# Viewpoint

## Plant water use theory should incorporate hypotheses about extreme environments, population ecology, and community ecology

## **Summary**

Plant water use theory has largely been developed within a plantperformance paradigm that conceptualizes water use in terms of value for carbon gain and that sits within a neoclassical economic framework. This theory works very well in many contexts but does not consider other values of water to plants that could impact their fitness. Here, we survey a range of alternative hypotheses for drivers of water use and stomatal regulation. These hypotheses are organized around relevance to extreme environments, population ecology, and community ecology. Most of these hypotheses are not yet empirically tested and some are controversial (e.g. requiring more agency and behavior than is commonly believed possible for plants). Some hypotheses, especially those focused around using water to avoid thermal stress, using water to promote reproduction instead of growth, and using water to hoard it, may be useful to incorporate into theory or to implement in Earth System Models.

#### Introduction

Predicting plant water use for species with differing traits and across environmental conditions is a central challenge for ecophysiology (Venturas et al., 2017; Kannenberg et al., 2022). Accurate predictions are critical for fundamental knowledge of plant biology, as well as for upscaling to ecosystem fluxes, for example, via Earth System Models (Christoffersen et al., 2016; Fisher et al., 2018). Current marginal gain theory for water use (Wang et al., 2020) has been developed from a foundation of stomatal optimization framed in terms of maximizing carbon gain (Cowan & Farquhar, 1977). This theory proposes that plants maximize  $A-\Theta$ , where A represents carbon gain and  $\Theta$  represents some carbon penalties or costs, for example, relating to soil drying, or hydraulic failure. If E is the amount of water plants use, then mathematically, maximization occurs when  $\frac{\partial A}{\partial E} = \frac{\partial \Theta}{\partial E}$ , that is, when the marginal gain equals the marginal penalty (Wang et al., 2020). Implementations of theory make different assumptions about A

and  $\Theta$ , for example, whether the timescale of each is instantaneous or temporally extended (i.e. whether A and  $\Theta$  are defined as a rate or a quantity), whether the biological scale is a single leaf or a whole plant, and whether the environment is constant or variable. Often, E is assumed to depend only on stomatal opening, so the term 'stomatal optimization theory' is commonly used interchangeably with 'marginal gain theory'.

Current iterations of marginal gain theory are widely considered reliable and generally suggest that the penalty depends on vapor pressure deficit, or soil water potential, or other similar variables (Wang et al., 2020; Kannenberg et al., 2022), leading to the development of marginal gain models that consider the entire soilplant-atmosphere continuum (Sperry et al., 2016; Wolf et al., 2016). For example, some empirical models based on vapor pressure deficit (Leuning, 1995; Medlyn et al., 2011) are widely used to parameterize Earth System Models, and predictions are often reliable at leaf and ecosystem scale (Franks et al., 2018). Nevertheless, substantial uncertainty remains about the biological and temporal scales over which this optimization is valid (Feng et al., 2022).

Plant water use may be more complex than what most implementations of marginal gain theory indicate. As a first example, theory often fails in hot environments. Previous efforts to model gas exchange at higher thermal stresses (Franks et al., 1997; Eamus et al., 2008) have yielded results that are at odds with stomatal optimization theory thought to apply at short timescales (Medlyn et al., 2011). Heat avoidance can occur, where water use is optimized not to maximize instantaneous carbon gain per unit water loss (i.e. marginal gain theory), but instead to prioritize evaporative cooling that avoids thermal stress/mortality at high temperatures (Chaves et al., 2016; Slot et al., 2016; Urban et al., 2017; Blonder & Michaletz, 2018; Aparecido et al., 2020; Marchin et al., 2022a) and thus enables sustained long-term carbon gain. As a second example, stomatal optimization theory thought to apply to isolated individuals can fail when plants are measured in community contexts. Recent models of stomatal optimization under competition (Wolf et al., 2016; Lu et al., 2020) indicate that water use strategies should change when neighbors compete for water. This prediction of water use shifting when growing with competitors vs alone has been upheld in at least two sets of experiments (Vysotskaya et al., 2011; Zenes et al., 2020).

## Conceptual framing

Marginal gain theory has been limited in four practical ways. First, validations of theory have been carried out primarily on a limited set of species, often crops (von Caemmerer & Evans, 1991; Harley et al., 1992). While these cases are clearly of high interest, they represent a limited and biased subset of plant functional and evolutionary diversity. Second, tests of theory have been limited in 4698137, 0, Downloaded from https://nph.onlinelibrary.wiley.com/doi/10.111/nph.18800, Wiley Online Library on [0603/2023]. See the Terms and Conditions (https://onlinelibrary.wiley.com/terms-and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons Licensean Conditions (https://onlinelibrary.wiley.com/terms-and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons Licensean Conditions (https://onlinelibrary.wiley.com/terms-and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons Licensean Conditions (https://onlinelibrary.wiley.com/terms-and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons Licensean Conditions (https://onlinelibrary.wiley.com/terms-and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons Licensean Conditions (https://onlinelibrary.wiley.com/terms-and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons Licensean Conditions (https://onlinelibrary.wiley.com/terms-and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons (https://onlinelibrary.wiley.com/terms-and-conditions) on the applicable Creative Commons (https://onlinelibrary.wiley.com/terms-

extreme (very hot and concurrently very wet or very dry) environments (Schulze *et al.*, 1973; Aparecido *et al.*, 2020; Grossiord *et al.*, 2020). This limitation is especially critical for accurate predictions in response to novel environmental conditions (Williams & Jackson, 2007) projected under global change. Third, most data come from measurements of leaves, isolated from the whole-plant context, or individual plants, isolated from their population or community contexts. While such data provide snapshots of behavior under controlled conditions, they may not accurately represent behavior in complex natural conditions. Fourth, theory may not yet provide a sufficient description of all the biological processes affecting water use. This reflects either a trade-off between model complexity and usefulness (Harrison *et al.*, 2021), or alternatively, incomplete process knowledge or data for parameterizing models.

Marginal gain theory is also limited by its foundational assumption that plant water use is a problem analogous to the rational actor problem in classical (Smith, 1776) and neoclassical economics (Jevons, 1879). Both propose that the leaf or plant (analogously, the individual) has evolved mechanisms (chooses actions) that maximize performance (maximize utility or diminish marginal utility) due to natural selection (self-interest). Both also assume that the optimization can be conceived in terms of a single measure and store of value, carbon (money). This assumption has been widely critiqued in economics (Hollis & Nell, 1975; Sen, 1977), as it leads to both inaccurate predictions of human behavior (Veblen, 1898; Fullbrook, 2004), and also negates the possibility of considering cooperative behavior and multiple incommensurable types of values relevant to decision-making (e.g. 'human wellbeing' or 'ecosystem health'; Jackson, 2016). Marginal gain theory for plant water use draws from this economic heritage and implicitly requires us to accept the premise that plant behavior can be collapsed to the single currency of carbon, and the single objective of getting as much of it as possible. This logic also produces the concepts of ecosystem service valuation (Reid et al., 2005) and natural capital (Costanza et al., 1997), which propose that moneybased prices or valuations can be given to values that may actually not be commensurable. Returning to plant water use, it therefore may be useful to cast off the shackles of old economic ideas, and embrace a broader perspective on the values of water to plants that are not defined as, or priced in terms of, carbon gain.

Our central hypothesis is that the value of water to a plant extends far beyond its immediate value for instantaneous carbon gain, requiring either extension of marginal gain theory to more biological and temporal contexts, or alternatively development of new theory that asks how water use influences fitness—not ecophysiological performance. We survey hypotheses that operate at different biological and temporal scales (illustrated in Fig. 1) that might cause current theory to fail, which are linked to performance in extreme environments ('E' hypotheses), to life history and population ecology ('P' hypotheses), and to species interactions and community context ('C' hypotheses). We then sketch a range of observed and yet unobserved hypotheses that are still on the frontier of current theory and that may run contrary to current theory's predictions.

Some of the hypotheses we propose are at odds with current understandings of how plants process information and ascribe a

level of agency to plants that is not widely considered reasonable at individual or population scale. There is also little evidence to date that plants have evolved mechanisms to enable such complex behavior. Nevertheless, these strategies remain an underinvestigated conceptual possibility. We used agentive terms within these hypotheses below because they are standard within game theory and computer science. Additionally, the limits of plant agency, behavior, communication/sensing, and social interactions remain poorly studied (Silvertown & Gordon, 1989; Karban, 2015; van Loon, 2016).

## Hypotheses relating to extreme environments

## E1. Avoiding thermal damage or mortality

High transpiration may reflect prioritization of leaf cooling to avoid heat damage or death in thermally extreme environments (Fig. 2a). This appears contrary to the stomatal optimization prediction (over short timescales) that stomata should close to prevent hydraulic failure and also appears maladaptive because of the reduced efficiency of photosynthesis at high temperatures. However, increasing transpiration at high temperatures actually can be optimal over long timescales if water is available and doing so prevents leaf thermal damage or mortality, that is, if it promotes future carbon gain, and offsets the loss of carbon invested in the construction of the leaf (Blonder & Michaletz, 2018). This type of optimization is difficult to capture in analytic models because leaf mortality is not reversible: if a leaf has exceeded a critical temperature even once, its future carbon gain is zero, regardless of the temperature the leaf later obtains.

