2.9 | 2001-2006 | (Bond-Lamberty et al., 2004) |
| CA_Oas | 53.62 | -106.19 | 530 | DBF | 0.34 | 428.53 | 5.1 | 1997-2010 | (Barr et al., 2004) |
| DE_Gri | 50.95 | 13.51 | 385 | GRA | 7.8 | 901 | 4.9 | 2004-2014 | (Prescher <i>et al.</i> , 2010) |
| DE_Hai | 51.08 | 10.45 | 430 | DBF | 8.3 | 720 | 5.5 | 2000-2009 | (Knohl <i>et al.</i> , 2003) |
| DE_Obe | 50.79 | 13.72 | 734 | ENF | 5.5 | 996 | 5.1 | 2008-2014 | (Zimmermann <i>et al.</i> , 2006) |
| DE_RuS | 50.86 | 6.45 | 102 | CRO | 10 | 700 | 5.4 | 2011-2014 | (Mauder et al., 2013) |
| DE_Seh | 50.87 | 6.45 | 103 | CRO | 9.9 | 693 | 5.4 | 2007-2010 | (Schmidt et al., 2012) |
| FR_Fon | 48.48 | 2.78 | 103 | DBF | 10.2 | 720 | 4.7 | 2012-2014 | (Bazot et al., 2013) |
| FR_Pue | 43.74 | 3.60 | 270 | EBF | 13.5 | 883 | 3.8 | 2001-2014 | (Rambal et al., 2004) |
| IT_Cpz | 41.71 | 12.38 | 68 | EBF | 15.6 | 780 | 3.9 | 2000-2007 | (Garbulsky et al., 2008) |
| NL_Loo | 52.17 | 5.74 | 25 | ENF | 9.8 | 786 | 4.7 | 1997-2014 | (Gioli <i>et al.</i> , 2004) |
| US_ARb | 35.55 | -98.04 | 424 | GRA | / | / | 3.5 | 2005-2006 | (Schmidt et al., 2011) |
| US_ARc | 35.55 | -98.04 | 424 | GRA | / | / | 3.5 | 2005 | (Schmidt et al., 2011) |
| US_CRT | 41.63 | -83.35 | 180 | CRO | 10.1 | 849 | 5.3 | 2011-2013 | (Chu et al., 2016) |
| US_Goo | 34.25 | -89.87 | 87 | GRA | 15.89 | 1425.77 | 4.2 | 2002-2006 | (Benjamin et al., 2017) |
| US_Ha1 | 42.54 | -72.17 | 340 | DBF | 6.62 | 1071 | 4.6 | 1994-2012 | (Barford et al., 2001) |
| US_Ne1 | 41.17 | -96.48 | 361 | CRO | 10.07 | 790.37 | 1.6 | 2002-2010 | (Suyker <i>et al.</i> , 2004) |
| US_Ne2 | 41.16 | -96.47 | 362 | CRO | 10.08 | 788.89 | 4.0(C ₃)/1.6(C ₄) | 2001-2012 | (Suyker <i>et al.</i> , 2004) |
| US_Ne3 | 41.18 | -96.44 | 363 | CRO | 10.11 | 783.68 | 4.0(C ₃)/1.6(C ₄) | 2001-2012 | (Suyker <i>et al.</i> , 2004) |
| US_NR1 | 40.03 | -105.55 | 3050 | ENF | 1.5 | 800 | 2.6 | 2006-2008 | (Arain et al., 2005) |
| US_SRG | 31.79 | -110.83 | 1291 | GRA | 17 | 420 | 3.7 | 2008-2014 | (Biederman <i>et al.</i> , 2016) |
| US_SRM | 31.82 | -110.87 | 1120 | WSA | 17.92 | 380 | 3.7 | 2011-2013 | (Scott et al, 2010) |
| US_Ton | 38.43 | -120.97 | 177 | WSA | 15.8 | 559 | 3.3 | 2003-2005 | (Fisher <i>et al.</i> , 2007) |
| US_UMB | 45.56 | -84.71 | 234 | DBF | 5.83 | 803 | 4.7 | 2000-2014 | (Curtis et al., 2007) |
| US_Wkg | 31.74 | -109.94 | 1531 | GRA | 15.64 | 407 | 3.9 | 2004-2012 | (Moran <i>et al.</i> , 2009) |

399 Note: Site code, Latitude (LAT, °), longitude (LON, °), elevation (m), plant functional type (PFT), MAP (mean annual precipitation), MAT
400 (mean annual temperature), g_l (parameter of the Ball stomatal conductance model), years used and corresponding reference are listed. The
401 PFTs include croplands (CRO), deciduous broadleaf forests (DBF), evergreen broadleaf forests (EBF), evergreen needleleaf forests (ENF),
402 grasslands (GRA) and woody savannas (WSA).

404 **Table 2.** Comparison of monthly and growing season *T:ET* among Scott and Biederman, 2017, Zhou *et al.*, 2015 and this study at the
405 US_SRG (2008-2015) and US_Wkg (2004-2015) sites.

| Site | Study | July | August | September | October | Growing Season | Difference (%) |
|--------|---------------------------|------|--------|-----------|---------|----------------|----------------|
| US_SRG | Scott and Biederman, 2017 | 0.54 | 0.62 | 0.54 | 0.51 | 0.55 | / |
| | This study | 0.55 | 0.63 | 0.55 | 0.52 | 0.56 | + 1.82 |
| | Zhou et al., 2015 | 0.64 | 0.73 | 0.6 | 0.6 | 0.64 | + 14.3 |
| US_Wkg | Scott and Biederman, 2017 | 0.26 | 0.58 | 0.54 | 0.45 | 0.46 | / |
| | This study | 0.35 | 0.52 | 0.48 | 0.46 | 0.45 | - 0.3 |
| | Zhou et al., 2015 | 0.48 | 0.66 | 0.66 | 0.69 | 0.62 | + 37.8 |

406

407 Note: The two grassland sites are in water-limited regions where mean annual precipitation is 420 and 407 mm. The difference (%) is

408 expressed as the percent of the *T:ET* difference values relative to the results of Scott and Biederman (2017).

409