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Is the reputation of *Eucalyptus* plantations for using more water than *Pinus* plantations justified?

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Abstract. The effect of *Eucalyptus* plantations on water balance is thought to be more severe than for commercial alternatives such as *Pinus* species. Although this perception is firmly entrenched, even in the scientific community, only four direct comparisons of the effect on the water balance of a *Eucalyptus* species and a commercial alternative have been published. One of these, from South Africa, showed that *Eucalyptus grandis* caused a larger and more rapid reduction in streamflow than *Pinus patula*. The other three, one in South Australia and two in Chile, did not find any significant difference between the annual evapotranspiration of *E. globulus* and *P. radiata* after canopy closure.

While direct comparisons are few, there are at least 57 published estimates of annual evapotranspiration of either the *Eucalyptus* or *Pinus* species. This paper presents a metaanalysis of these published data. Zhang et al. (2004) fitted a relationship between the vegetation evaporation efficiency and the climate wetness index to published data from catchment studies and proposed this approach for comparing land uses. We fitted this model to the published data for *Eucalyptus* and *Pinus* and found that the single parameter of this

model did not differ significantly between the two genera $(p = 0.48)$. This was also the case for all parameters of an exponential relationship between evapotranspiration and rainfall ($p = 0.589$) and a linear relationship between the vegetation evaporation index and rainfall ($p = 0.155$).

These results provide strong evidence that, for a given climate wetness index, the two genera have similar annual water use. The residuals compared to the model of Zhang et al. (2004) were significantly correlated with soil depth for *Eucalyptus*, but this was not the case for *Pinus*. For *Eucalyptus*, the model overestimates the vegetation evaporation efficiency on deep soils and underestimates the vegetation evaporation efficiency on shallow soils.

1 Introduction

There are now more than 23 Mha of *Eucalyptus* plantations in the temperate and tropical zones of the world (Keenan et al., 2015; MacDicken et al., 2016). These plantations extend from near the Equator to approximately 43◦ of latitude north and south and play an important and growing role in minimising the gap between global demand for wood products and the supply (Kanninen, 2010). These *Eucalyptus* plantations are mostly established in seasonally dry climate zones (dry tropics, sub-tropics, and Mediterranean climate types). This, and the reputation of *Eucalyptus* for high rates of water use when compared to alternatives, means that wherever large-scale planting of *Eucalyptus* has occurred, it has been associated with concern, debate, and often protest about the effect of these plantations on the security of water supply (Albaugh et al., 2013). Afforestation with *Pinus* and other genera has also resulted in concern about changes in local hydrology (Huber and Iroumé, 2001; Little et al., 2009) but has not been associated with the same level of polemic or controversy as the planting of *Eucalyptus*.

In 2010, plantations managed for wood production occupied a total land area of 109 Mha (Kanninen, 2010). Approximately 35 % of these plantations were of *Pinus* species, while 10 % were *Eucalyptus* (Kanninen, 2010). The annual increase in production plantations between 2010 and 2015 was 1.2 %. During this time, the total area of *Pinus* plantations remained virtually unchanged, and much of the global increase was in either *Eucalyptus* plantations or other short-rotation options such as *Acacia* (Payn et al., 2015). The global trends in plantations are towards *Eucalyptus* or species managed on short rotations to grow pulp or biomass for energy. While these global trends are important, the conflict associated with the establishment of *Eucalyptus* plantations and the potential for reduced water availability manifests locally. In South Africa and South Australia, these concerns have resulted in legislation to either regulate water use (Greenwood, 2013) or planting (Albaugh et al., 2013). The effects of *Eucalyptus* on water are currently being actively debated in Chile, where Arauco SA (the largest plantation grower in Chile and the second-largest pulp producer in the world) plan to replace approximately 250 000 ha of *P. radiata* plantations with *Eucalyptus*. In China, regional governments are supporting research to investigate the water benefits of mixed plantings of local species with *Eucalyptus*. It is also likely that the global goal of reduced $CO₂$ emissions will intensify the debate about *Eucalyptus* water use. Given the dominance of the global plantation estates by species of *Pinus* and *Eucalyptus*, and the direct substitution of *Pinus* with *Eucalyptus*, a quantitative comparison between the water use characteristics of these two genera is timely.

The evidence that plantations use more water than grasslands or dryland crops is very strong (Zhang et al., 2001; Zhang, 2004). Similarly, there is evidence that plantations use more water, and therefore generate less streamflow, than natural forest in Chile (Huber et al., 2008), Brazil (Almeida et al., 2007; Meinzer et al., 1999) and Spain (Rodriguez Suarez et al., 2014). The magnitude of the difference between plantations and natural forest is less than that observed between plantations and annual pastures (Zhang et al., 2004).

While there is a perception that *Eucalyptus* use more water than alternative commercial plantation options such as *Pinus*, three of four published comparisons of the water use (defined as evapotranspiration) reported no difference between the water use of species from these two genera. The evidence for higher rates of water use by *Eucalyptus* is mostly from South Africa, where, in a paired catchment study, Scott and Lesch (1997) showed that, at least in the early stages of growth, *Eucalyptus grandis W. Hill.* used up to 92 mm more water per year than *Pinus patula Schiede ex Schltdl. et Cham*. In another direct comparison of the water use of a *Pinus* and *Eucalyptus* species in plantations, Benyon et al. (2006) found that the annual water use of plantations of *E. globulus Labill.* and *P. radiata D. Don.*, with or without access to shallow fresh groundwater, were not significantly different. Recent stand- and catchment-scale comparisons of *P. radiata* and *E. globulus* in central Chile have not observed significant differences between the average annual water use of *P. radiata* and *E. globulus* (Iroumé et al., 2021; White et al., 2021). Given these contrasting results, and the trend towards more planting of *Eucalyptus*, it is important to understand when and why differences might occur in the water balance of *Pinus* and *Eucalyptus* plantations.