Low transpiration rates prevent evaporative cooling and lead to high temperatures in leaves (Gates, 1968; Monteith & Unsworth, 2013) and also limit advective heat transfer by xylem sap, which can lead to high temperatures in branch and stem phloem, xylem, and sapwood (Swanson, 1994). High leaf and branch temperatures, if sustained, could cause permanent (or costly-to-repair) biochemical damage, structural changes and reduced hydraulic conductivity of xylem (Michaletz et al., 2012), or cell and tissue necrosis (Teskey et al., 2015). Failure to evaporatively cool can also cause mortality. For example, during a hot drought in Australia, crown dieback in urban trees was negatively correlated with the critical temperatures for photosystem II dysfunction, while leaf water potentials showed trees were not severely water stressed (Marchin et al., 2022b). This suggests that dieback was driven by heat stress (not embolism) resulting from reduced transpirational cooling that enabled leaf temperatures to exceed a critical threshold.

Cooling is likely to be important in plants with relatively carbon-expensive and/or evergreen leaves, in environments that experience transiently high temperatures, especially when combined with high soil water availability (Aparecido *et al.*, 2020). For example, stomatal responses to vapor pressure deficit and temperature can become decoupled in some desert plants (Schulze *et al.*, 1973). There are likely more efficient ways to keep leaves cool via trait evolution over longer timescales, or alternatively higher thermal tolerances to occur via phenotypic plasticity or evolution. These latter approaches are likely taken by many hot-adapted species,

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## Categories of plant water use hypotheses

E: Extreme environments P: Population ecology

C: Community ecology

M: Marginal gain

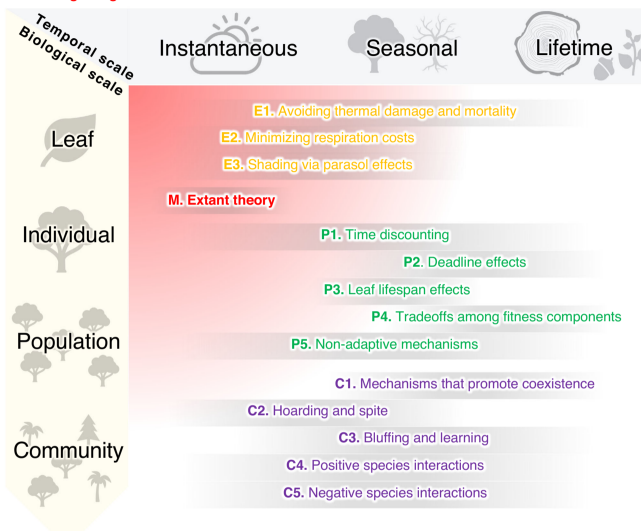


Fig. 1 Overview of hypotheses considered in this manuscript. Hypotheses are arranged by biological scale (vertical) and temporal scale (horizontal) at which they might operate. They are colored by category: E, extreme environments (orange); P, population ecology (green); C, community ecology (purple). They are also contrasted with extant theory: M, marginal gain (red).

which can see leaf temperatures reach 48-49°C associated with midday depressions of photosynthesis and transpiration, while CAM plants can exceed 50°C leaf temperature during midday, and some cacti obtain temperatures > 50°C, with heat tolerances up to 69°C (Smith et al., 1984). For example, the Rubisco activase isoform in Agave remains active at up to 50°C, 10°C higher than Oryza isoforms (Shivhare & Mueller-Cajar, 2017).

Several underlying physiological processes could generate this cooling response. The most straightforward possibility is stomatal opening at high temperatures. Sensory mechanisms remain unknown, but may include the detection of unfolded proteins

and changes in membrane fluidity (Hayes et al., 2021). Abscisic acid (ABA) may play a role, as its accumulation is linked to reduced transpiration and increased hydraulic conductivity during drought stress (Muhammad Aslam et al., 2022). Alternatively, other signal transduction mechanisms could cause high-temperature stomatal opening (Kostaki et al., 2020). Another speculative possibility is stomatal popping, in which pressurized hot air in intercellular spaces forces guard cell opening (Brix et al., 1992; Aparecido et al., 2020). This is unlikely because such popping would occur only once, after which depressurization would occur and prevent further water flux. Alternatively, transpiration fluxes could arise



(a) Avoiding thermal damage and mortality (E1)



(c) Deadline effects (P2)



(d) Tradeoffs among fitness components (P4)

(b) Shading via parasol effects (E3)



(e) Hoarding and spite (C2)



(f) Positive species interactions (C4)



Fig. 2 Speculative examples of the multiple values of water to plants. Photographs are selected to illustrate concepts, not necessarily because they directly confirm a hypothesis. (a) In southeastern Arizona, a cottonwood tree (Populus fremontii) growing in hot conditions with abundant water may use water for evaporative cooling instead of photosynthesis (Hypothesis E1). (b) In a moist tropical forest in Borneo, water used to maintain the top layer of a forest canopy may provide shading to photosynthetically active leaves in lower layers (Hypothesis E3). (c) In the northern Sonoran Desert where winter rain is the only moisture source before spring/foresummer drought, water may be used as quickly as possible to support seed production before the drought deadline occurs (Hypothesis P2). (d) In a tomato plant, fitness may be enhanced by allocating water to higher fruit production even if it reduces water available for photosynthesis and growth (Hypothesis P4). Image in public domain. (e) In the southern Sonoran Desert, succulent or deep-rooted plants may hoard water (via vegetative storage or hydraulic descent) to prevent it from being used by competitors (Hypothesis C2). (f) In the alpine zone of the Rocky Mountains, plants may use water to help neighbors in a facilitative or mutualistic relationship, for example, by leaf transpiration that co-creates a more humid microclimate in dry sites, or via investment in flowers and flower transpiration that attracts shared pollinators (Hypothesis C4). Photo credits: (a) Kevin Hultine; (b, c, e, f) Benjamin Blonder; (d) public domain.

from high cuticular conductance, which is known to increase at higher leaf temperatures, due to changes in the physical properties of cuticular waxes. This last cooling mechanism seems most plausible and has been described in desert (Bueno et al., 2019) and tropical (Slot et al., 2021) species, though it is unclear yet whether it is adaptive.

Cooling can also occur via leaf positioning rather than transpiration. For example, in Piper auritum, a tropical species associated with high light conditions in canopy gaps (Chiariello et al., 1987), high water use results in midday wilting. This reduces the direct sunlight that hits the leaves in their exposed habitat, which in turn decreases leaf temperatures. The reduction in incident irradiance caused photosynthesis to decrease, but less so than transpiration, so water use efficiency is improved by the avoidance of midday sun. The continued photosynthesis, albeit at reduced rate, suggests that this strategy enables optimization of time-integrated carbon capture. Similar behavior is observed in wheat, where leaf erectness and leaf rolling behavior are breeding targets for improving heat tolerance (Hunt et al., 2018).

Cooling requires available water, which can depend both on precipitation but also species properties like maximum rooting depths and capacitance. For example, in hot conditions, deeply rooted evergreen woody plant taxa, Quercus turbinella and Rhus ovata, were shown to achieve high leaf-level transpiration rates that were decoupled from rates of carbon uptake (Aparecido et al., 2020). Similarly, stem gas exchange measurements conducted on mature Carnegiea gigantea (saguaro cactus) in the Sonoran Desert during midsummer yielded daytime transpiration rates that at times equaled nighttime transpiration rates during Phase I of the CAM cycle (Bronson et al., 2011). These patterns indicate that under heat stress, giant saguaros either actively transpire water stored in their succulent stems by opening their stomata during the day, in turn losing carbon fixed at night potentially as a mechanism to reduce stem surface temperatures – or experience daytime water loss through the cuticle as surface temperatures increase. As another example, transient shallow soil moisture conditions enable the desert shrub, creosote (Larrea tridentata) to take a riskier, anisohydric hydraulic strategy following precipitation pulses in both winter and summer; while adopting a more conservative, isohydric strategy in other periods when shallow soil moisture is absent (Guo et al., 2020). Likewise, warm-adapted cottonwood (Populus fremontii) - a desert riparian tree species increases stomatal conductance during the warmest time of the year, unlike cool-adapted genotypes that show no seasonal changes in stomatal conductance (Blasini et al., 2022). Consequently, warm-adapted genotypes maintain cooler leaves than cool-adapted genotypes, which improves carbon balance and protects against leaf thermal damage.