Studies in Brazil (Lima et al., 1990) and Tasmania, Australia (Honeysett et al., 1996), have shown that, when planted in deep soils and with regular inputs of rainfall or irrigation, *Eucalyptus* plantations can use water at a rate that approaches the energy limit. Similar high rates of water use have also been observed in *P. radiata* plantations in southern Australia (Benyon et al., 2006) and in Chile (Huber and Iroumé, 2001). Studies in China have found that the annual rate of water use by *Eucalyptus* can be substantially less than both rainfall and available energy (Lane et al., 2004; Ren et al., 2019). This occurs during the dry season and has also been observed in *Pinus* species (Myers et al., 1998). Notwithstanding these similarities, it has been observed that the water use of *Pinus* species decreases more rapidly with the onset of water stress than is the case with commercial *Eucalyptus* alternatives for the same site (Teskey and Sheriff, 1996).

Reviews of the water use potential of *Eucalyptus* have highlighted the variability in reported rates of both transpiration and evapotranspiration (Albaugh et al., 2013; Shi et al., 2012), yet there has been no systematic attempt to determine if the high rate of water use observed in some studies is a characteristic of *Eucalyptus* in plantations or if it has more to do with the conditions that prevailed in those studies. Most of the published studies of water balance, with a couple of exceptions (Mendham et al., 2011; Scott and Lesch, 1997), have reported water balance measurements made within a single rotation, and most studies cover only a small proportion of that rotation. It is likely that plantations must eventually reach a long-term equilibrium with the local climate and that, except in circumstances where trees have access to off-site water such as a regional aquifer (see O'Grady et al., 2011, for a meta-analysis), these high rates of water use, often observed early in the first rotation, will not be sustained. What is needed is to determine if the longer-term equilibrium water balance of catchments planted with *Eucalyptus* will be associated with different levels of water storage, and therefore streamflow, from those under alternative species options for wood production plantations (McDonnell, 2017).

While there are only four direct comparisons of the annual water balance of *Pinus* and *Eucalyptus*, there are many studies that quantify annual water use by either a *Eucalyptus* or a *Pinus* species. These studies, and their estimates of water use, are very situation specific. Comparison of alternative land uses is complicated by the dominant role that climate and hydrogeology play in determining the local water balance. While vegetation cover has a smaller effect on the catchment water balance than either climate or hydrogeology, it is the part of the system that can be actively managed. If studies are available for the two genera from a comparable range of annual rainfall and evaporative environments, then comparison might be possible through normalising water use (evapotranspiration) with respect to potential or energy-limited evaporation and plotting this as a function of the climate wetness index (ratio of rainfall to potential evaporation). While this approach has previously been used to compare the water use of forests with dryland agriculture (Zhang et al., 2004), the normalisation of both axes with respect to potential evaporation may mask the effect of vegetation on evapotranspiration.

In this study, we collated published annual water balance estimates for plantations with either *Eucalyptus* and/or a *Pinus* species. To test the null hypothesis that the evaporation of commercial plantations of *Pinus* and *Eucalyptus* was the same, we fitted three models to the data, including the model described by Zhang et al. (2004), an exponential relationship between evapotranspiration and rainfall, and a linear relationship between the vegetation evaporation efficiency and rainfall. We also test the hypothesis that variation from the first model is determined by variation in soil depth.

2 Methods

This paper presents a meta-analysis of published measurements of the water balance of *Eucalyptus* and *Pinus* plantations in tropical and temperate regions. The focus of this analysis is on post-canopy closure plantations in a notional equilibrium with the site. The behaviour of the plantations is quantified by comparing an index of the function of the crop (the vegetation evaporation efficiency, VEE) with an index of climate wetness, in the manner proposed by Budyko (1974) and applied by Zhang et al. (2004), to compare forests with dryland agricultural systems.

2.1 Definitions of terms

The terms evapotranspiration, water use, potential evaporation, vegetation evaporation efficiency, and climate wetness index have various meanings in the scientific literature, and to avoid ambiguity, they are defined here as they are used in this paper.

2.1.1 Evapotranspiration and water use

Evapotranspiration (ET) and water use are used in this paper to describe the total evaporation from a vegetated land surface. They are the sum of the transpiration of all plants $(T;$ the evaporation through leaf and other plant surfaces of water drawn from the soil and transported to sites of evaporation through the xylem), water intercepted by plant canopies and evaporated without reaching the ground (interception, I), and evaporation of water directly from soil and litter (often called soil evaporation, E_s). All these processes are affected by the choice of crop and by the management of that crop and should therefore be included as part of the water use of that vegetation.

2.1.2 Potential evaporation (PET)

Evapotranspiration (ET) by any land use is situation specific; it is affected by the climate (energy and rainfall), the structure and function of the vegetation, and by characteristics of the soil and the litter. In this paper, for the purposes of comparison, estimates of water use or evapotranspiration are normalised relative to measures of the local water supply (rainfall) and potential evaporation, which represents the energy-limited maximum rate of evaporation. There are numerous measures of reference or potential evaporation, including the Penman potential evaporation (Penman, 1949), FAO56 reference evaporation (Allen et al., 2005), pan evaporation, and Priestley–Taylor potential evaporation (Priestley and Taylor, 1972). They are all intended to represent the maximum possible rate of evaporation by a land surface covered with vegetation. In this paper, potential evaporation (PET) always refers to Priestley–Taylor potential evaporation (see the notes under data analysis below to see how the Priestley– Taylor PET was calculated for each site). We have used the coefficient 1.26 in the Priestley–Taylor equation; this coefficient accounts for the extra roughness of forests when compared with short crops and pastures (Eichinger et al., 1996). The evapotranspiration of plantations may still, of course, exceed this measure of PET. This may be the case if there is an additional source of energy such as the advection or movement of hot air into the forest. This might occur at the edge of a plantation, especially of it is adjacent to an area of land from which there is a large sensible heat flux. The choice of method for calculating PET is less important than applying the same method for all calculations in this analysis.

2.1.3 Water and energy limit, vegetation evaporation efficiency (k) , and climate wetness index (CWI)

The climate imposes limits on evapotranspiration. Evapotranspiration cannot exceed the amount of water available, which is usually limited to rainfall but may include irrigation and soil stored water and groundwater (O'Grady et al., 2011). Similarly, although evapotranspiration may exceed the calculated PET under some circumstances, it is ultimately limited by available energy.

The relationship between the ratio of actual evapotranspiration to potential evaporation and the climate wetness index (CWI, which is the ratio of rainfall to potential evaporation; Budyko, 1974) provides a simple way of partitioning rainfall between evaporation and runoff. Within this framework, evapotranspiration is water limited when it is less than rainfall and energy limited when it exceeds rainfall. The ratio of evapotranspiration to potential evaporation is termed the evaporation efficiency of a surface (Komatsu, 2003). For example, the ratio of evaporation from a soil to the potential soil evaporation is referred to as the soil evaporation efficiency (Merlin et al., 2016). In this paper, the ratio of evapotranspiration to reference evaporation for commercial plantations of *Eucalyptus* and *Pinus* species is referred to as their vegetation evaporation efficiency (VEE). A more evaporation-efficient plantation converts a relatively greater proportion of available energy to latent rather than sensible heat.

Zhang et al. (2004) developed a simple model that predicted the vegetation evaporation efficiency (VEE) as a function of the climate wetness index (CWI). This model is given by Eq. (1) (Eq. A22 in Zhang et al., 2004) below and includes the parameter c (an empirical catchment characteristic), which captures the effect of hydrogeology and vegetation cover on the vegetation evaporation efficiency.

$$
VEE = 1 + CWI - (1 + CWIc)^{\frac{1}{c}}.
$$
 (1)

2.2 Meta-analysis of published studies

While direct comparisons of the water balance of *Eucalyptus* and *Pinus* plantations are few, there are a reasonable number of previously published estimates of either streamflow or evapotranspiration. These data were collated and used in the meta-analysis described below. The studies included are described in some detail in the Supplement, and the main features are summarised in Tables 1 and 2. A list of potentially suitable references was first found by conducting a series of searches of the Web of Science and Google Scholar. The following searches were conducted:

- 1. title contains (evapotranspiration or water use) and (eucalypt or eucalyptus);
- 2. title contains (evapotranspiration or water use) and (pine or pinus);
- 3. paper contains (evapotranspiration or water use) and (eucalypt or eucalyptus); and
- 4. paper contains (evapotranspiration or water use) and (pine or pinus).

The first two searches yielded fewer than 100 papers in total. The latter two found many thousands of articles. The 200 most relevant in each search were checked to decide their suitability. For inclusion, the paper must measure or estimate evapotranspiration by a *Eucalyptus* or *Pinus* species for at least 1 year. Only planted forests managed primarily for wood production were included. Agroforestry systems were excluded, as were measurements made prior to canopy closure. Native forests and burned forests and plantations with access to the water table were also excluded. Several of the studies covered multiple years. A single value of rainfall and evaporation was calculated as the average of all the years in each study. Sometimes a paper reported multiple estimates of evapotranspiration for forests in the same location and growing under the same conditions. In these cases, average values were calculated for the multiple sites.

After applying these criteria to the articles found in the above searches, a total of 30 *Pinus* and 27 *Eucalyptus* stands were included in the meta-analysis. The location, rainfall data, and evapotranspiration data are provided in the Supplement. The estimates of evapotranspiration were made using one of four methods. The method applied in each study is indicated in Table 1.

2.2.1 Method 1 – measurement and addition of component fluxes

At the stand or plot scale, evapotranspiration (water use) is the sum of evaporation from the soil and leaf litter (E_s) , evaporation of rainfall intercepted by the vegetation canopy (I) , and transpiration or the direct uptake of water by the trees and the evaporation of this water through the leaf surface (T) . Evapotranspiration can therefore be calculated as the sum of the component processes.

2.2.2 Method 2 – one-dimensional water balance

Provided there is no leakage or runoff, evapotranspiration (ET) can be calculated in stand-scale studies as the sum of rainfall (P) and the change in the soil water content (ΔS) between two measurements.

$$
ET = P + \Delta S. \tag{2}
$$

2.2.3 Method 3 – catchment water balance

For a catchment, if there is no change in the amount of water stored in the soil or the groundwater (ΔS) , then evapotranspiration (ET) is simply the difference between rainfall and streamflow (Q) . Over long time periods, it is often assumed that the change in storage is negligible; this is less valid when the period of the estimate is reduced or if the annual total rainfall has a clear temporal trend.

$$
ET = Q - P + \Delta S. \tag{3}
$$

aFull rotation.