## E2. Minimizing respiration costs

Nighttime transpiration is observed in many species when environments are hot (Dawson et al., 2007; Yu et al., 2019). Reduced nighttime transpiration is recognized as a valuable target trait for breeding crops with high water use efficiency, for example Coupel-Ledru et al. (2016). Nighttime transpiration in C<sub>3</sub> and C<sub>4</sub> plants is challenging to explain from an optimality perspective as it results in high water costs and no instantaneous carbon gain. An alternative explanation for this observation involves optimization of carbon gain over longer timescales. Cooling leaves at night via transpiration may reduce respiration rates, which increase exponentially with temperature. This strategy may complement or trade off with acclimation of respiration to elevated nighttime temperature et al., 2021). While the amount of cooling achieved by low rates of nighttime transpiration is likely small, the effect on carbon flux may nonetheless be significant. Reductions in nighttime carbon losses from respiration may offset low net carbon gains during the daytime when photosynthesis is limited at high temperatures and respiration is high. This might occur in hot environments where nighttime respiration is a major cost, and only for C<sub>3</sub> and C<sub>4</sub> species (Wang et al., 2021). Nighttime transpiration has also been linked to refilling of xylem embolisms (Zeppel et al., 2014), though this may occur only under high water potentials (Klein et al., 2018). Such an effect would in turn

benefit daytime carbon fixation, as it would enable more sustained stomatal opening.

## E3. Shading via parasol effects

In extremely hot or bright environments, upper canopy sun leaves may photosynthesize suboptimally at the leaf level, but still contribute significantly to overall plant carbon assimilation (Ishii *et al.*, 2004; Fig. 2b). While the morphology of sun leaves often helps maximize convective heat losses to the atmosphere (Vogel, 2009), such leaves may become too hot or light-saturated to efficiently carry out photosynthesis for much of the daylight hours and may also require a high water supply in order to maintain open stomata, cool via transpiration, and avoid thermal mortality. However, this upper canopy layer may provide sufficient shading and associated cooling benefits to produce a canopy microclimate that enables lower-canopy shade leaves to photosynthesize at a lower water cost. This hypothesis has been supported in computer models of woody species architecture (de Haldat du Lys *et al.*, 2022).

There is limited empirical support for positive effects on carbon gain of shading in a Puerto Rican tropical wet forest (Miller et al., 2021) and also in a temperate tree experiment (Kothari et al., 2021). While leaves under moderate shade may have lower maximum photosynthesis rates, the lower maximum tissue temperature and lower leaf-to-air vapor pressure deficit may enable them to avoid midday stomatal depression and maintain photosynthesis during most of the day. Thus the standard carbon-for-water optimization could still occur, but at the scale of the whole plant, not the scale of a single leaf. This strategy might occur if the relative benefit of lower-canopy leaves exceeds the relative cost of sacrificial upper canopy leaves that act as a parasol for the overall plant. Alternatively, sun leaves, which are typically much shorter-lived than shade leaves (Reich et al., 2004), might operate at high photosynthetic rate for a while when they are young, and then continue their role as parasols until they are replaced by new leaves. Regardless, measurements of photosynthetic capacity that focus on sun leaves may obscure optimization occurring at whole-plant level.

## Hypotheses relating to population ecology

## P1. Time discounting

In plants with 'fast' life histories or strong competition among species over a common growing season, there may be substantial benefits to carbon gained earlier in ontogeny (e.g. photosynthetic cotyledons and early leaf flushes). Carbon acquired earlier in life can be reinvested in the deployment of additional photosynthetic tissue, which further accelerates growth (Chabot & Hicks, 1982). This is because, from an economic perspective, the time discounting rate for carbon could be large, and costs incurred later may not be as important as benefits obtained earlier (Westoby et al., 2000). This perspective is widely discussed in the leaf lifespan literature (Kikuzawa & Lechowicz, 2006; Falster et al., 2012; Castorena et al., 2022) and can be incorporated into stomatal optimization models, though most models do not include it (Wang

et al., 2020). Thus, high water use early on in a life cycle or in a growing season may be optimal, even if it later leads to drawdowns in soil moisture, stem hydraulic failure, or leaf mortality. For example, many facultative CAM plants begin life using C<sub>3</sub> photosynthesis, because they prioritize rapid early growth over water use efficiency (Winter et al., 2011).

#### P2. Deadline effects

Related to the time discounting mechanisms, species may sometimes maximize fitness by exploiting limited resources as rapidly as possible, for example, desert winter annuals where lifespans are limited by the onset of an arid foresummer, or in a savanna where locust outbreaks may occur driving fatal levels of herbivory (Fig. 2c). As in a workplace analogy, any species that meets its deadline (completes its life cycle) is a winner, while all others that cannot meet the deadline are losers (Aronson *et al.*, 1992; Peñuelas *et al.*, 2004). This is effectively a special case of time discounting where the discount rate becomes infinite after a certain amount of time. In such a case, water use may reach levels high enough to exhaust water resources and risk vegetative tissue damage or death, so long as it leads to successful reproduction within a minimum time interval.

## P3. Leaf lifespan effects

Species differing in their leaf lifespan (deciduous vs evergreen) may have different water use strategies (Schulze, 1982). In tropical species, there is often a trade-off between drought avoidance (deciduousness) and hydraulic safety (Oliveira et al., 2021). However, much of this data comes from seedlings, so empirical knowledge remains limited. Deciduousness may also influence water use in ways unrelated to hydraulic safety. Drought-deciduous species do not have leaves for part of the year and benefit from not needing to pay the water and carbon costs of maintaining them (via transpiration and respiration, respectively; Vico et al., 2017). However, they also lose out on the opportunity to use water at other times of year to support carbon uptake, as well as on the opportunity to vegetatively compete with other species that could otherwise use the same shared pool of soil water (note that this argument is reversed when snowy/wintry conditions rather than drought are considered). Deciduous species also may take more aggressive water use strategies (Zeppel et al., 2014) when in the presence of evergreen species or when they have evolved under competition from evergreen species.

## P4. Trade-offs among fitness components

In plants where fitness is most demographically sensitive to variation in reproduction or survival, rather than to growth, water may be used in ways that deprioritize carbon gain in leaves (Fig. 2d). This could drive apparently nonoptimal stomatal regulation at the leaf level if allocation is not correlated with signals known to drive stomatal regulation, such as variation in stem water potential. For example, allocation of carbon towards

osmotic regulation (e.g. for freezing responses) or storage (at the end of a season), or allocation of water towards fruit production, could lead to low leaf water use or leaf death, even when this strategy is optimal for fitness overall.

Some of these strategies might be expected to be more common in species where selection may be stronger for nongrowth fitness components. During drought, plants might be more limited by water than by carbon, so they may prioritize allocation to reproduction over that to leaves, not only because of fitness but also because water loss may be higher from leaves than from flowers or fruits. In stressful conditions, prioritization of water for fruit growth has been reported in Solanum lycopersicum (tomato; Harrison Day et al., 2022) while prioritization of transpiration in flowers occurs in Glycine max (soybean; Sinha et al., 2022), and prioritization of storage occurs in Picea abies (spruce). There is also increased allocation to reproduction when water is limited during seasonal and El Niño drought in tropical forest (Detto et al., 2018). Similarly, some annual (monocarpic) crop species need to initiate whole-plant senescence to remobilize and transfer assimilates to grains (Yang & Zhang, 2006). The amount of carbohydrates fixed before senescence can be particularly important for grain filling (Asseng & Van Herwaarden, 2003). Therefore, leaves of annual plants may maximize short-term assimilation in the phase just before grain filling and senescence.

Additionally, selection may favor whole-plant survival even when it impacts leaf performance. For example, using advective heat transfer of xylem sap described previously, *Pinus ponderosa* seedlings have been observed to maintain stem temperatures as much as 15°C below ambient air temperature, enabling them to survive in open forests where soil surface temperatures could exceed 75°C (Kolb & Robberecht, 1996). The lethal temperature for stems was 63°C, so the elevated stomatal conductance found in needles of surviving plants appears critical for whole-plant survival. As another example, *Helianthemum squamatum* switches between surface and deep water sources in dry conditions. This reduces water use efficiency, because the switching behavior has high costs that lead to reduced nutrient uptake, which then reduces leaf photosynthetic capacity (Querejeta *et al.*, 2021).

## P5. Nonadaptive mechanisms

It is also possible that very high or low water use may be maladaptive and occur as a necessary cost driven by other processes that influence population fitness. A range of mechanisms are possible, and increasingly being recognized. Limited phloem export capacity or sink limitation may mean that water is often not transpired when it is available and could be used (Fatichi *et al.*, 2014), because any resulting carbon gain would result in the accumulation of reaction products and inhibit further reactions (e.g. driving the commonly observed late afternoon depression in photosynthesis). This also could occur if plants are adapted to lower levels of atmospheric [CO<sub>2</sub>] than they currently experience. The same outcome can result from sink limitation; reduced sink activity results in carbohydrate accumulation in leaves, which triggers the downregulation of photosynthesis and transpiration (Quereix *et al.*, 2001; Li *et al.*, 2007).

Additionally, the temperature dependence of cuticular conductance described previously could lead to higher water use than would be adaptive, either because it is an unavoidable biophysical reality (Slot *et al.*, 2021) or because of trade-offs with photosynthetic capacity (Machado *et al.*, 2021).