Post-canopy closure only.

c

Includes pre- and post-canopy closure.

Table 2. Summary of the studies included in the meta-analysis (see the Supplement for more details and the references for each study). This table indicates the number of studies included by country or continent, species, and climate zone.

2.2.4 Method 4 – eddy covariance (flux towers)

Properly located flux towers can be used to estimate the net carbon and water flux (evapotranspiration) above an ecosystem. The instruments on these towers measure the total solar and net radiation and partition this to latent (evapotranspiration) and sensible heat flux (air temperature change) and heat storage changes in soil and biomass. The covariances of high-frequency measurements of air temperature, humidity, and $CO₂$ are used to calculate the total evaporation and carbon exchange between the atmosphere and the underlying vegetation (Aubinet et al., 2012). Measurements are typically made on a 30 min time interval to represent fluxes from an upwind surface area or footprint. The area of the footprint is dependent on the strength of the turbulence in the air, a function of wind speed and surface roughness elements, and the height of the instruments; thus, the location of the land surface influencing the measurements changes through time. Eddy covariance measurements give total fluxes from the contributing footprint and are thus useful for total ecosystem energy, water, and carbon balances. However, partitioning the fluxes between different contributing vegetation and soil components requires additional measurements, such as sap flow, rain throughfall, and soil evaporation. Also, the measurements are unreliable during periods of stable air and low turbulence, such as still cold nights, but, for the purposes of the analyses in this paper, these are periods typically with very low water fluxes and have only minor influence on the total system water balance. There is substantial literature describing these methods and complementary measurements. A detailed description is beyond the scope of this paper but can be found in Wilson et al. (2001), where the method is compared with alternatives.

2.3 Variations at two sites

A study by Scott and Lesch (1997) in three catchments on the Mpumalanga escarpment in South Africa reported more rapid changes in streamflow after planting of *E. grandis* than after planting of *P. patula*. The soil was very deep, and it is probable, though this was not measured, that evapotranspiration exceeded rainfall and that this was more pronounced in the *E. grandis* than the *P. patula*. To allow for this effect, we assumed a storage of 100 mm m^{-1} of soil and a rate of root extension of 2 m yr−¹ for *E. grandis*, after Dye (1996), and 1 m yr−¹ in *P. patula*. This relative rate is consistent with the observation that streamflow ceased 5 and 10 years, respectively, after the planting of *E. grandis* and *P. patula* (Scott and Lesch, 1997).

Another study included here was made at Lewisham in Tasmania, Australia, by Honeysett et al. (1996). In this study, the effect of drought on the water relations and water balance of *E. globulus* and *E. nitens* was investigated using irrigated controls and rainfed plots. The irrigated treatments were excluded from this meta-analysis. However, to avoid mortality, the rainfed treatments received some supplementary irrigation. This irrigation is included in the rainfall figure in Table 1 and in the Supplement.

2.4 Derived climate and vegetation indices

In each of the papers included in this analysis, evapotranspiration (ET) was estimated from the measurement of other variables by one of the four methods described above. Rainfall data were available for all the studies included in this review. Time series climate data from the 0.5◦ grid point closest to each site were also downloaded for the duration of each experiment (Climate Research Unit Time Series v4.03; Harris et al., 2020). Net radiation was calculated for the location, after Hargreaves and Samani (1985), and then the Priestley– Taylor PET was calculated as follows:

$$
\lambda \text{PET} = 1.26 \left[\frac{s}{s + \gamma} \right] R_{n},\tag{4}
$$

where R_n is net radiation in watts per square metre (W m⁻²), λ is the latent heat of vaporisation of water (2245 kJ kg⁻¹), s is the slope of the relationship between saturated vapour pressure and temperature (kPa $\mathrm{e}^{\circ}C^{-1}$), and γ is the psychrometric constant ($kPa \, ^{\circ}C^{-1}$). These constants are temperature dependent; s was calculated using the empirical model in Eq. (5) (Monteith and Unsworth, 2013), and γ was calculated using Eq. (6) in which T_a and P_a are the average daily air temperature (calculated as the average of T_{max} and T_{min}) and atmospheric pressure (assumed to be 101.3 kPa). c_p is the specific heat of dry air (1.013 kJ kg $^{\circ}C^{-1}$), and ε is the ratio of the molecular weight of water to dry air (0.622).

$$
s = 0.04145e^{0.06088T_a}
$$
 (5)

$$
\gamma = \frac{c_{\rm p} P_{\rm a}}{\lambda \varepsilon}.\tag{6}
$$

For each measurement year at each study location, the vegetation evaporation efficiency (VEE) and the climate wetness index were also calculated using Eqs. (7) and (8), respectively.

$$
VEE = \frac{ET}{PET}
$$
 (7)

$$
CWI = \frac{P}{PET}.
$$
\n(8)

2.5 Meta-analysis

In total, three models were fitted to the data using the nonlinear regression function in R (R-Core-Team, 2013), Eq. (1), an exponential relationship between ET and P (Eq. 9), and a linear relationship between VEE and P (Eq. 9).

$$
ET = ETmax + bekP VEE = VEEmin + dP.
$$
 (9)

In each case, the parameters and the coefficients of determination, R^2 , values were calculated for each genus separately and for the pooled data (R-Core-Team, 2013). Analysis of variance was also completed to test for a significant difference between *Pinus* and *Eucalyptus* in the parameters of each model (R-Core-Team, 2013). The residuals (predicted minus observed) from the first model (Eq. 1) were plotted against soil depth for the sites where these data were available. Linear regression was used to explore the relationship between annual transpiration and annual evapotranspiration. Simple t tests for non-paired observations were used to test for differences between the genera in annual evapotranspiration and the ratio of evapotranspiration to rainfall.