Alternatively, correlated selection (e.g. due to genetic linkage) could occur, resulting in nonoptimal water use being a necessary consequence of strong selection for other traits. However, there seems to be limited evidence for this given the omnigenic basis of many complex traits in plants (Boyle *et al.*, 2017). Or, in some abiotically nonstressful environments, neither water use nor carbon gain may be limiting to performance, when for example, competition for pollinators or dispersers may be a priority. In such cases, water use strategies may be variable and inexplicable from an optimality perspective (Gould & Lewontin, 1979). While it seems unlikely that selection does not act strongly on water use, the alternative hypothesis should be rejected based on evidence rather than assumption.

## Hypotheses relating to community ecology

## C1. Mechanisms that promote coexistence

The fitness of a population may be influenced by community context. Mechanisms that promote coexistence effectively increase the long-term fitness of a population. Therefore, mechanisms influencing coexistence may lead to selection on water use behavior for species when they occur with other species, whether the underlying species interactions are positive or negative. Coexistence mechanisms are processes that yield higher intraspecific competition relative to interspecific competition when each species is rare, meaning that species are able to recover from low densities without becoming extinct (Chesson, 2000). These coexistence mechanisms necessarily operate at scales beyond that of the individual or population.

Temporal niche partitioning is one such mechanism. If water is a key resource, then some species may perform better only when water availability is consistent over time and space, while others may perform better when water availability is variable over time and space. Thus, some species may appear to use nonoptimal water use strategies when measured in a constant environment, when in fact their strategy is optimal for population dynamics when environments fluctuate (Chesson *et al.*, 2004).

Spatial niche partitioning is another mechanism. If species exploit water at different soil depths, there may be reductions in interspecific competition mediated by shifts in species' water use efficiencies or timing of water use (as droughts affect each depth differently) and potential impacts on ecosystem water use and community diversity. Evidence for complementarity and partitioning remains mixed (Verheyen *et al.*, 2008; Bachmann *et al.*, 2015; Guderle *et al.*, 2018). Alternatively, hydraulic redistribution of moisture between soil depths (Richards & Caldwell, 1987; Caldwell *et al.*, 1998) could instead drive facilitation or increased interspecific competition (Dawson, 1993). Deeply rooted plants have the potential to engineer the thermal environment of whole communities, not only by providing added

soil water via hydraulic lift (Dawson, 1993) but also by modifying the humidity of the shared near-surface environment in a community.

The storage effect is an additional mechanism that can operate if water availability varies over time or space. If some species have differential responses to water availability, then in 'good' years they may be able to store these gains as larger population sizes or seedbanks, while having a relatively small negative effect on their populations in 'bad' years. This mechanism is known to operate in Sonoran Desert annual plants (Venable & Pake, 1999), which vary widely in their water use efficiency (Angert *et al.*, 2009). Succulents might also be able to store water for multiple months or years before using it, yielding a lagged version of the storage effect. This population buffering mechanism means that species that have nonoptimal water use within bad years may be buffered from negative consequences due to their success in good years.

## C2. Hoarding and spite

In other cases, fitness may not be maximized by coexistence, but rather by resource preemption (Fig. 2e). Species may succeed by using soil water to cause the local extinction of other species. The balance between facilitation and competition is known to shift as soil water becomes more limiting (Holmgren *et al.*, 1997; Haberstroh & Werner, 2022). Resource preemption may thus arise only under stressful circumstances (i.e. not likely in very wet environments, or in those with many species present, or for small-sized plants).

Resource preemption can occur via hoarding. Plants could rapidly acquire soil moisture and then store it in tissues or via capacitance, after which they can transpire it at any future time while also preventing other plants from using it, as might occur for desert plants like agaves, various columnar cacti, or baobabs. Alternatively, they could redistribute it to deeper soil (hydraulic descent) and make it unavailable to shallow-rooted competitors but available to themselves. This may occur primarily in environments where precipitation events are rare. Hoarding does occur for carbon and other nutrients in plant–fungal interactions, wherein fungi withhold transfer of mineral nutrients until the plant provides a higher amount of carbon in return (van't Padje *et al.*, 2021a,b), suggesting hoarding may also occur among plants.

Resource preemption can also occur via spite. Aggressive water use can manifest as exuberant transpiration. General theory for spite has been developed (Hamilton, 1970), but these ideas are not yet included in models of plant water use. However, if a species uses soil water quickly, it could make that water unavailable to other species. Spite would only be successful in cases where one species is able to cause greater harm to all other species than it causes to itself through such aggressive soil water use. Such an idea is consistent with the minimum resource level concepts in  $R^*$  resource competition theory (Tilman, 1982). A consequential hypothesis is that high nighttime transpiration (Wang *et al.*, 2021) or hydraulic descent (Hultine *et al.*, 2003) may be examples of spiteful water use by drought-tolerant species to harm drought-intolerant species.

## C3. Bluffing and learning

In a community context, if all species benefit from using a common resource (water), the optimal strategies (at individual or population level) may differ strongly from those in a single-species context. From this perspective, water use strategies could be viewed as a game in which the opponent's strategy is only partially observable. The game can have a Nash equilibrium, that is, a stable outcome where each species is better off retaining its strategy than switching to a different strategy. This has been demonstrated in one stomatal regulation model (Wolf et al., 2016) where competition causes changes in water use efficiency under drought, and has been considered in other allocation models (Farrior et al., 2013; Lu et al., 2020). It has also been shown empirically in Pinus taeda (Zenes et al., 2020), Lactuca sativa (lettuce) and Solanum lycopersicum (tomato; Vysotskaya et al., 2011). The broader game theory of competitive or cooperative depletion of water resources has also been considered in human contexts (Madani, 2010) but not widely applied to plants.

In a Nash equilibrium, the optimal strategies are often mixed, in that they involve randomizing choice of action over some probability distribution. This occurs because the actions taken by other species may influence the value of the actions taken by a focal species; additionally, each individual may have limited information available about the resources available to other individuals and may not know their actual strategy. Rather, each individual may only be able to respond to observed sequences of actions taken by another individual, for example, via rhizosphere processes including sensing soil moisture drawdown near roots, or chemosensing of other species' root hormones and exudates (Jackson, 1997; Hinsinger *et al.*, 2005) or of other volatile cues (Jin *et al.*, 2021).

In simple terms, this means that, in community contexts, species may deploy unusual behaviors. They may bluff with their water use, by using more or less water than they would otherwise, to send a false signal to others or force others to take certain actions. They may also invest in learning, by exploring the environment through yet-unknown means to better predict the current or future levels of water availability. For example, species that pre-form buds years in advance like *Veratrum tenuipetalum* (Iler & Inouye, 2013) are effectively gambling on the future state of the environment being favorable for photosynthesis.

#### C4. Positive species interactions

A fundamental premise of much water use theory and the ideas outlined previously is that the most common interactions among plant species are negative and often competitive. However, the assumption of competition as the fundamental process may reflect our biases more than reality (Simha *et al.*, 2022). Instead, beginning from a viewpoint of abundance (Kimmerer, 2020) and mutualism (Bronstein, 2015), sharing water may actually be mutually beneficial in many cases, especially when indirect interactions between plants occur that are mediated by other species (Fig. 2f). As conceptual examples, a first species could provide water to a second species that in turn transpires it and provides beneficial cool and

shaded conditions to the first species; or the second species could attract insects that would also pollinate or defend the first species.

Species may use water in ways that appear nonoptimal from an individual performance perspective or from a competitive perspective, because they produce cooperative (positive) interactions. Species that are most able to access water could redistribute it to other species that need it (e.g. via hydraulic lift, or evaporative cooling affecting a whole community), and in turn receive a nonwater benefit they cannot provide for themself. Temperate trees using water to support their own growth provide shade to some other species that in turn grow faster than they would alone (Kothari *et al.*, 2021). Or, savanna trees bring up water from deeper soil layers and redistribute it in the more superficial root zone (i.e. hydraulic lift), improving grass quality and attracting herbivores (Treydte *et al.*, 2007, 2011). Grass decomposition and addition of urine and feces may in turn improve nutrient availability for the trees.

## C5. Negative species interactions

Mycorrhizal fungi have been implicated in changes in various aspects of plant water use, including increased aquaporin function, and increased root hydraulic conductivity, generally resulting in improved performance of mycorrhizal plants under drought (Lehto & Zwiazek, 2011). They thus generally are considered to yield positive interactions with plants. However, in the process of maximizing their own fitness, these fungi could lead a plant into adopting a nonoptimal water use strategy in a parasitic interaction. Mycorrhizal fungi are known to exploit multiple species of plants to extract the best price for their mineral nutrients when demand is highest (Whiteside et al., 2019). Direct transfer of water taken up by arbuscular mycorrhizal fungi to host plants has also recently been demonstrated (Kakouridis et al., 2022). Increases in plant transpiration and photosynthesis also have been observed when fungal partners are present for Linum usitatissimum (flax; Drüge & Schonbeck, 1993) and Citrus jambhiri (rough lemon; Levy et al., 1983), suggesting that some aspects of plant water use may be mediated by exchanges with fungal partners. Whether and when these interactions are negative instead of positive remains underexplored.

Endophytes, organisms living inside leaves, may also have negative interactions with host plants through causing nonoptimal water use for their hosts. Minimum leaf conductance is almost twice as high in *Theobroma cacao* (cacao) leaves with natural abundances of endophytes than in leaves without endophytes (Arnold & Engelbrecht, 2007). All plants in natural ecosystems contain leaf endophytes (Rodriguez *et al.*, 2009), but densities and species assemblages vary widely, even across leaves within individual trees (Arnold & Herre, 2003). Water loss through leaf cuticles or from leaky stomata may thus be affected by the abundance and perhaps identity and functional properties of leaf endophytes, independent of the short-term carbon economy of the leaves.

Epiphylls, plants growing on top of the leaf surface can also have negative interactions with their host plants through water use. As leaves of evergreen plants age, they may accumulate epiphylls to the extent that exposed leaves only receive 15–45% of the sunlight they would receive in the absence of epiphylls (Coley *et al.*, 1993). As a result, the photosynthetic capacity of these shaded leaves is reduced (Anthony *et al.*, 2002) and presumably also their water use. The retention of such leaves that are unlikely to fix significantly more carbon than they use in respiration appears nonoptimal from both a carbon and water use perspective, unless there are other not yet understood benefits of epiphyll presence, for example, uptake of foliar water (Rosado & Almeida, 2020) or nitrogen (Bentley, 1987), or benefits of maintaining old leaves, for example, taking up space and shading competitors.

# Implications for leaf and earth system models under climate change

These hypotheses, if supported widely by empirical data, would require substantial revisions to extant water use theory and the leaf and earth system models they are used within. Many process-based leaf models use the Ball *et al.* (1987) or Medlyn *et al.* (2011) representation of stomatal conductance, in which stomatal conductance is calculated based on a suite of environmental variables, and notably, the photosynthetic rate and a parameter that identifies the relationship between stomatal conductance and photosynthesis (often referred to as  $g_1$  in these models, representing the inverse of water use efficiency). The  $g_1$  value can change within and across species, and by plant functional category if used in a larger-scale model. The value may also respond to processes described by any of the above hypotheses, though such effects are not yet implemented.

Errors in the plant water use theory currently used in many Earth System Models (ESMs) can have important consequences for understanding changes in water resources, heat and precipitation extremes, and ecosystem functioning under changing climate. This is because the soil–plant–atmosphere continuum representations in ESMs often use similar or the same marginal gain theory used in leaf-scale models to calculate ecosystem stomatal conductance (Clark et al., 2011; Massoud et al., 2019; Koven et al., 2020). There are known inaccuracies in ecosystem water fluxes simulated by ESMs (Li et al., 2021), which ultimately can influence other simulated quantities like soil moisture and precipitation. Even when average patterns and trends are simulated reasonably well, inaccuracies can manifest in other functions and time scales, such as predicting future heatwave intensities and durations (Kala et al., 2016).

A key question is whether adding complexity to ESMs would be useful when upscaled predictions of water flux are needed (Medlyn et al., 2017). Because ecosystem scale predictions are already usually successful and are made without including detailed vegetation dynamics, it is unlikely that the population and community ecology hypotheses we outlined would require incorporation into theory used in ESMs. However, the extreme environment hypotheses should likely be integrated into water use theory given existing evidence that current theory fails in some of these cases, and because of the forecasted higher prevalence of extreme environments in the near future. We suggest that incorporation of cooling (E1) into theory will be most productive. However, as ESMs

continue to be used to answer more complex questions related to feedback between vegetation and climate, some of the population and community context hypotheses may also become relevant. We speculate that trade-offs among fitness components (P4) and resource hoarding (C2) are the two highest priorities.

#### Conclusion

We have highlighted numerous hypotheses that indicate the multiple values of water to plants and that have varying degrees of support based on empirical observations. Not all these hypotheses may be common, nor equally important. However, in rejecting or supporting the hypotheses underlying them, we may come closer to a more complete theory of plant water use. More significantly, many of these hypotheses may be relevant for some species and not others, or in some environments or communities and not others, suggesting that it will be difficult to generate a single water use theory applicable in all situations.

We are not seeking to challenge the relevance of extant theory. It is correct, or close enough to correct, for most scenarios that have been of interest to date. Rather, we aim to highlight the many additional values of water relevant to more types of plants and environments than this theory has considered, and in doing so, to challenge the implicit use of simple economic ideas within ecophysiology. The alternative ideas we proposed may have potentially large consequences for individual plants, for community dynamics, and for ecosystem fluxes. By expanding beyond ecophysiology to incorporate population and community ecology – with all their uncertainties and complexities, we may achieve a more complete understanding of plant water use.

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## **Competing interests**

None declared.

#### **Author contributions**

BWB conceived the manuscript and wrote the first draft. BCP, MS and BWB created the figures. LMTA, KRH, DL, STM, BCP, MS and KW contributed sections of the text.

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#### References

- Angert AL, Huxman TE, Chesson P, Venable DL. 2009. Functional tradeoffs determine species coexistence via the storage effect. Proceedings of the National Academy of Sciences, USA 106: 11641–11645.
- Anthony PA, Holtum JAM, Jackes BR. 2002. Shade acclimation of rainforest leaves to colonization by lichens. Functional Ecology 16: 808–816.
- Aparecido LM, Woo S, Suazo C, Hultine KR, Blonder B. 2020. High water use in desert plants exposed to extreme heat. *Ecology Letters* 23: 1189–1200.
- Arnold AE, Engelbrecht BM. 2007. Fungal endophytes nearly double minimum leaf conductance in seedlings of a neotropical tree species. *Journal of Tropical Ecology* 23: 369–372.
- Arnold AE, Herre EA. 2003. Canopy cover and leaf age affect colonization by tropical fungal endophytes: ecological pattern and process in *Theobroma cacao* (Malvaceae). *Mycologia* 95: 388–398.
- Aronson J, Kigel J, Shmida A, Klein J. 1992. Adaptive phenology of desert and Mediterranean populations of annual plants grown with and without water stress. *Oecologia* 89: 17–26.
- Asseng S, Van Herwaarden AF. 2003. Analysis of the benefits to wheat yield from assimilates stored prior to grain filling in a range of environments. *Plant and Soil* 256: 217–229.
- Bachmann D, Gockele A, Ravenek JM, Roscher C, Strecker T, Weigelt A, Buchmann N. 2015. No evidence of complementary water use along a plant species richness gradient in temperate experimental grasslands. *PLoS ONE* 10: e0116367.
- Ball JT, Woodrow IE, Berry JA. 1987. A model predicting stomatal conductance and its contribution to the control of photosynthesis under different environmental conditions. In: *Progress in photosynthesis research*. Dordrecht, the Netherlands: Springer, 221–224.
- Bentley BL. 1987. Nitrogen fixation by epiphylls in a tropical rainforest. *Annals of the Missouri Botanical Garden* 74: 234–241.
- Blasini DE, Koepke DF, Bush SE, Allan GJ, Gehring CA, Whitham TG, Day TA, Hultine KR. 2022. Tradeoffs between leaf cooling and hydraulic safety in a