3 Results

3.1 The plantations used in the meta-analysis

In total, 27 *Eucalyptus* and 30 *Pinus* sites were included in the meta-analysis. The details of these sites are summarised in three tables. The most detailed information is in the Supplement, together with the measured and calculated climatic data, estimated evapotranspiration, and the detailed results of the data analysis. The papers from which the data were taken are listed in Table 1, with the rainfall data, species studied, and the method used to estimate evapotranspiration. Table 2 summarises the range of climatic conditions and evaporation rates by species and indicates the number of studies for each species by country or continent.

The analysis included sites from tropical, dry tropical, subtropical, warm temperate, cool temperate, Mediterranean, and montane climates, with both genera represented in all but one climate type and in most locations. There is a bias of *Pinus* studies to the United States and of *Eucalyptus* to Australia (Table 2). Species of *Eucalyptus* represented in order of decreasing number of estimates were *E. globulus* (10), *E. nitens* (H. Deane & Maiden) *Maiden* (7), *E. urophylla S. T. Blake* (3), *E. grandis* (2), *E. urophylla* x *grandis* (2), *E. urophylla* x *globulus* (2), and *E. saligna Sm.* (1) (Table 1). Estimates for species of *Pinus* were made for *P. radiata* (18), *P. taeda L.* (5), *P. patula* (2), a mixed stand of *P. taeda* and *P. palustris Miller* (1), a mixed stand of *P. elliottii Engel.* and *P. palustris* (1), *P. elliottii* (1), *P. caribaea* var. *hondurensis W. H. Barrett and Golfari* (1), and *P. strobus* L. (1) (Table 1). Thus, each genus is represented by species from tropical, sub-tropical, and temperate environments.

3.2 Annual rainfall and evapotranspiration

The annual rainfall at the 24 *Eucalyptus* sites ranged from 489 mm at one of the South Australian sites to 2088 mm at a site in the Rio Grande do Sul in southern Brazil. The range of rainfall was similar for the 27 *Pinus* sites and varied from 600 mm at a South Australian site to 2081 mm at a site near Valdivia in south–central Chile. Interestingly, both the low-rainfall site in South Australia and the high-rainfall site in Chile were planted with *P. radiata*. The situation was similar for the average annual potential evaporation, which ranged from 1005 to 2008 mm at the *Eucalyptus* sites and from 1021 to 2004 mm at the *Pinus* sites (see the Supplement). The median annual rainfall for the *Eucalyptus* and *Pinus* sites, respectively, was 940 and 927 mm, while average potential evaporation was 1480 and 1551 mm (Table 2). Thus, the range and median conditions covered by the sites included in this meta-analysis were very similar for both genera.

Annual rates of evapotranspiration reported for *Eucalyptus* species were between 488 mm at a low-rainfall site in South Australia planted with *E. globulus* (Benyon et al.,

Figure 1. Box-and-whisker plots of annual evapotranspiration for the *Eucalyptus* and the *Pinus* sites. The three horizontal lines in the box show the median and 25th and 75th percentile values. The whiskers show the minimum and maximum values, and x indicates the mean values. The associated labels indicate the actual values.

2006) and 1345 mm at a site in Brazil planted with *E urophylla x E. grandis* (Soares and Almeida, 2001). The lowest and highest annual evapotranspiration for *Pinus* species were 355 mm for *P. radiata* at Jonkershoek, Western Cape, South Africa (Lesch and Scott, 1997), and 1291 mm for *P. strobus* in North Carolina (Ford et al., 2007).

The minimum, mean, median, and maximum rates of evapotranspiration were all slightly greater for the *Eucalyptus* sites than for the *Pinus* sites (Fig. 1), but this difference was not significant ($p = 0.24$). When evapotranspiration was divided by rainfall, the mean values of the ratio for the two genera were nearly identical at 0.77 and 0.76 (Fig. 2). The ratio of evapotranspiration to rainfall varied from 0.45 to 1.31 in *Eucalyptus* and from 0.44 to 1.2 in *Pinus* species. At one site in South Africa (Lesch and Scott, 1997), the rate of evapotranspiration by *E. grandis* exceeded rainfall by 31 % (Fig. 2). At the same site, evapotranspiration by *P. patula* exceeded rainfall by 19 % (Fig. 2).

3.3 The effect of genus (*Eucalyptus* and *Pinus*) on the parameters of three models

Genus (*Eucalyptus* or *Pinus*) did not have a significant effect on any parameter in any of the three models (Table 3). Models 2 and 3 were included to check if normalisation with respect to potential evaporation in Model 1 (Eq. 1, Zhang et al., 2004) was masking the effect of genus on evapotranspiration. The results of the analysis of covariance for Model 2 and Model 3 suggest that this was not the case. However, the lower p value for the effect of genus on the slope parameter of Model 3 is noteworthy and was the result of two to three points of high leverage associated with sites where the soil was very deep $(> 20 \,\text{m})$ or where some irrigation was applied during summer. The full results of the analysis and figures for Models 2 and 3 are included in the Supplement.