- dominant arid land riparian tree species. Plant, Cell & Environment 45: 1664–1681.
- Blonder B, Michaletz ST. 2018. A model for leaf temperature decoupling from air temperature. *Agricultural and Forest Meteorology* 262: 354–360.
- Boyle EA, Li YI, Pritchard JK. 2017. An expanded view of complex traits: from polygenic to omnigenic. *Cell* 169: 1177–1186.
- Brix H, Sorrell BK, Orr PT. 1992. Internal pressurization and convective gas flow in some emergent freshwater macrophytes. *Limnology and Oceanography* 37: 1420– 1433.
- Bronson DR, English NB, Dettman DL, Williams DG. 2011. Seasonal photosynthetic gas exchange and water-use efficiency in a constitutive CAM plant, the giant saguaro cactus (*Carnegiea gigantea*). *Oecologia* 167: 861–871.
- Bronstein JL. 2015. Mutualism. Cary, NC, USA: Oxford University Press.
- Bueno A, Alfarhan A, Arand K, Burghardt M, Deininger A-C, Hedrich R, Leide J, Seufert P, Staiger S, Riederer M. 2019. Temperature effects on the cuticular transpiration barrier of two desert plants with water-spender and water-saver life strategies. *Journal of Experimental Botany*. 70: 1613–1625.
- von Caemmerer S, Evans JR. 1991. Determination of the average partial pressure of CO<sub>2</sub> in chloroplasts from leaves of several C<sub>3</sub> plants. Functional Plant Biology 18: 287–305.
- Caldwell MM, Dawson TE, Richards JH. 1998. Hydraulic lift: consequences of water efflux from the roots of plants. *Oecologia* 113: 151–161.
- Castorena M, Olson ME, Enquist BJ, Fajardo A. 2022. Toward a general theory of plant carbon economics. *Trends in Ecology & Evolution* 37: 829–837.
- Chabot BF, Hicks DJ. 1982. The ecology of leaf life spans. Annual Review of Ecology and Systematics 13: 229–259.
- Chaves MM, Costa JM, Zarrouk O, Pinheiro C, Lopes CM, Pereira JS. 2016. Controlling stomatal aperture in semi-arid regions the dilemma of saving water or being cool? *Plant Science* 251: 54–64.
- Chesson P. 2000. Mechanisms of maintenance of species diversity. Annual Review of Ecology and Systematics 31: 343–366.
- Chesson P, Gebauer RL, Schwinning S, Huntly N, Wiegand K, Ernest MS, Sher A, Novoplansky A, Weltzin JF. 2004. Resource pulses, species interactions, and diversity maintenance in arid and semi-arid environments. *Oecologia* 141: 236– 253.
- Chiariello NR, Field CB, Mooney HA. 1987. Midday wilting in a tropical pioneer tree. *Functional Ecology* 1: 3–11.
- Christoffersen BO, Gloor M, Fauset S, Fyllas NM, Galbraith DR, Baker TR, Kruijt B, Rowland L, Fisher RA, Binks OJ et al. 2016. Linking hydraulic traits to tropical forest function in a size-structured and trait-driven model (TFS vs 1-Hydro). Geoscientific Model Development 9: 4227–4255.
- Clark D, Mercado L, Sitch S, Jones C, Gedney N, Best M, Pryor M, Rooney GG, Essery RLH, Blyth E et al. 2011. The Joint UK Land Environment Simulator (JULES), model description–part 2: carbon fluxes and vegetation dynamics. Geoscientific Model Development 4: 701–722.
- Coley PD, Kursar TA, Machado J-L. 1993. Colonization of tropical rain forest leaves by epiphylls: effects of site and host plant leaf lifetime. *Ecology* 74: 619–623.
- Costanza R, d'Arge R, De Groot R, Farber S, Grasso M, Hannon B, Limburg K, Naeem S, O'Neill RV, Paruelo J et al. 1997. The value of the world's ecosystem services and natural capital. Nature 387: 253–260.
- Coupel-Ledru A, Lebon E, Christophe A, Gallo A, Gago P, Pantin F, Doligez A, Simonneau T. 2016. Reduced nighttime transpiration is a relevant breeding target for high water-use efficiency in grapevine. *Proceedings of the National Academy of Sciences, USA* 113: 8963–8968.
- Cowan IR, Farquhar GD. 1977. Stomatal function in relation to leaf metabolism and environment. Symposia of the Society for Experimental Biology 31: 471–505.
- Dawson TE. 1993. Hydraulic lift and water use by plants: implications for water balance, performance and plant–plant interactions. *Oecologia* 95: 565–574.
- Dawson TE, Burgess SS, Tu KP, Oliveira RS, Santiago LS, Fisher JB, Simonin KA, Ambrose AR. 2007. Nighttime transpiration in woody plants from contrasting ecosystems. *Tree Physiology* 27: 561–575.
- Detto M, Wright SJ, Calderón O, Muller-Landau HC. 2018. Resource acquisition and reproductive strategies of tropical forest in response to the El Niño–Southern Oscillation. *Nature Communications* 9: 1–8.
- Drüge U, Schonbeck F. 1993. Effect of vesicular-arbuscular mycorrhizal infection on transpiration, photosynthesis and growth of flax (*Linum usitatissimum* L.) in relation to cytokinin levels. *Journal of Plant Physiology* 141: 40–48.

- Eamus D, Taylor DT, Macinnis-Ng C, Shanahan S, De Silva L. 2008. Comparing model predictions and experimental data for the response of stomatal conductance and guard cell turgor to manipulations of cuticular conductance, leaf-to-air vapour pressure difference and temperature: feedback mechanisms are able to account for all observations. Plant, Cell & Environment 31: 269-277.
- Falster DS, Reich PB, Ellsworth DS, Wright IJ, Westoby M, Oleksyn J, Lee TD. 2012. Lifetime return on investment increases with leaf lifespan among 10 Australian woodland species. New Phytologist 193: 409-419.
- Farrior CE, Dybzinski R, Levin SA, Pacala SW. 2013. Competition for water and light in closed-canopy forests: a tractable model of carbon allocation with implications for carbon sinks. American Naturalist 181: 314-330.
- Fatichi S, Leuzinger S, Körner C. 2014. Moving beyond photosynthesis: from carbon source to sink-driven vegetation modeling. New Phytologist 201: 1086-1095.
- Feng X, Lu Y, Jiang M, Katul G, Manzoni S, Mrad A, Vico G. 2022. Instantaneous stomatal optimization results in suboptimal carbon gain due to legacy effects. Plant, Cell & Environment 45: 3189-3204.
- Fisher RA, Koven CD, Anderegg WR, Christoffersen BO, Dietze MC, Farrior CE, Holm JA, Hurtt GC, Knox RG, Lawrence PJ et al. 2018. Vegetation demographics in earth system models: a review of progress and priorities. Global Change Biology 24: 35-54.
- Franks PJ, Bonan GB, Berry JA, Lombardozzi DL, Holbrook NM, Herold N, Oleson KW. 2018. Comparing optimal and empirical stomatal conductance models for application in Earth system models. Global Change Biology 24: 5708-
- Franks PJ, Cowan IR, Farquhar GD. 1997. The apparent feedforward response of stomata to air vapour pressure deficit: information revealed by different experimental procedures with two rainforest trees. Plant, Cell & Environment 20: 142-145.
- Fullbrook E. 2004. A guide to what's wrong with economics. London, UK: Anthem Press.
- Gates DM. 1968. Transpiration and leaf temperature. Annual Review of Plant Physiology 19: 211-238.
- Gould SJ, Lewontin RC. 1979. The spandrels of san marco and the panglossian paradigm: a critique of the adaptationist programme. Proceedings of the Royal Society of London. Series B: Biological Sciences 205: 581-598.
- Grossiord C, Buckley TN, Cernusak LA, Novick KA, Poulter B, Siegwolf RT, Sperry JS, McDowell N. 2020. Plant responses to rising vapor pressure deficit. New Phytologist 226: 1550-1566.
- Guderle M, Bachmann D, Milcu A, Gockele A, Bechmann M, Fischer C, Roscher C, Landais D, Ravel O, Devidal S et al. 2018. Dynamic niche partitioning in root water uptake facilitates efficient water use in more diverse grassland plant communities. Functional Ecology 32: 214-227.
- Guo JS, Hultine KR, Koch GW, Kropp H, Ogle K. 2020. Temporal shifts in iso/ anisohydry revealed from daily observations of plant water potential in a dominant desert shrub. New Phytologist 225: 713-726.
- Haberstroh S, Werner C. 2022. The role of species interactions for forest resilience to drought. Plant Biology 24: 1098-1107.
- de Haldat du Lys A, Millan M, Barczi J-F, Caraglio Y, Midgley GF, Charles-Dominique T. 2022. If self-shading is so bad, why is there so much? Short shoots reconcile costs and benefits. New Phytologist 237: 1684-1695.
- Hamilton WD. 1970. Selfish and spiteful behaviour in an evolutionary model. Nature 228: 1218-1220.
- Harley PC, Thomas RB, Reynolds JF, Strain BR. 1992. Modelling photosynthesis of cotton grown in elevated CO<sub>2</sub>. Plant, Cell & Environment 15: 271-282.
- Harrison Day BL, Carins-Murphy MR, Brodribb TJ. 2022. Reproductive water supply is prioritized during drought in tomato. Plant, Cell & Environment 45: 69-
- Harrison SP, Cramer W, Franklin O, Prentice IC, Wang H, Brännström Å, Boer H, Dieckmann U, Joshi J, Keenan TF et al. 2021. Eco-evolutionary optimality as a means to improve vegetation and land-surface models. New Phytologist 231: 2125-2141.
- Hayes S, Schachtschabel J, Mishkind M, Munnik T, Arisz SA. 2021. Hot topic: thermosensing in plants. Plant, Cell & Environment 44: 2018–2033.
- Hinsinger P, Gobran GR, Gregory PJ, Wenzel WW. 2005. Rhizosphere geometry and heterogeneity arising from root-mediated physical and chemical processes. New Phytologist 168: 293-303.

- Hollis M, Nell E. 1975. Rational economic man: a philosophical critique of neoclassical economics. Cambridge, UK: Cambridge University Press.
- Holmgren M, Scheffer M, Huston MA. 1997. The interplay of facilitation and competition in plant communities. Ecology 78: 1966-1975.
- Hultine KR, Cable WL, Burgess SSO, Williams DG. 2003. Hydraulic redistribution by deep roots of a Chihuahuan Desert phreatophyte. Tree Physiology 23: 353-360.
- Hunt JR, Hayman PT, Richards RA, Passioura JB. 2018. Opportunities to reduce heat damage in rain-fed wheat crops based on plant breeding and agronomic management. Field Crops Research 224: 126-138.
- Iler AM, Inouye DW. 2013. Effects of climate change on mast-flowering cues in a clonal montane herb, Veratrum tenuipetalum (Melanthiaceae). American Journal of Botany 100: 519-525.
- Ishii HT, Tanabe S, Hiura T. 2004. Exploring the relationships among canopy structure, stand productivity, and biodiversity of temperate forest ecosystems. Forest Science 50: 342-355.
- Jackson M. 1997. Hormones from roots as signals for the shoots of stressed plants. Trends in Plant Science 2: 22-28.
- Jackson T. 2016. Prosperity without growth: foundations for the economy of tomorrow. Abingdon, UK: Routledge.
- Jevons WS. 1879. The theory of political economy. London, UK: Macmillan.
- Jin J, Zhao M, Gao T, Jing T, Zhang N, Wang J, Zhang X, Huang J, Schwab W, Song C. 2021. Amplification of early drought responses caused by volatile cues emitted from neighboring tea plants. Horticulture Research 8: 243.
- Kakouridis A, Hagen JA, Kan MP, Mambelli S, Feldman LJ, Herman DJ, Weber PK, Pett-Ridge J, Firestone MK. 2022. Routes to roots: direct evidence of water transport by arbuscular mycorrhizal fungi to host plants. New Phytologist 236: 210-221.
- Kala J, De Kauwe MG, Pitman AJ, Medlyn BE, Wang Y-P, Lorenz R, Perkins-Kirkpatrick SE. 2016. Impact of the representation of stomatal conductance on model projections of heatwave intensity. Scientific Reports 6: 23418.
- Kannenberg SA, Guo JS, Novick KA, Anderegg WR, Feng X, Kennedy D, Konings AG, Martínez-Vilalta J, Matheny AM. 2022. Opportunities, challenges and pitfalls in characterizing plant water-use strategies. Functional Ecology 36: 24-37.
- Karban R. 2015. Plant sensing and communication. In: Plant sensing and communication. Chicago, IL, USA: University of Chicago Press.
- Kikuzawa K, Lechowicz MJ. 2006. Toward synthesis of relationships among leaf longevity, instantaneous photosynthetic rate, lifetime leaf carbon gain, and the gross primary production of forests. American Naturalist 168: 373-383.
- Kimmerer RW. 2020. The serviceberry: an economy of abundance. Emergence
- Klein T, Zeppel MJB, Anderegg WRL, Bloemen J, De Kauwe MG, Hudson P, Ruehr NK, Powell TL, von Arx G, Nardini A. 2018. Xylem embolism refilling and resilience against drought-induced mortality in woody plants: processes and trade-offs. Ecological Research 33: 839-855.
- Kolb PF, Robberecht R. 1996. High temperature and drought stress effects on survival of Pinus ponderosa seedlings. Tree Physiology 16: 665-672.
- Kostaki K-I, Coupel-Ledru A, Bonnell VC, Gustavsson M, Sun P, McLaughlin FJ, Fraser DP, McLachlan DH, Hetherington AM, Dodd AN et al. 2020. Guard cells integrate light and temperature signals to control stomatal aperture. Plant Physiology 182: 1404-1419.
- Kothari S, Montgomery RA, Cavender-Bares J. 2021. Physiological responses to light explain competition and facilitation in a tree diversity experiment. Journal of Ecology 109: 2000-2018.
- Koven CD, Knox RG, Fisher RA, Chambers JQ, Christoffersen BO, Davies SJ, Detto M, Dietze MC, Faybishenko B, Holm J et al. 2020. Benchmarking and parameter sensitivity of physiological and vegetation dynamics using the Functionally Assembled Terrestrial Ecosystem Simulator (FATES) at Barro Colorado Island, Panama. Biogeosciences 17: 3017-3044.
- Lehto T, Zwiazek JJ. 2011. Ectomycorrhizas and water relations of trees: a review. Mycorrhiza 21: 71-90.
- Leuning R. 1995. A critical appraisal of a combined stomatal-photosynthesis model for C3 plants. Plant, Cell & Environment 18: 339-355.
- Levy Y, Syvertsen JP, Nemec S. 1983. Effect of drought stress and vesiculararbuscular mycorrhiza on citrus transpiration and hydraulic conductivity of roots. New Phytologist 93: 61–66.

- Li J, Miao C, Wei W, Zhang G, Hua L, Chen Y, Wang X. 2021. Evaluation of CMIP6 global climate models for simulating land surface energy and water fluxes during 1979–2014. *Journal of Advances in Modeling Earth Systems* 13: e2021MS002515.
- Li WD, Duan W, Fan PG, Yan ST, Li SH. 2007. Photosynthesis in response to sink source activity and in relation to end products and activities of metabolic enzymes in peach trees. *Tree Physiology* 27: 1307–1318.
- van Loon LC. 2016. The intelligent behavior of plants. *Trends in Plant Science* 21: 286–294
- Lu Y, Duursma RA, Farrior CE, Medlyn BE, Feng X. 2020. Optimal stomatal drought response shaped by competition for water and hydraulic risk can explain plant trait covariation. *New Phytologist* 225: 1206–1217.
- Machado R, Loram-Lourenço L, Farnese FS, Alves RDFB, de Sousa LF, Silva FG, Filho SCV, Torres-Ruiz JM, Cochard H, Menezes-Silva PE. 2021. Where do leaf water leaks come from? Trade-offs underlying the variability in minimum conductance across tropical savanna species with contrasting growth strategies. *New Phytologist* 229: 1415–1430.
- Madani K. 2010. Game theory and water resources. Journal of Hydrology 381: 225–238
- Marchin RM, Backes D, Ossola A, Leishman MR, Tjoelker MG, Ellsworth DS. 2022a. Extreme heat increases stomatal conductance and drought-induced mortality risk in vulnerable plant species. Global Change Biology 28: 1133–1146.
- Marchin RM, Esperon-Rodriguez M, Tjoelker MG, Ellsworth DS. 2022b. Crown dieback and mortality of urban trees linked to heatwaves during extreme drought. *Science of the Total Environment* 850: 157915.
- Massoud EC, Xu C, Fisher RA, Knox RG, Walker AP, Serbin SP, Christoffersen BO, Holm JA, Kueppers LM, Ricciuto DM et al. 2019. Identification of key parameters controlling demographically structured vegetation dynamics in a land surface model: CLM4.5 (FATES). Geoscientific Model Development 12: 4133–4164
- Medlyn BE, de Kauwe MG, Lin Y-S, Knauer J, Duursma RA, Williams CA, Arneth A, Clement R, Isaac P, Limousin JM et al. 2017. How do leaf and ecosystem measures of water-use efficiency compare? New Phytologist 216: 758–770.
- Medlyn BE, Duursma RA, Eamus D, Ellsworth DS, Prentice IC, Barton CV, Crous KY, de Angelis P, Freeman M, Wingate L. 2011. Reconciling the optimal and empirical approaches to modelling stomatal conductance. *Global Change Biology* 17: 2134–2144.
- Michaletz ST, Johnson EA, Tyree MT. 2012. Moving beyond the cambium necrosis hypothesis of post-fire tree mortality: cavitation and deformation of xylem in forest fires. *New Phytologist* 194: 254–263.
- Miller BD, Carter KR, Reed SC, Wood TE, Cavaleri MA. 2021. Only sun-lit leaves of the uppermost canopy exceed both air temperature and photosynthetic thermal optima in a wet tropical forest. Agricultural and Forest Meteorology 301: 108347.
- Monteith JL, Unsworth MH. 2013. Principles of environmental physics, 4th edn. Boston, MA, USA: Academic Press.
- Muhammad Aslam M, Waseem M, Jakada BH, Okal EJ, Lei Z, Saqib HSA, Yuan W, Xu W, Zhang Q. 2022. Mechanisms of abscisic acid-mediated drought stress responses in plants. *International Journal of Molecular Sciences* 23: 1084.
- Oliveira RS, Eller CB, Barros FDV, Hirota M, Brum M, Bittencourt P. 2021. Linking plant hydraulics and the fast–slow continuum to understand resilience to drought in tropical ecosystems. *New Phytologist* 230: 904–923.
- van't Padje A, Oyarte Galvez L, Klein M, Hink MA, Postma M, Shimizu T, Kiers ET. 2021a. Temporal tracking of quantum-dot apatite across *in vitro* mycorrhizal networks shows how host demand can influence fungal nutrient transfer strategies. *The ISME Journal* 15: 435–449.
- van't Padje A, Werner GDA, Kiers ET. 2021b. Mycorrhizal fungi control phosphorus value in trade symbiosis with host roots when exposed to abrupt 'crashes' and 'booms' of resource availability. New Phytologist 229: 2933–2944.
- Peñuelas J, Filella I, Zhang X, Llorens L, Ogaya R, Lloret F, Comas P, Estiarte M, Terradas J. 2004. Complex spatiotemporal phenological shifts as a response to rainfall changes. *New Phytologist* 161: 837–846.
- Quereix A, Dewar RC, Gaudillere J-P, Dayau S, Valancogne C. 2001. Sink feedback regulation of photosynthesis in vines: measurements and a model. *Journal of Experimental Botany* 52: 2313–2322.
- Querejeta JI, Ren W, Prieto I. 2021. Vertical decoupling of soil nutrients and water under climate warming reduces plant cumulative nutrient uptake, water-use efficiency and productivity. *New Phytologist* 230: 1378–1393.