Figure 2. Box-and-whisker plots of the ratio of the evapotranspiration to rainfall for the *Eucalyptus* and the *Pinus* sites. The three horizontal lines in the box show the median and 25th and 75th percentile values. The whiskers show the minimum and maximum values, and x indicates the mean values.

3.4 Vegetation evaporation efficiency as a function of the climate wetness index (*Eucalyptus* and *Pinus*)

In Fig. 3, the vegetation evaporation efficiency for each study site is plotted as a function of the climate wetness index. For both the *Eucalyptus* and *Pinus* sites, there is a strong, positive correlation between the vegetation evaporation efficiency and the climate wetness index. For the *Eucalyptus* sites, the model of Zhang et al. (2004; Eq. 1) explained 66 % of the variation in the vegetation evaporation efficiency, while for *Pinus* this decreased to 63% . The parameter c in the model described by Eq. (1) fitted to the data was 2.84 for *Eucalyptus* and 2.64 for *Pinus*. While this may be an important difference, it was not statistically significant ($p = 0.50$), and the value for c when the relationship was fitted to the pooled data was 2.74, and the R^2 was 0.69. Figure 4 shows the ratio of the predicted vegetation evaporation efficiency for *Eucalyptus* to the predicted vegetation evaporation efficiency for *Pinus* as a function of climate wetness index. The maximum proportional effect of genus on the vegetation evaporation efficiency of 3.5 % is predicted to occur where the climate wetness index is 1.

3.5 The effect of soil depth

While the relationships in Fig. 3 are significant for both genera, there is nonetheless substantial scatter. The soil depth was not provided in all the papers included in this analysis. When the residuals (observed minus predicted) were plotted as a function of the soil depth, the relationship was significant for the *Eucalyptus* sites (Fig. 5) but not for the *Pinus* sites (data not shown). A linear relationship with soil depth explained 57 % of the error for *Eucalyptus* and indicated that the model shown in Fig. 3, for c of 3.1, overestimated the vegetation evaporation efficiency in shallow soils and under-

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Table 3. The effect of genus on the parameters of three models for estimating the evapotranspiration of *Eucalyptus* and *Pinus* in plantations.

	p value (species)			Parameter value (pooled data)		
Model $1, Eq. (1)$	c			c		
$VEE = 1 + CWI - (1 + CWIc)^{\frac{1}{c}}$	0.5			2.74		
Model 2 , Eq. (9) $ET = ET_{max} + be^{kP}$	ET_{max} 0.38	$b \qquad k$ 0.62	0.74	ET_{max} 978.5	h -11060	ĸ -0.00804
Model 3, Eq. (10) $VEE = VEE_{min} + dP$	VEE_{min} 0.55	d 0.16		VEE_{min} 0.264	d 0.00029	

Figure 3. The vegetation evaporation efficiency as a function of the climate wetness index (a Budyko plot) for 57 (27 *Eucalyptus* and 30 *Pinus*) published studies. The solid grey lines are the water limit (evapotranspiration is equal to rainfall) and the energy limit (evapotranspiration is equal to potential evaporation). The dotted and dashed lines are for Eq. (1) fitted separately to the data for *Eucalyptus* and *Pinus*.

estimated it in deep soils (Fig. 5), with the model having zero residual with a soil depth around 10 m.

3.6 Transpiration as a proportion of evapotranspiration

A subset of the studies, again indicated in the Supplement, also provided estimates of transpiration made using sap flow sensors. For both *Eucalyptus* and *Pinus*, there was a strong linear relationship between transpiration and evapotranspiration, with an approximate slope of 0.5 (Fig. 6).

4 Discussion

The results of the meta-analysis of the published records of evapotranspiration for *Eucalyptus* and *Pinus* species in this paper suggest that, for a given climate wetness index, the

Figure 4. The ratio of the vegetation evaporation efficiency (VEE) for *Eucalyptus* to the vegetation evaporation efficiency for *Pinus* plotted as a function of the climate wetness index. The vegetation evaporation efficiency was predicted using the separate relationships for the two genera in Fig. 3.

Figure 5. The residuals from Fig. 3 for the *Eucalyptus* sites as a function of soil depth. The model in Fig. 3, with a value for c of 2.84, overestimates the observed value of VEE in shallow soils and underestimates VEE in deep soils.

Figure 6. The relationship between annual transpiration and annual evapotranspiration for the subset of sites where transpiration was measured using sap flow sensors.

water use of *Eucalyptus* and *Pinus* plantations is not significantly different ($p = 0.50$). This does not mean that there are not circumstances, or periods within a rotation, when *Eucalyptus* will use more water than the alternatives. The water balance of plantations and alternatives is very situation specific, and our focus should be on understanding the sources of variation rather than generalising about one land use option. The work of Scott and Lesch (1997) and the results of White et al. (2009) from three *E. globulus* plantations established in southwestern Australia highlight the potential of *Eucalyptus* plantations to exceed the water limit early in the rotation on deep soils. This is an issue that warrants deeper understanding and the development of management strategies. The results of the meta-analysis suggest that the average annual water use by the two genera will be similar over large areas and long time periods (decades). They do not, however, preclude periods of high water use by *Eucalyptus*.