- Reich PB, Stefanski A, Rich RL, Sendall KM, Wei X, Zhao C, Hou J, Montgomery RA, Bermudez R. 2021. Assessing the relevant time frame for temperature acclimation of leaf dark respiration: a test with 10 boreal and temperate species. *Global Change Biology* 27: 2945–2958.
- Reich PB, Uhl C, Walters MB, Prugh L, Ellsworth DS. 2004. Leaf demography and phenology in Amazonian rain forest: a census of 40 000 leaves of 23 tree species. *Ecological Monographs* 74: 3–23.
- Reid WV, Mooney HA, Cropper A, Capistrano D, Carpenter SR, Chopra K, Dasgupta P, Dietz T, Duraiappah AK, Hassan R et al. 2005. Ecosystems and human well-being-synthesis: a report of the millennium ecosystem assessment. Washington, DC, USA: Island Press.
- Richards JH, Caldwell MM. 1987. Hydraulic lift: substantial nocturnal water transport between soil layers by *Artemisia tridentata* roots. *Oecologia* 73: 486–489.
- Rodriguez RJ, White JF Jr, Arnold AE, Redman AR. 2009. Fungal endophytes: diversity and functional roles. New Phytologist 182: 314–330.
- Rosado BHP, Almeida LC. 2020. The importance of phyllosphere on foliar water uptake. *Trends in Plant Science* 25: 1058–1060.
- Schulze ED, ed. 1982. Plant life forms and their carbon, water and nutrient relations. In: *Physiological plant ecology II*. Berlin, Germany: Springer, 615– 676.
- Schulze E-D, Lange OL, Kappen L, Buschbom U, Evenari M. 1973. Stomatal responses to changes in temperature at increasing water stress. *Planta* 110: 29–42.
- Sen AK. 1977. Rational fools: a critique of the behavioral foundations of economic theory. *Philosophy & Public Affairs* 6: 317–344.
- Shivhare D, Mueller-Cajar O. 2017. *In vitro* characterization of thermostable CAM Rubisco activase reveals a Rubisco interacting surface loop. *Plant Physiology* 174: 1505–1516.
- Silvertown J, Gordon DM. 1989. A framework for plant behavior. Annual Review of Ecology and Systematics 20: 349–366.
- Simha A, Hoz CP-D, Carley L. 2022. Moving beyond the "diversity paradox": the limitations of competition-based frameworks in understanding species diversity. *American Naturalist* 200: 89–100.
- Sinha R, Zandalinas SI, Fichman Y, Sen S, Zeng S, Gómez-Cadenas A, Joshi T, Fritschi FB, Mittler R. 2022. Differential regulation of flower transpiration during abiotic stress in annual plants. New Phytologist 235: 611–629.
- Slot M, Garcia MN, Winter K. 2016. Temperature response of CO<sub>2</sub> exchange in three tropical tree species. *Functional Plant Biology* 43: 468–478.
- Slot M, Nardwattanawong T, Hernández GG, Bueno A, Riederer M, Winter K. 2021. Large differences in leaf cuticle conductance and its temperature response among 24 tropical tree species from across a rainfall gradient. New Phytologist 232: 1618–1631.
- Smith A. 1776. The wealth of nations. London, UK: W. Strahan & T. Cadell.Smith SD, Didden-Zopfy B, Nobel PS. 1984. High-temperature responses of North American cacti. *Ecology* 65: 643–651.
- Sperry JS, Wang Y, Wolfe BT, Mackay DS, Anderegg WR, McDowell NG, Pockman WT. 2016. Pragmatic hydraulic theory predicts stomatal responses to climatic water deficits. *New Phytologist* 212: 577–589.
- Swanson RH. 1994. Significant historical developments in thermal methods for measuring sap flow in trees. Agricultural and Forest Meteorology 72: 113–132.
- Teskey R, Wertin T, Bauweraerts I, Ameye M, McGuire MA, Steppe K. 2015. Responses of tree species to heat waves and extreme heat events. *Plant, Cell & Environment* 38: 1699–1712.
- Tilman D. 1982. Resource competition and community structure. Princeton, UK: Princeton University Press.
- Treydte AC, Heitkönig IM, Prins HH, Ludwig F. 2007. Trees improve grass quality for herbivores in African savannas. Perspectives in Plant Ecology, Evolution and Systematics 8: 197–205.
- Treydte AC, van der Beek JG, Perdok AA, van Wieren SE. 2011. Grazing ungulates select for grasses growing beneath trees in African savannas. *Mammalian Biology* 76: 345–350.
- Urban J, Ingwers MW, McGuire MA, Teskey RO. 2017. Increase in leaf temperature opens stomata and decouples net photosynthesis from stomatal conductance in *Pinus taeda* and *Populus deltoides* × *nigra*. *Journal of Experimental Botany* 68: 1757–1767.
- Veblen T. 1898. Why is economics not an evolutionary science? The Quarterly Journal of Economics 12: 373–397.

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- Venable DL, Pake CE. 1999. Population ecology of Sonoran Desert annual plants. In: *The ecology of Sonoran Desert plants and plant communities.* Tucson, AZ, USA: University of Arizona Press, 115–142.
- Venturas MD, Sperry JS, Hacke UG. 2017. Plant xylem hydraulics: what we understand, current research, and future challenges. *Journal of Integrative Plant Biology* 59: 356–389.
- Verheyen K, Bulteel H, Palmborg C, Olivié B, Nijs I, Raes D, Muys B. 2008. Can complementarity in water use help to explain diversity–productivity relationships in experimental grassland plots? *Oecologia* 156: 351–361.
- Vico G, Dralle D, Feng X, Thompson S, Manzoni S. 2017. How competitive is drought deciduousness in tropical forests? A combined eco-hydrological and ecoevolutionary approach. *Environmental Research Letters* 12: 65006.
- Vogel S. 2009. Leaves in the lowest and highest winds: temperature, force and shape. New Phytologist 183: 13–26.
- Vysotskaya L, Wilkinson S, Davies WJ, Arkhivpova T, Kudoyarova G. 2011. The effect of competition from neighbours on stomatal conductance in lettuce and tomato plants. *Plant, Cell & Environment* 34: 729–737.
- Wang Y, Anderegg WR, Venturas MD, Trugman AT, Yu K, Frankenberg C. 2021.

  Optimization theory explains nighttime stomatal responses. *New Phytologist* 230: 1550–1561.
- Wang Y, Sperry JS, Anderegg WR, Venturas MD, Trugman AT. 2020. A theoretical and empirical assessment of stomatal optimization modeling. *New Phytologist* 227: 311–325.
- Westoby M, Warton D, Reich PB. 2000. The time value of leaf area. *American Naturalist* 155: 649–656.
- Whiteside MD, Werner GD, Caldas VE, van't Padje A, Dupin SE, Elbers B, Bakker M, Wyatt GAK, Klein M, Hink MA et al. 2019. Mycorrhizal fungi

- respond to resource inequality by moving phosphorus from rich to poor patches across networks. *Current Biology* **29**: 2043–2050.
- Williams JW, Jackson ST. 2007. Novel climates, no-analog communities, and ecological surprises. Frontiers in Ecology and the Environment 5: 475–482.
- Winter K, Garcia M, Holtum JA. 2011. Drought-stress-induced up-regulation of CAM in seedlings of a tropical cactus, *Opuntia elatior*, operating predominantly in the  $C_3$  mode. *Journal of Experimental Botany* 62: 4037–4042.
- Wolf A, Anderegg WR, Pacala SW. 2016. Optimal stomatal behavior with competition for water and risk of hydraulic impairment. Proceedings of the National Academy of Sciences, USA 113: E7222–E7230.
- Yang J, Zhang J. 2006. Grain filling of cereals under soil drying. New Phytologist 169: 223–236.
- Yu K, Goldsmith GR, Wang Y, Anderegg WRL. 2019. Phylogenetic and biogeographic controls of plant nighttime stomatal conductance. *New Phytologist* 222: 1778–1788.
- Zenes N, Kerr KL, Trugman AT, Anderegg WR. 2020. Competition and drought alter optimal stomatal strategy in tree seedlings. Frontiers in Plant Science 11: 478.
- Zeppel MJB, Lewis JD, Phillips NG, Tissue DT. 2014. Consequences of nocturnal water loss: a synthesis of regulating factors and implications for capacitance, embolism and use in models. *Tree Physiology* 34: 1047–1055.

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