The range of annual rainfall, climate wetness indices, and annual evapotranspiration in the published studies was similar for the 27 *Eucalyptus* and 30 *Pinus* sites included in meta-analysis (Tables 1 and 2 and the Supplement). Only a few sites had climate wetness indices more than 1.5. These were Jijia and Hetou in China, Huape and Valdivia in central Chile, and Coweeta in North Carolina. In the case of the Chinese sites, Lane et al. (2004) and Ren et al. (2019) concluded that plantations of *Eucalyptus* would neither have an important effect on water resources nor on water security in this part of China. Notwithstanding this conclusion, there is still a lot of investment made to quantify the water use of *Eucalyptus* in these regions. Wherever the climate wetness index exceeds 1.5, the amount of streamflow will always be substantial, even in lower-rainfall years (White et al., 2016). Thus, rather than annual water balance, the focus should be on water quality and dry season flow to better understand the effect of land use change, including the planting of *Eucalyptus*, on water security.

For the published *Eucalyptus* and *Pinus* studies analysed here, there was a strong positive correlation between evapotranspiration and rainfall and therefore between the vegetation evaporation efficiency and the climate wetness index (Fig. 3). The coefficient, or catchment characteristic, c was greater in *Eucalyptus* (2.84) than in *Pinus* (2.64), but the difference between the two genera was not statistically significant ($p = 0.50$). When this result was discussed with colleagues in the forestry sector or with people in the forest research community, it was met with responses ranging from mild surprise to disbelief. The belief that *Eucalyptus* uses more water than any of the alternative crops is very firmly entrenched, even though it does not seem to have a firm scientific foundation. Given that the meta-analysis presented in this paper produced a result that was counter to the prevailing view, it is very important to consider the direct and corroborative evidence that either support or oppose this observation. The following paragraphs attempt to provide a mechanistic basis for the observation that, while under some circumstances *Eucalyptus* can use water much more rapidly than *Pinus* (Scott and Lesch, 1997), the average behaviour of the two genera appears similar (Benyon and Doody, 2015; Fig. 3). This mechanistic basis is then used to indicate under which circumstances the effects of plantations of *Pinus* or *Eucalyptus* species on water resources should be evaluated and actively managed.

The key to understanding the patterns of water use in *Eucalyptus* and *Pinus* plantations lies in the hydraulic architecture of the two genera and in the way that this affects the relationship between water use and carbon gain. There are some consistent differences between the group of *Eucalyptus* and *Pinus* species that are grown in commercial plantations. First, and very importantly, *Pinus* species are gymnosperms, and their water conducting elements are tracheids, while in *Eucalyptus* water is transported in vessels. The maximum hydraulic conductivity of angiosperms exceeds that of conifers with almost no overlap in the ranges (Sperry et al., 2006). It is the diameter of the vessels that affords angiosperms greater maximum hydraulic conductance (Sperry et al., 2006). It is also known that, in the *Eucalyptus* genus vessel size, the maximum hydraulic conductivity of the xylem is correlated with climate wetness (Pfautsch et al., 2016) so that the major plantation species can have hydraulic conductivities among the highest in the plant kingdom. Leaf conductance and maximum photosynthetic capacity scale directly with the hydraulic conductivity of the xylem (Hubbard et al., 2001; Tyree, 2003).

Thus, plantation *Eucalyptus* species, the most important of which are from the *Symphyomyrtus* subgenus and grow naturally in the wetter fringes of the Australian continent, have a higher maximum hydraulic conductivity, water use, and photosynthetic capacity than commercially grown *Pinus* species (Whitehead and Beadle, 2004). In the early growth phase, *Symphyomyrtus Eucalyptus* species also have a much higher specific leaf area (ratio of leaf area to mass) than

Pinus, and this results in more rapid canopy development and the potential for faster early growth and water use, such as that observed by Scott and Lesch (1997). This can, of course, only happen if there is water available to support this growth and canopy development, and this can be supplied by rainfall throughout the year or by additional sources of water stored in deep soil (Dye and Olbrich, 1992; Scott and Lesch, 1997; White et al., 2014), shallow groundwater (Benyon et al., 2006; Brooksbank et al., 2011; Eamus et al., 2000; O'Grady et al., 2011), or applied as irrigation (Honeysett et al., 1996). If *Eucalyptus* plantations are grown on deep soils and in regions where the climate wetness index is much less than one (potential evaporation exceeds rainfall), then, by virtue of their hydraulic architecture, they have the potential to affect the water balance more than alternatives.

The capacity of *Eucalyptus* for high instantaneous sap velocities that are associated with elevated photosynthetic capacity also affects the seasonal patterns of water use in *Eucalyptus* compared to *Pinus*. Transpiration of *Eucalyptus* species increases rapidly in spring and is associated with high maximum stem and leaf conductivity (White et al., 1999). The relative behaviour of *E. globulus* and *P. radiata* is well understood, making them good exemplars. They are also two plantation species of great global importance that are grown in similar areas, including in central Chile. In Chile and Australia, *P. radiata* is known to be capable of surviving more severe droughts than *E. globulus*, and plantations of the species therefore extend into drier areas than *E. globulus* both in Chile and in Australia. The greater drought tolerance of *P. radiata* than *E. globulus* is mediated by a much stronger stomatal response to soil drying (Mitchell et al., 2014). In situations where the amount of soil water storage imposes an upper limit on annual use, although this store of water will be completely depleted by both species, it will then be used earlier in the growing season by *Eucalyptus*. Thus, the period of peak physiological activity and growth in *Eucalyptus* is associated with lower average temperatures and more moderate air saturation deficits. This pattern of water use, biased towards spring and early summer, can result in very efficient water use growth and wood production (White et al., 2015). This behaviour of the *Eucalyptus* is closer to a mimic of the seasonal water use pattern of an annual species. This mechanism underlies the greater water use efficiency of *Eucalyptus* species than of the *Pinus* but is also associated with an increased risk of mortality (White et al., 2003, 2009) if the soil water runs out. It also underlies the high rates of water use sometimes observed in deep soils (Scott and Lesch, 1997) and the high leverage in Model 3 of *Eucalyptus* grown on deep soil (see the Supplement).

At equilibrium, *Eucalyptus* and *Pinus* species generally have different seasonal patterns of water use. Nonetheless, the average annual water use does not differ significantly between the two genera amongst the published studies presented in Fig. 3. This observation is entirely consistent with the observed hydraulic architecture of these two genera in the field. Radiation interception and absorption, and therefore productivity and evapotranspiration in forests, including plantations, are strongly correlated with leaf area index. Battaglia et al. (1998) proposed that, after the canopy closes, plantations will arrive at an equilibrium leaf area index that maximises the net primary productivity. They further demonstrated that the value of this optimum leaf area index is strongly influenced by the climate wetness; higher optimum values of leaf area index were observed in wetter situations. The value of this optimum leaf area index tends to be higher for a given climate wetness in *Pinus* species than in *Eucalyptus* species. For those experiments included in this analysis that reported leaf area index, the average value for *Pinus* was approximately 4, which is nearly a full unit greater than the average value for the *Eucalyptus* plantations.

In comparing *Eucalyptus* and *Pinus* in commercial plantations, it is important to compare at least one, and possibly more, full crop rotation. *Pinus* is generally managed for solid wood production and therefore on a longer rotation than *Eucalyptus*, which is usually, but not exclusively, grown for pulpwood production. Around the world, the time from planting to harvest of *Pinus* species is between 2 and 3 times that of the *Eucalyptus* in the same location. In Chile, for example, *Eucalyptus* is harvested after about 12 years, while *Pinus* is grown for about 25 years. *Pinus* is usually grown for solid wood or veneer production and is therefore thinned at least once and is often pruned to produce clear wood. After the harvesting of the first *Eucalyptus* crop, a *Pinus* plantation on the same location would remain standing and operating at, or near, the water limit. For a period of between 2 and 3 years after the *Eucalyptus* harvest, the evapotranspiration of the *Pinus* will therefore exceed that of the *Eucalyptus*. This is evident in the results of Scott and Lesch (1997), who compared *E. grandis* with *P. patula*. The frequency of the harvest of *Eucalyptus* will be a key factor affecting the comparative water balance of *Pinus* and *Eucalyptus* plantations. Paradoxically, more frequent harvests will increase the average streamflow from *Eucalyptus* plantations relative to *Pinus*. It has been demonstrated that the effects of thinning on the water balance are transient, lasting for a maximum of 1 year in both *Pinus* and *Eucalyptus* (Scott and Lesch, 1997; White et al., 2014).

The proportion of evapotranspiration that occurs as transpiration was approximately 0.5 for both *Pinus* and *Eucalyptus* across a wide range of climate wetness indices (Fig. 6). This means that the annual partitioning of evapotranspiration to fluxes other than transpiration is similar for these two genera. The partitioning of these other fluxes to understorey transpiration, soil evaporation, and interception may have important implications for ecosystem productivity and efficiency. The water use efficiency of wood production is directly correlated with the ratio of transpiration to other fluxes (White et al., 2015). In a study that compared *E. globulus* and *P. radiata*, Benyon and Doody (2015) observed that interception was more than half the non-transpirational fluxes in *P. radi-* *ata* and less than half in *E. globulus*. This variation in partitioning is a direct consequence of the previously noted tendency for *Pinus* to have a higher leaf area index than *Eucalyptus* and the greater canopy storage per unit leaf area in *Pinus* than in broadleaved species (Iida et al., 2005). A weakness of this analysis and of the literature on water balance is the exclusion of stemflow from most water balance studies. It is likely that stemflow will contribute more to throughfall in *Eucalyptus* (7 % of rainfall) than in *Pinus* (2 % to 5 %; Crockford and Richardson, 1990). This difference is approximately equivalent in magnitude to the observed, albeit nonsignificant, difference between the genera in this analysis.

5 Conclusion

Water use by vegetation is very situation specific. The comparison between *Eucalyptus* and *Pinus* depends on the age of the plantation, the length of the rotation, the seasonality of rainfall, and the depth of the soil. In this paper, a metaanalysis of published estimates of evapotranspiration by *Pinus* and *Eucalyptus* species in commercial plantations did not find a significant difference between the genera. Specifically, while there was a small, but systematic, difference of about 3 % in water use between the genera (see Figs. 5 and 6), this analysis finds that, for a given climate wetness index, the evapotranspiration by *Pinus* and *Eucalyptus* was statistically the same. Moreover, our understanding of the hydraulic architecture and stomatal physiology of pines and eucalypts suggests that, although the long-term average behaviour may be similar, there will be differences in their temporal pattern of water use both within and between years. *Eucalyptus* will use more water than *Pinus* early in the growing season and in the early years of the rotation. On deep soils, this may result in lasting differences but under most circumstances the total effect on the water balance will be similar. The reputation of much higher water use by *Eucalyptus* may stem partly from the observation of vigorous early growth of *Eucalyptus* and the many studies on young plantation stands.

Code and data availability. Codes and data sets are provided in the Supplement.

Supplement. The supplement related to this article is available online at: [https://doi.org/10.5194/hess-26-5357-2022-supplement.](https://doi.org/10.5194/hess-26-5357-2022-supplement)

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